Systemic HIV and SIV latency reversal via non-canonical NF-κB signalling in vivo

Long-lasting, latently infected resting CD4+ T cells are the greatest obstacle to obtaining a cure for HIV infection, as these cells can persist despite decades of treatment with antiretroviral therapy (ART). Estimates indicate that more than 70 years of continuous, fully suppressive ART are needed to eliminate the HIV reservoir. Alternatively, induction of HIV from its latent state could accelerate the decrease in the reservoir, thus reducing the time to eradication. Previous attempts to reactivate latent HIV in preclinical animal models and in clinical trials have measured HIV induction in the peripheral blood with minimal focus on tissue reservoirs and have had limited effect. Here we show that activation of the non-canonical NF-κB signalling pathway by AZD5582 results in the induction of HIV and SIV RNA expression in the blood and tissues of ART-suppressed bone marrow–liver–thymus (BLT) humanized mice and rhesus macaques infected with HIV and SIV, respectively. Analysis of resting CD4+ T cells from tissues after AZD5582 treatment revealed increased SIV RNA expression in the lymph nodes of macaques and robust induction of HIV in almost all tissues analysed in humanized mice, including the lymph nodes, thymus, bone marrow, liver and lung. This promising approach to latency reversal—in combination with appropriate tools for systemic clearance of persistent HIV infection—greatly increases opportunities for HIV eradication.

Latently infected cells carrying an integrated replication-competent provirus that contribute to viral rebound after the interruption of ART (termed the HIV reservoir) are not detected and eliminated by the immune system or current therapeutics. Therefore, the HIV reservoir has been targeted by approaches to reverse latency and induce viral antigen production (that is, 'HIV reactivation') –, which renders infected cells susceptible to virus-induced cell death or clearance by the immune system. Previous approaches to HIV reactivation have been modestly effective and have not demonstrated reactivation of HIV in resting CD4+ T cells in tissues.

HIV induction in vitro by SMAC mimetics

The lack of specificity of molecules that activate the NF-κB pathway as latency-reversal agents (LRAs) often leads to toxicities that prevent clinical implementation. We tested the induction of HIV and...
ncNF-κB signalling pathway, AZD5582 has limited pleotropic effects, which may translate to fewer off-target effects.

**Latency reversal in BLT humanized mice**

BLT mice were infected with HIV-1JR-CSF (Supplementary Table 1) and supplied with ART11–18 (Fig. 2a, b). Mice then received a single intraperitoneal injection of 3 mg kg⁻¹ AZD5582 or vehicle. No changes in plasma HIV RNA levels were detected in vehicle-control-treated mice at 24 or 48 h nor in AZD5582-treated mice 24 h after AZD5582 administration (Fig. 2c). However, 48 h after AZD5582 treatment increased HIV RNA expression was detected in the plasma of 3 out of 6 (50%) and 3 out of 4 (75%) mice in two independent experiments (Fig. 2c). These data demonstrate that a single dose of AZD5582 can induce HIV production, resulting in significant viraemia (up to 1,574 HIV RNA copies per ml plasma) in ART-treated BLT mice (Supplementary Table 2).

The hallmark of HIV persistence in humans is the presence of inducible HIV in resting CD4⁺ T cells. Therefore, resting cells from primary (bone marrow and thymic organoid) and peripheral blood (lymph node and spleen) immune tissues were isolated from HIV-infected ART-suppressed BLT mice 48 h after treatment with vehicle control or AZD558219–22. The levels of HIV RNA in resting CD4⁺ T cells from AZD5582-treated mice were 11-fold (bone marrow, \( P = 0.0201 \)), 21-fold (thymic organoid, \( P = 0.0038 \)), 12-fold (lymph node, \( P = 0.0004 \)), 1.4-fold (spleen, \( P = 0.0426 \)), 24-fold (liver, \( P = 0.0145 \)) and 3.2-fold (lung, \( P = 0.0029 \)) higher than controls (Fig. 2d). These results were confirmed in a second independent experiment (Extended Data Fig. 2). No notable differences in cell-associated HIV DNA were noted between mice treated with vehicle control or AZD5582 (Supplementary Table 3). These results demonstrate that AZD5582 induces systemic HIV RNA production in resting CD4⁺ T cells, indicative of latency reversal in this important cellular source of persistent HIV infection. We also isolated cells from the peripheral blood, female reproductive tract and brain of HIV-infected ART-suppressed BLT mice 48 h after treatment with AZD5582 or vehicle control. As too few CD4⁺ T cells were available for cell sorting from these compartments, RNA was extracted from total cells and analysed for the presence of HIV RNA in each tissue. The levels of HIV RNA were significantly higher in the female reproductive tract (3.4-fold, \( P = 0.0152 \)) and brain (8.7-fold, \( P = 0.0147 \)) of AZD5582-treated mice compared with vehicle controls, but not in the blood (\( P = 0.3095 \)) (Fig. 2e). Together, these results show that AZD5582 treatment induces systemic HIV RNA production in BLT mice.

**Pharmacodynamics and safety in BLT mice**

Target engagement after treatment with AZD5582 was confirmed ex vivo by the degradation of cIAP1 (proximal) and p100 (distal), targets of SMAC in the ncNF-κB pathway (Extended Data Fig. 3a). In vivo target engagement was demonstrated in resting CD4⁺ T cells isolated from thymus, spleen, lymph nodes, liver, lung and bone marrow of BLT mice treated with a single dose of 3 mg kg⁻¹ AZD5582 or vehicle control (Extended Data Fig. 3b). These results were confirmed by immunohistochemical analysis of the thymic organoid of HIV-infected ART-suppressed BLT mice. This analysis showed a marked reduction in cIAP1 expression in the thymic organoid of AZD5582-treated mice (Extended Data Fig. 3c).

To study off-target or immune-mediated toxicities of AZD5582, we measured serum chemistry, T cell activation and a panel of plasma cytokines after in vivo treatment of immunocompetent BALB/c mice. AZD5582 administration resulted in mild and transient increases in alanine aminotransferase and aspartate aminotransferase that resolved a few days after treatment. No changes in serum chemistries were noted (Supplementary Table 4). In addition, no differences were noted in the levels of activated (CD38 'HLA-DR') CD4⁺ or CD8⁺ T cells in BLT mice treated with AZD5582 or vehicle control (Supplementary Table 5) or in plasma levels of 41 human cytokines and chemokines that serve as indicators of systemic
activation and inflammation \(^{31}\) (Supplementary Table 6). Together, these results demonstrate that AZD5582 does not cause generalized toxicity or activation of the immune system in the BLT model.

### Latency reversal in rhesus macaques

We next evaluated the latency-reversal activity of AZD5582 in 21 Mamu*08* and Mamu*17* rhesus macaques infected with SIV\(_{mac239}\) and treated with a potent ART regimen comprising tenofovir disoproxil fumarate, emtricitabine, and dolutegravir initiated 8 weeks after infection (Fig. 3a and Supplementary Table 7) \(^{22,23}\). Suppression of SIV viraemia below 60 copies per ml (standard assay limit of detection) was achieved in all macaques in 2–20 weeks and ART was continued for 55–67 weeks before further treatment (Fig. 3b). On the basis of pharmacokinetic and pharmacodynamic data from uninfected macaques (Extended Data Fig. 4a) as well as protocols for SMAC mimetics used in oncology, intravenous infusions of 0.1 mg kg\(^{-1}\) AZD5582 were administered weekly to 12 SIV-infected ART-suppressed rhesus macaques for 3 or 10 weeks (Fig. 3a). Nine SIV-infected ART-suppressed rhesus macaques served as controls (Fig. 3a). Plasma concentrations of AZD5582 measured after the first, third, sixth, and tenth dose showed that drug exposures in SIV-infected ART-suppressed macaques were consistent across the treatment period and comparable to those observed in uninfected rhesus macaques (Extended Data Fig. 4b).

Latency reversal, defined as on-ART viraemia increasing from less than 60 copies per ml of plasma to more than 60 copies per ml of plasma after AZD5582 treatment, was observed as early as 96 h after the first dose and reached levels as high as 1,390 copies per ml in SIV-infected rhesus macaques (Fig. 3c, d). On-ART viraemia >60 copies per ml of plasma was observed in 5 out of 12 rhesus macaques (42%), corresponding to 5 out of 9 rhesus macaques (55%) that received 10 doses of AZD5582 (Fig. 3c, d). Multiple instances of sustained viraemia >60 copies per ml between AZD5582 doses were observed. Out of 140 viral load measurements performed on the 5 macaques that exhibited on-ART viraemia during AZD5582 treatment, 64 were >60 copies per ml (46%); in the macaque with the greatest frequency of reactivation, this proportion was 15 out of 28 (53%). Longitudinal examination of plasma virus by single-genome sequencing analysis of the SIV\(_{mac239}\) env gene in all rhesus macaques that experienced AZD5582-induced on-ART viraemia was performed at four selected time points: 2 weeks after infection (near peak viraemia), 8 weeks after infection (immediately before ART initiation), and at 2 time points separated by 26–42 days during AZD5582 treatment. Phylogenetic analyses showed several patterns of virus reactivation (Extended Data Fig. 5). In two rhesus macaques (RD116 and RK116), most of the reactivated virus sequences were phylogenetically closer to sequences at eight weeks after infection rather than peak viraemia and were unique, indicating that the variants produced during AZD5582 treatment originated from multiple cells that were seeded at the time of ART initiation \(^{24}\). In two other rhesus macaques (RK116 and RD116), a large fraction of the viruses produced during AZD5582 treatment showed identical sequences, suggesting that latency reversal occurred from a single cell or a clonally expanded population of infected cells. These clones clustered with both peak and pre-ART time points and were accompanied by additional unique sequences. In one rhesus macaque (RLy15), a single virus sequence was amplified at each time point during AZD5582 treatment and these were both phylogenetically similar to sequences found before ART treatment. Taken together, these results indicate that AZD5582 induced virus reactivation from a diverse population of cells, some of which may be clonally expanded \(^{22,23}\).

We quantified cell-associated SIV RNA and SIV DNA in resting CD4\(^+\) T cells sorted from SIV-infected ART-suppressed rhesus macaques treated or not with AZD5582. Cell-associated SIV RNA levels in resting CD4\(^+\) T cells isolated from lymph nodes were significantly higher in macaques that received ten doses of AZD5582 compared with controls (P = 0.0148) (Fig. 3e). A similar trend was observed in resting CD4\(^+\) T cells isolated from the spleens of a subgroup of six macaques that were euthanized. Levels of cell-associated SIV DNA in resting CD4\(^+\) T cells were similar in each compartment across groups (Fig. 3e). To further understand whether latency reversal induced by AZD5582 resulted in a perturbation of the overall level of infected CD4\(^+\) T cells, we performed longitudinal measurements of cell-associated SIV DNA in total (rather than resting) CD4\(^+\) T cells isolated from lymph nodes and blood as well as quantitative viral outgrowth assays using CD4\(^+\) T cells from lymph nodes and spleen at the end of the treatment period (Extended
confirmed by western blot analysis of the degradation of p100 to p52.

Pharmacodynamics in rhesus macaques

Pharmacological target engagement of the ncNF-κB pathway was confirmed by western blot analysis of the degradation of p100 to p52 in lymph-node mononuclear cells after in vivo exposure to AZD5582 (Extended Data Fig. 4c) and in splenocytes treated ex vivo with AZD5582 (Extended Data Fig. 4d–g). Transcriptional profiling of CD4+ T cells from the peripheral blood and lymph nodes isolated from AZD5582-treated rhesus macaques showed a distinct effect of AZD5582 on gene expression based on principal component (Fig. 4a) and DAVID pathway (Extended Data Fig. 7a) analyses. Enrichment of NF-κB targets after AZD5582 treatment was demonstrated by gene-set enrichment analysis in lymph-node mononuclear cells after in vivo exposure to AZD5582.

Despite the high level of virus reactivation induced by AZD5582, these experiments did not reveal a consistent reduction in the total or replication-competent SIV reservoir compared with controls.
(GSEA) (Fig. 4b; a heat map of differentially expressed gene targets of NF-κB is shown in Extended Data Fig. 7b). The hallmark ncNF-κB signalling genes NFKB2 and RELB were significantly upregulated in CD4⁺ T cells from the peripheral blood and lymph nodes after AZD5582 treatment, whereas cNF-κB signalling molecules were mostly unafected except for the inhibitor NFKBIA (Fig. 4c). The baculoviral IAP repeating containing 3 (BIRC3) gene, which encodes cIAP2 and regulates both the cNF-κB and ncNF-κB pathways, was also significantly upregulated in CD4⁺ T cells from the peripheral blood and lymph nodes (Fig. 4c). Notably, activation of ncNF-κB signalling genes was evident in rhesus macaques with and without on-ART viremia of more than 60 copies per ml (Fig. 4c and Extended Data Fig. 7c) and GSEA showed similar changes in overall gene expression when AZD5582-treated rhesus macaques were grouped according to the presence or absence of on-ART viremia of more than 60 copies per ml (Supplementary Table 8), suggesting that the lack of a virological response measured by the standard viral load assay was not due to compromised target engagement.

We next used an ultrasensitive assay to measure SIV RNA in plasma, comparing 2–3 baseline time points during ART in the 4 weeks before AZD5582 treatment (when plasma SIV RNA was less than 60 copies per ml in all rhesus macaques using the standard viral load assay) and 3–4 time points during AZD5582 treatment (Extended Data Fig. 8a). We found significantly higher plasma SIV RNA values during the period of AZD5582 treatment compared with pre-treatment (Extended Data Fig. 8b, P = 0.008), with 8 out of 12 rhesus macaques demonstrating ≥2 SIV RNA measurements above the median of their baseline values. This group of eight rhesus macaques with evidence of latency reversal induced by AZD5582 could be segregated from the four rhesus macaques with unchanged viral loads based on two parameters: higher levels of plasma SIV RNA immediately before ART initiation and SIV DNA in peripheral blood (top) and lymph nodes (bottom) of SIV-infected rhesus macaques before treatment with AZD5582 and 48 h after the final dose (n = 12, red) and control rhesus macaques (n = 9, blue). Statistical significance was determined using a two-sided Mann–Whitney U-test.

Safety and immune effects in rhesus macaques

The potential for AZD5582 toxicity in SIV-infected ART-suppressed rhesus macaques was examined and transient increases in liver enzymes (aspartate aminotransferase and γ-glutamyltransferase) were observed (Extended Data Fig. 9a). Total white-blood cell counts were decreased in all rhesus macaques 48 h after the first dose of AZD5582 but returned to normal levels when measured before the second dose and over time (Extended Data Fig. 9b). After the seventh and eighth doses, one macaque experienced a reaction characterized by fever, emesis, fatigue and lack of appetite, and had elevated liver enzyme and creatinine levels and bandsaemia as shown by laboratory examination. All abnormalities resolved within a two-week period; however, this rhesus macaque did
not receive further doses. Notably, weights did not significantly fluctuate over the course of the treatment period (Extended Data Fig. 9c). In summary, 97 doses of AZD5582 were administered to 12 SIV-infected ART-suppressed rhesus macaques, of which 95 were well-tolerated and 2 resulted in a mild adverse reaction.

Markers of T cell activation (both CD4+ and CD8+) were assessed in AZD5582-treated rhesus macaques and controls, and expression of CCR5, HLA-DR and PD-1 on CD4+ T cells from the blood and lymph nodes was similar before and after treatment with AZD5582 (Fig. 4d, e). Levels of intracellular Ki-67 expression were increased in CD4+ T cells in the blood and CD8+ T cells in the blood and lymph nodes after AZD5582 treatment (Fig. 4d, e). CD8+ T cell expression of HLA-DR was higher in the blood after exposure to AZD5582 (Fig. 4e). These results indicate that AZD5582 does not induce global CD4+ T cell activation in rhesus macaques but the combined pharmacological and virological effects may have a stimulatory effect on CD8+ T cells. Longitudinal analyses of CD4+ T cell counts and frequencies, CD4+ T cell viability, CD4+ T cell subset frequencies and their Ki-67 expression are shown in Extended Data Fig. 10a–d. The increased Ki-67 observed within CD4+ T cells after AZD5582 treatment may provide a note of caution as the proliferation of latently infected memory CD4+ T cells is hypothesized to contribute to the maintenance of the viral reservoir over time25,27; however, we did not find an increase in infected cells after AZD5582 treatment (Fig. 3e and Extended Data Fig. 6).

We next determined whether AZD5582 would impair SIV-specific T cell responses in SIV-infected ART-suppressed rhesus macaques, a potential adverse outcome of treatment that has been suggested for other LRAs28. However, the frequency of SIV Gag- or Env-specific CD8+ T cells measured by IFNγ ELISPOT did not decrease after AZD5582 treatment (Extended Data Fig. 10e), and CD8+ T cell polyfunctionality and proliferative responses were largely unaffected by AZD5582 treatment ex vivo, with the exception of IL-2 IFNγ TNF triple-positive cells, which were slightly reduced (P = 0.0476) (Extended Data Fig. 10f, g). Furthermore, longitudinal assessment of plasma levels of inflammatory cytokines and chemokines did not reveal any marked changes induced by AZD5582 (Extended Data Fig. 10h). Overall, our work in the rhesus macaque model indicates that AZD5582 treatment is safe in most macaques and can induce appreciable increases in plasma SIV RNA as well as SIV RNA expression in resting CD4+ T cells from the lymph nodes during ART.

**Discussion**

Eradication of HIV infection after prolonged viral suppression is the focus of intense research and latency reversal has been a cornerstone in the search for HIV eradication approaches in humans. Various approaches have been investigated, including latency-reversing agents and implications for drug and treatment design. JCI Insight 3, e120352 (2018).

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Methods

Experimental design
The purpose of this study was to determine the efficacy of the SMAC-mimetic AZD5882 as an HIV and SIV LRA in vivo. To this end, two animal models of HIV-1 infection were used: the HIV-infected BLT humanized mouse model and the SIV-infected rhesus macaque (Macaca mulatta) nonhuman primate (NHP) model. In each model, animals were infected with HIV-1 or SIV and viremia was durably suppressed with ART. AZD5882 was administered to ART-suppressed animals that were then assessed for changes associated with the reactivation of the viral reservoir. Mice were maintained under specific-pathogen-free conditions by the Division of Comparative Medicine at the University of North Carolina, Chapel Hill. Mouse experiments were conducted in accordance with NIH guidelines for the housing and care of laboratory animals and in accordance with protocols reviewed and approved by the Institutional Animal Care and Use Committee (IACUC) at the University of North Carolina, Chapel Hill. Healthy rhesus macaques for pharmacokinetic studies were housed at GloxsoSmithKline and all procedures were conducted in accordance with the GloxsoSmithKline policy on the care, welfare and treatment of laboratory animals and were reviewed by the IACUC at GloxsoSmithKline. Rhesus macaques infected with SIV were housed at the Yerkes National Primate Research Center and treated in accordance with Emory University and Yerkes National Primate Research Center IACUC regulations (PROTO201800308). Animal-care facilities are accredited by the US Department of Agriculture and the Association for Assessment and Accreditation of Laboratory Animal Care International.

Preparation of Jurkat HIV–luciferase cell clones
Cell clones Jurkat-C16 and Jurkat-I15 were prepared by infecting Jurkat cells (Jurkat clone E6-1 cells, American Type Culture Collection TIB-152, authenticated by morphological identification and virus-susceptibility profiles, tested for mycoplasma by the supplier) with a full-length, infectious HIV-1NL4-3-based virus engineered to express a luciferase reporter in place of the HIV-1 nef gene (NLCH-Luci). The Jurkat-N6 cell clone was generated using the same virus as described above with an additional mouse heat-stable antigen (HSA) reporter located just downstream of the luciferase open-reading frame and separated by a T2A element (NLCH-Luci-HSA). NLCH, provided by R. Swanstrom, is the parent molecular infectious clone used to make the Jurkat clones and is a modification of HIV-1NL4-3 (GenBank U26942) in which flanking sequences were removed. All viruses were derived by transfection of human embryonic kidney 293 cells (HEK 293T, European Collection of Authenticated Cell Cultures, authenticated by morphological identification, tested for mycoplasma by the supplier) with 1 μg of the HIV-1NL4-3-derived infectious molecular plasmid DNA using the FuGENE HD Transfection reagent (Promega) according to the manufacturer’s recommendations. Supernatants were collected 48 h after transfection, passed through a 0.2-μm filter, and used to infect wild-type Jurkat cells. After infection, cells expressing high levels of HIV-encoded mouse HSA were removed using biotin-labelled rat anti-mouse CD24 antibody (clone MI/69, BD Biosciences) that was adsorbed to streptavidin-labelled magnetic Dynabeads M-280 (Life Technologies) according to the manufacturer’s recommendations. Negatively selected HIV-infected Jurkat cells were then limit-diluted to 0.5 cells per well in 96-well plates, and individual cell clones were expanded for 2–4 weeks in culture in the presence of 500 nM efavirenz. Clones were profiled for baseline reporter level and responsiveness to benchmark LRAs, with C16, I15 and N6 representing the most quiescent clones profiled for baseline reporter level and responsiveness to benchmark LRAs, with C16, I15 and N6 representing the most quiescent

Cell culture and Jurkat HIV–luciferase assay
Jurkat HIV–luciferase clones were maintained in Roswell Park Memorial Institute (RPMI) 1640 medium (Gibco, Life Technologies) containing 10% (v/v) fetal bovine serum (FBS; SAFC, Sigma-Aldrich), 25 μM 1′-penicillin and 25 μg/ml streptomycin (Gibco, Life Technologies), and were split 1:4 every 3–4 days to maintain a cell density of around 0.3–1 million cells per ml. The Jurkat clones were maintained with the addition of 500 nMefavirenz to the medium. Three Jurkat cell clones (C16, I15 and N6), each of which contained one or two integrated HIV proviruses that expressed the luciferase reporter gene, were added at equal amounts for a total of 5,000 cells per well to 384-well plates containing compound titrations. Dose–response testing was performed on compounds dissolved in dimethyl sulfoxide (DMSO; Fisher Scientific) dispensed in duplicate serial threefold, 14-point titrations using a D300e Digital Droplet Dispenser (Hewlett-Packard) to give final assay concentrations ranging from 10 μM to 2.1 pM in 50 μl of medium with a final concentration of 0.5% DMSO (v/v). Cells and compound were incubated at 37 °C for 48 h, unless otherwise indicated, followed by the addition of 20 μl of Steady-Glo Luciferase (Promega). Luminescence resulting from the induction of virally expressed luciferase was measured using an EnVision 2102 Multilabel Plate Reader (Perkin Elmer). Dose–response relationships were analysed with GraphPad Prism (v.6) using a four-parameter logistic regression model to calculate the concentration of compound that provides the half-maximal response and the maximal percentage activation compared to the vehicle control.

Immunoblot analyses
For the immunoblot assays, 10 μg of cell lysate was loaded per well into 4–20% Tris-Glycine SDS–PAGE gels. Proteins from the SDS–PAGE gels were transferred to Turbo Midi PVDF Transfer Packs (BioRad) using the ‘Mixed MW’ protocol for one Midi Format Gel (constant 2.5 A up to 25 V, for 7 min) of the Trans-Blot Turbo Transfer System (BioRad) with pre-made Trans-Blots according to the manufacturer’s instructions. After transfer, PVDF membranes were blocked in 5% bovine serum albumin (BSA) in 1× Tris-buffed saline (TBS) (BioRad) with 0.1% Tween-20 for 1 h at room temperature with gentle rocking. Primary antibodies were added and incubated overnight at 4 °C (anti-cIAP1, 1:1,000 (Abcam); anti-pI100/p52, 1:1,000 (Cell Signaling Technology); anti-kBα, 1:1,000 (Cell Signaling Technology); anti-cIAP2, 1:1,000 (Abcam); and anti-actin-HRP conjugate, 1:3,000 (Abcam). After staining with primary antibodies, the membrane was washed three times with 1× TBS and 0.1% Tween-20, 10 min each wash. After washing, the membrane was incubated in 5% BSA in 1× TBS and 0.1% Tween-20 with the appropriate secondary antibody for 2 h at room temperature. After staining with secondary antibodies, the membrane was washed twice for 10 min with 1× TBS and 0.1% Tween-20 followed by a 10-min wash with 1× TBS. The membrane was then patted dry with filter paper and an image was captured of the undeveloped membrane on a ChemiDoc MP Imaging System using Image Laboratory software (v.6.0.1, BioRad). Sufficient ECL reagent (GE Healthcare) was used to cover the membrane and a series of images was taken with increasing exposure times until the luminescence from the developed membrane saturated the image. The developed membrane was then washed 3 times with 1× TBS for 5 min to remove the residual ECL reagent and then stored at 4 °C in sufficient 1× TBS to submerge the entire membrane. Densitometry of images of the developed membrane was then carried out using Image Laboratory software (v.6.0.1, BioRad). Some membranes were stripped for 1 min with One Minute Plus Western Blot Stripping Buffer (GM Biosciences) and then washed 3 times for 10 min with 1× TBS. The stripped membranes were then blocked in 5% BSA in 1× TBS and 0.1% Tween-20 for 1 h and re-probed overnight with a new primary antibody. To normalize samples for loading a 1:20,000 dilution of β-actin (Abcam) was run on the stripped membranes.

Target gene RT–qPCR
We treated 2 million normal donor CD4+ T cells with a range of concentrations of AZD5882. Total RNA was isolated using the RNEasy Mini kit (Qiagen) according to the manufacturer’s instructions. The following

Software

The series of images was taken with increasing exposure times until the luminescence from the developed membrane saturated the image. The developed membrane was then washed 3 times with 1× TBS for 5 min to remove the residual ECL reagent and then stored at 4 °C in sufficient 1× TBS to submerge the entire membrane. Densitometry of images of the developed membrane was then carried out using Image Laboratory software (v.6.0.1, BioRad). Some membranes were stripped for 1 min with One Minute Plus Western Blot Stripping Buffer (GM Biosciences) and then washed 3 times for 10 min with 1× TBS. The stripped membranes were then blocked in 5% BSA in 1× TBS and 0.1% Tween-20 for 1 h and re-probed overnight with a new primary antibody. To normalize samples for loading a 1:20,000 dilution of β-actin (Abcam) was run on the stripped membranes.
outgrowth assays were performed as previously described\cite{32,33} with HPRT1 (RT–qPCR; Fast Virus 1-Step Master Mix, Applied Biosystems) was used and Hs02800695, Hs00174517_m1 (TaqMan primer probe sets were sourced from Applied Biosystems: >1 and adjusted \( P \) < 0.05. Graphs and summary tables were built in R using ggplot. Gene-set enrichment was performed using GSEA (v.2.2.3) and GO analysis (GO PANTHER v.11.1)\cite{39}. Results shown are the mean responses of the four donors tested.

**Generation and maintenance of BLT mice**

BLT mice were prepared as previously reported\cite{38,40,41}. In brief, a 1–2-mm piece of human liver tissue was sandwiched between two pieces of autologous thymus tissue (Advanced Bioscience Resources) under the kidney capsule of subcutaneously irradiated (200 cGy) 12–15-week-old female NOD.Cg-Pkdcd\textsuperscript{-/-}Il2rg\textsuperscript{-/-}\textit{Il6r}\textsuperscript{-/-}\textit{Il15r}\textsuperscript{-/-}SzJ (NSG; The Jackson Laboratory) mice. After implantation, mice were transplanted intravenously with CD34\textsuperscript{+} haematopoietic stem and progenitor cells isolated from autologous human liver tissue. Human immune cell reconstitution was monitored in the peripheral blood of BLT mice by flow cytometry every 3–4 weeks\cite{39}. For the study that examined the effect of AZD5582 administration on T cell activation, the mean weight of the mice was 26.88 g and they were approximately 1 year of age at the initiation of the study. For the study that examined the effect of AZD5582 administration on plasma and tissue viraemia during ART suppression, the mean weight of the mice used was 23.14 g and the mice were approximately 7 months old at the initiation of the study. Mice were randomized for assignment to either experimental or control groups (https://www.random.org/).

**HIV infection of BLT mice**

Stocks of HIV-1\textsubscript{in.CD4} were prepared as follows. The proviral clone was transfected into HEK293T cells using Lipofectamine 2000 (Invitrogen) following the manufacturer’s protocols. Viral supernatant was collected 48 h after transfection. Viral supernatant was titered by infecting TZM-bl cells (NIH AIDS Reagent Program, authenticated by morphological identification and virus-susceptibility profiles, tested for mycoplasma by the supplier) at multiple dilutions. Virus-containing medium was removed the next day and replaced with fresh Dulbecco’s modified Eagle medium (DMEM; Thermofisher) plus 10% FBS and the incubation continued for 24 h. The cells were fixed and stained with 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside and blue cells were counted directly to determine infectious particles per ml. Each titre of these viral stocks was performed in triplicate and at least two different titre determinations were performed for each virus stock. Exposure of BLT mice to HIV-1\textsubscript{in.CD4} was conducted by tail vein injection with 3 \times 10\textsuperscript{4} tissue culture infectious units of virus. Plasma viral load in peripheral blood of infected mice was monitored longitudinally by RT–qPCR using TaqMan RNA to-C\textsubscript{+}, 1-step kit (Applied Biosystems). The sequences of the forward and reverse primers and the TaqMan probe for PCR amplification and detection of HIV gag RNA were: 5′-CATGTTTTCA\textsuperscript{GCAATGTTTCTAAGAGAAG-3′}, 5′-TGCTTGATGTCCCCCCACT-3′ and 5′-FA M-CCACCCCCCAAGATTAAACACCAT-GCTAA-Q-3′, respectively. For viral load analysis, 40 μl of plasma was collected and analysed with a sensitivity of 350 copies per ml. All samples were run and analysed on an ABI 7500 Fast Real Time PCR System (Applied Biosystems).

**SIV infection of NHPs**

In total, 21 male and female Indian rhesus macaques, 3–6 years of age, with the exclusion of MamuB*08* and MamuB*17* macaques, were included in this study (Supplementary Table 7). Rhesus macaques were infected intravenously with 3 \times 10\textsuperscript{7} TCID\textsubscript{50} (50% tissue culture infective dose) of SIVmac259. The SIVmac259 Stock was titrated in vitro for viral infectivity by standard end-point titration on CEMx174 cells. The TCID\textsubscript{50} was calculated as previously described\cite{42}. Standard SIVmac259 plasma viral-load quantification was performed regularly throughout the study and the three times per week during the AZD5582 treatment period in the Translational Virology Core Laboratory of the Emory Center for AIDS Research using a standard qPCR assay (limit of detection of 60 copies per ml of plasma) as previously described\cite{43}. Ultra-sensitive SIVmac259 plasma viral-load quantification (limit of detection of 3 copies per ml of plasma) was performed for 2–3 time points before
AZD5582 treatment and 3–4 time points during AZD5582 treatment as previously described.\(^4,4^5\)

**ART and AZD5582 treatment of BLT mice and NHPs**

ART was administered to BLT mice using irradiated Teklad chow containing emtiracetabine (1,500 mg kg⁻¹), tenofovir disoproxil fumarate (1,560 mg kg⁻¹) and raltegravir (600 mg kg⁻¹) (Research Diets). Both the AZD5582 and the vehicle control (10% sterile captisol dissolved in sterile distilled water) were prepared fresh for each administration. AZD5582-2HCl was obtained from ChemieTek and dissolved at 5 mg ml⁻¹ (1,560 mg kg⁻¹) and raltegravir (600 mg kg⁻¹) (Research Diets). Both ART was administered to BLT T mice using irradiated Teklad chow containing emtricitabine (1,500 mg kg⁻¹), tenofovir disoproxil fumarate (1,560 mg kg⁻¹) and raltegravir (600 mg kg⁻¹) (Research Diets). Both ART and AZD5582 treatment of BLT mice and NHPs...parasitemia was initiated 56 days after infection that consisted of 2 reverse transcriptase inhibitors, tenofovir disoproxil fumarate (5.1 mg ml⁻¹) and emtricitabine (40 mg ml⁻¹) plus the integrase inhibitor dolutegravir (2.5 mg ml⁻¹). ART was administered once daily at 1 ml kg⁻¹ body weight via the subcutaneous route. Peak plasma viral load (measured by the standard assay) and plasma viral load before LRA intervention (as measured by the ultrasensitive assay) were controlled for when allocating macaques into experimental groups. In total, 12 rhesus macaques were treated with AZD5582 and 9 rhesus macaques served as controls. AZD5582 was infused weekly intravenously at 0.1 mg kg⁻¹. Three rhesus macaques received 3 doses of AZD5582 and were euthanized 48 h after the last dose. Nine rhesus macaques received 10 doses of AZD5582 and 3 macaques were euthanized 48 h after the last dose. Among the control rhesus macaques, 4 macaques received a weekly placebo infusion, 2 macaques were euthanized 48 h after 3 infusions and 2 macaques were euthanized 48 h after 10 infusions. The remaining five control macaques received ART only.

**Resting CD4⁺ T cell enrichment**

Human resting CD4⁺ T cells were enriched from total cells isolated from BLT mouse tissues as follows. Each tissue from each mouse was processed individually and then tissues were pooled for immunomagnetic sorting. Each pooled tissue was first enriched for human cells with the EasySep Mouse/Human Chimera Kit (Stem Cell Technologies) and then for resting CD4⁺ T cells with a human custom selection kit that included the following antibodies: CD8, CD4, CD14, CD19, CD20, CD36, CD56, CD123, glycophrin A, CD66b, CD25 and HLA-DR (all Stem Cell Technologies). Flow cytometry was performed before and after the enrichment to confirm the efficacy of the sort and purity of the sorted samples.

Resting CD4⁺ T cells from rhesus macaques were isolated from the peripheral blood, bone marrow, lymph node and spleen. Before sorting, CD4⁺ T cells were enriched using magnetic beads and column purification (Miltenyi Biotech). Enriched CD4⁺ T cells were then stained with previously determined volumes of the following fluorescently conjugated antibodies: CD3 AF700 (clone SP34-2), CD8 APC-Cy7 (clone SK1), CD69 PE-CFS94 (clone FN50), HLA-DR PerCP-Cy5.5 (clone G46-6) (all from BD Bioscience) and CD4 Brilliant Violet (BV) 650 (clone OKT4) and CD25 PE-Cy7 (clone BC96) (both from Biolegend). Resting CD4⁺ T cells were defined as CD3⁺CD4⁺CD8⁻CD69⁻HLA-DR⁻. Sorting was performed on a FACSAria LSR II (BD Biosciences) equipped with FACSDiva software.

**Plasma cytokine and chemokine analysis**

Human plasma cytokine and chemokine analysis in BLT mice was performed by the University of North Carolina, Chapel Hill Center for AIDS Research Virology Core Laboratory. Plasma was tested undiluted in single wells using a Millplex MAP kit (Millipore, HCYTMAG-60K-PX41) on a Luminex MAGPIX instrument. The following markers were tested: EGF, eotaxin, FGF-2, FLT-3L, fractalkine, G-CSF, GM-CSF, GRO, IFNα2, IFNγ, IL-1α, IL-1β, IL-1RA, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-12p40, IL-12p70, IL-13, IL-17A, IP-10, MCP-1, MCP-3, MDC, MIP-1α, MIP-1β, PDGF-AA, PDGF-AB/BB, RANTES, sCD40L, TGFB, TNF, TNFβ and VEGF. Rhesus plasma levels of proinflammatory cytokines and chemokines were evaluated using the NHP MSD V-Plex assay systems developed by Mesoscale Discovery. The two validated kits used were the V-PLEX Plus Proinflammatory Panel 1 NHP Kit (K15056D) that evaluates IFNγ, IL-10, IL-1β, IL-2, IL-6 and IL-8, and a custom Chemokine NHP Kit that evaluates IP-10, MCP-1 and MIP-1β (K15055G). The manufacturer-provided protocol was followed with a few modifications. As the plasma samples were infectious, Triton X-100 at a final concentration of 1% was added not only to the subject samples but also to the assay diluents provided with the MSD kit before use. Calibration standards were reconstituted according to the protocol provided with the assay diluent that had the Triton X-100 added to it so that all standards and samples had the same components present. Plasma (60 μl) was diluted twofold with assay diluent and then 50 μl of the diluted sample was added to the assay plate along with 50 μl of the calibrator according to the manufacturer’s protocol. The plate was then covered and incubated on a shaker for 2 h. After the 2-h incubation, plates were washed at least 6 times with the supplied wash buffer and then 25 μl of detection antibodies was added according to the kit protocol. Subsequently, the plate was covered and incubated on a shaker for 2 h at room temperature. After the detection incubation was finished, plates were then washed 6 or 150 μl of 2× read buffer was added to each well and plates were analysed on the Sector s600 MSD plate reader. Data analysis was performed using the MSD Discovery Workbench analysis software.

**Cell-associated HIV RNA quantification in BLT mice**

For tissue RNA analysis, RNA was extracted using QiAamp viral RNA columns (Qiagen) according to the manufacturer’s protocol including an optional treatment with RNase-free DNase and analysed using one-step reverse-transcriptase qPCR (ABI custom TaqMan Assays-by-Design). Known quantities of HIV gag RNA standards were run in parallel, creating a standard curve for HIV gag and sample RNA was quantified by extrapolation from the standard curve. All samples were run and analysed on an ABI 7500 Fast Real-Time PCR System (Applied Biosystems). Owing to the relatively low number of human cells found in the brain, HIV RNA levels were quantified for only three vehicle control- and three AZD5582-treated mice.

**Immunohistochemical analysis in BLT mice**

Tissues for immunohistochemical analysis were collected from BLT mice and fixed in 10% formalin for 16–24 h at 4 °C. Samples were then embedded in paraffin, cut into 5-μm sections and mounted onto poly-t-lysin-coated glass slides. After paraffin removal, antigen retrieval (DIVA Decloaker, Biocare Medical) and blocking of nonspecific immunoglobulin-binding sites (Background Sniper, Biocare Medical), tissue sections were stained with polyclonal goat IgG (R&D Systems) negative control antibodies. Tissue sections were imaged with a Nikon Eclipse C5i microscope using Nikon Elements BR software (v.4.30.01) and a Nikon Digital Sight DS-Fi2 camera.

**Serum chemistry analysis in mice**

After treatment with AZD5582 for 24 h, serum chemistry analysis in female 20-week-old BALB/cj mice (The Jackson Laboratory) was performed by the University of North Carolina, Chapel Hill Animal Histopathology Core Laboratory of the Lineberger Comprehensive Cancer Center. All clinical chemistry was performed on an Alfa Wassermann Vet Axcel analyser using Alfa Wassermann reagents. Automated assays for each analyte were as follows and performed according to the...
manufacturer’s instructions: alkaline phosphatase, Tietz-optimized Bowers and McCombs assay; aspartate aminotransferase, Henry modification of Karn’s assay; alanine aminotransferase, Henry modification of Wroblewski and LaDue assay; creatinine, Jaffe reaction; blood urea nitrogen, enzymatic assay; albumin, Doumas and Briggs modification of the bromresol green dye method; amylase, based on the use of chromogenic 2-chloro-p-nitrophenol linked with maltotriose; calcium, calcium-arsenazo assay phosphorus, based on the method of Daly and Ertingshausen with modifications by Armador and Urban; total bilirubin, Walters and Gerarde modification of the DMSO method; total protein, modification of Weichselbaum’s biuret reagent.

**Immunophenotyping of BLT mice by flow cytometry**

Immunophenotyping was performed on peripheral-blood samples collected longitudinally and at study end point, on blood and mononuclear cells isolated from the tissues of BLT mice. All flow cytometry data were collected on either BD LSR Fortessa or BD FACSCanto instruments using BD FACS Diva software (v.6.1.3) and data were analysed with FlowJo software (v.10.4.2). Antibodies used for longitudinal monitoring of human cells in peripheral blood include anti-CD45 APC (clone HI30, BD Biosciences), anti-CD3 FITC (clone HLT3a, BD Biosciences), anti-CD4 PE (clone RPA-T4, BD Biosciences) and anti-CD8 PerCP (clone SK1, BD Biosciences). Flow cytometry gating for the expression of lineage-specific antigens on human leukocytes was performed as follows. Step 1, forward- and side-scatter properties were used to set a live-cell gate. Step 2, live cells were then analysed for expression of the human pan-leukocyte marker CD45. Step 3, human leukocytes were then analysed for human CD3+ T cells. Step 4, T cells were analysed for human CD4 and CD8 expression (Supplementary Fig. 2a).

At collection, peripheral blood and cells isolated from each individual tissue were stained with antibodies to detect human CD45, CD3, CD4, CD8, CD38 (anti-CD38 APC, clone HB7, BD Biosciences) and HLA-DR (anti-HLA-DR PE, clone TU36, BD Biosciences) to assess T cell activation. The following gating strategy was used. Step 1, forward- and side-scatter properties were used to set a live cell gate. Step 2, live cells were then analysed for expression of the human pan-leukocyte marker CD45. Step 3, human leukocytes were subsequently analysed for human CD3+ T cells. Step 4, T cells were analysed for human CD4 and CD8 expression. Step 5, either CD4+ or CD8+ T cells were analysed for the expression of CD38 and HLA-DR (Supplementary Fig. 2b).

To determine the success of the sort and the purity of the sorted human resting CD4+ T cells, the following antibodies were used to analyse pre-sort and post-sort samples: anti-CD3 BV421 (clone UCHT1, BD Biosciences), anti-HLA-DR PerCP (clone L243, BD Biosciences), anti-CD4 BV605 (clone RPA-T4, BD Biosciences), anti-CD8 APC-Cy7 (clone SK1, BD Biosciences), anti-CD25 APC (clone 2A3, BD Biosciences) and anti-CD45 V500 (clone HI30, BD Biosciences). Antibodies used as isotype controls: anti-mouse IgG1x APC (clone MOPC-21, BD Biosciences), anti-mouse IgG2ax PerCP (clone X39, BD Biosciences), anti-mouse IgG1x PE (clone MOPC-21, BD Biosciences) and anti-mouse IgG2ax FITC (clone G155-178, BD Biosciences). Flow cytometry gating was performed as follows. Step 1, live cells were gated on forward scatter and side scatter. Step 2, human haematopoietic cells were gated based on expression of human CD45. Step 3, human T cells were gated based on expression of CD3. Step 4, T cell subsets were gated based on the expression of human CD8 and CD4. Step 5, expression of activation markers (CD25 and HLA-DR) were analysed on the surface of CD4+ T cells (Supplementary Fig. 2c).

**NHP sample collection and processing**

EDTA-anticoagulated blood samples were collected regularly and used for a complete blood count, routine chemical analysis and immunostaining. Plasma was separated by centrifugation within 1 h of phlebotomy. At the end of the studies, tissue samples were collected, including lymph nodes (21 rhesus macaques), spleen (10 rhesus macaques) and bone marrow (11 rhesus macaques). After 2 washes in RPMI and removal of connective and fat tissues, lymph nodes were ground using a 70-μm cell strainer. PBMCs and bone-marrow mononuclear cells were prepared by density-gradient centrifugation. CD4+ T cells were negatively selected from fresh or frozen cell suspensions using magnetically labelled microbeads and subsequent column purification according to the manufacturer’s protocol (Miltenyi Biotec).

**AZD5582 pharmacokinetics in NHPs**

Three healthy male rhesus macaques of Indian origin (aged 6–7 years, 10.5–13.0 kg) were used for the study. Fasted rhesus macaque received 0.1 mg kg−1 of AZD5582 in 10% capisol and ≤5% DMSO (dose volume 0.25 ml kg−1 at 0.40 mg ml−1), filtered during administration (0.22-μm PES in-line filter, Millex), via a 30-min saphenous vein infusion (infusion pump, Harvard Apparatus). Blood from femoral venipuncture or saphenous catheters was collected to obtain samples for plasma pharmacokinetics (K2 EDTA microtainers spun at 13,000 rpm for 5 min to obtain plasma), flow cytometry (Cytochex tubes (Streck)) and PBMC isolation (CPT blood collection tubes, BD Biosciences) processed according to the manufacturers’ instructions and a dry pellet of cells was snap-frozen on dry ice and stored at −80 °C at various times after administration. AZD5582 was extracted from macaque plasma samples with an isotopically labelled internal standard (rilpivirine-d6) using protein precipitation. The compound was then eluted from a Waters Atlantis T3 (50 mm × 2.1 mm, 3 μm particle size) analytical column under reverse-phase conditions and detected on an AB Sciex API-5000 triple quadrupole mass spectrometer under Turbolonspray mode. Standards were prepared in singlet and quality controls in duplicate, and a calibration curve was generated using a weighted 1/(x^2) linear regression of the concentration (x) versus the analyte/intermediate standard peak area ratio (y). Concentrations of quality controls and study samples were calculated from this calibration curve using Sciex Analyst Software (v.1.6.2). The acceptance criterion of the assay was ±25% of the nominal concentration for standards and quality controls, and the quantifiable range was 2–2,000 ng ml−1.

**Ex vivo analysis of AZD5582 activity in NHP cells**

Splenocytes from rhesus macaques were initially processed as above and frozen in PBS and 10% DMSO. Cryopreserved splenocytes were thawed and exposed to a range of concentrations of AZD5582 for either 48 h continuously or for 1 h after which the drug was removed by washing and cells were further cultured for 47 h. In preparation for western blot analysis, cells were lysed on ice with intermittent vortex mixing using NP40 Cell Lysis Buffer (Invitrogen, FNN0021) supplemented with appropriately diluted 10× Protease Inhibitor Cocktail (Sigma, P-2714), 1 mM PMSF (Sigma, 93482) and 1 mM DTT (Sigma, 43816) to create a final concentration of 1 mM each in the complete lysis buffer. The lysed cells were then centrifuged at 13,000 rpm for 10 min at 4 °C and then supernatants (soluble fraction lysates) were removed and stored at −80 °C or used immediately for protein concentration and western blot analysis.

**Flow cytometry assay for p100 protein**

Whole blood collected in Cyto-Chex blood collection tubes (Myriad RBM) was added to ACK lysis buffer to remove red-blood cells, washed with PBS and resuspended in a staining cocktail that included anti-CD3 BV421 (clone SP34-2, BD Biosciences), anti-CD16 BV605 (clone 5G8, BD Biosciences), anti-CD4 BV711 (clone L200, BD Biosciences), anti-CD14 BV786 (clone M5E2, BD Biosciences), anti-CD123 PerCP-Cy5.5 (clone 7G3, BD Biosciences), anti-CD20 PE-CF594 (clone MOPC-21, BD Biosciences) and anti-CD8 APC-Cy7 (clone SK1, BD Biosciences). After surface-staining, cells were washed twice with PBS, permeabilized using the Cytofix/Cytoperm kit (BD Biosciences, kit used as directed) and stained intracellularly with anti-p100 antibody.
infected cells were quantified using a limiting dilution culture assay in the repository Core Laboratory of the Emory Center for AIDS Research. Latently Replication-competent SIV reservoirs were measured by the Viral copies, respectively, for display purposes. RNA were assigned a level of 1 copy per million cells or million CD4 RNA wells were used in all analyses. Samples with undetectable SIV DNA or CTCCAGCACAGTCTATAAGAAAGAGG-3′. The means of two replicate were Rh-CD4-F 5′-ACATCGTGGTGCTAGCTTTCCAGA-3′, Rh-CD4-R Mastermix II (ThermoScientific). The CD4 primer and probe sequences quantified by qPCR of the resultant cDNA using Taqman Universal gag the High Capacity cDNA Reverse Transcription kit (ThermoScientific) associated RNA quantification, RNA was reverse-transcribed using nuclease (TaqMan) assay with SIV gag was performed by serially diluting this cDNA in independent PCR reactions to identify a dilution in which amplification occurred in <30% of the total number of reactions. PCR amplification was performed with 1× PCR buffer, 2 mM MgSO4, 0.2 mM of each deoxynucleoside triphosphate, 0.2 μM of each primer and 0.025 μl Platinum Taq High Fidelity polymerase (Invitrogen) in a 20-μl reaction. First-round PCR was performed with primer SIVenvF1 5′-CTCCCTCCAGGACTGAC-3′ and anti-sense primer SIVenvR1 under the following conditions: 1 cycle of 94 °C for 2 min, 35 cycles at 94 °C for 15 s, 55 °C for 30 s and 68 °C for 5 min, followed by a final extension of 68 °C for 10 min. Next, 1 μl from the first-round PCR product was added to a second-round PCR reaction that included the sense primer SIVenvF2 5′-TATAATAGACTGAGACACCCTTGAGGGGACG-3′ and anti-sense primer SIVenvR2 5′-ATAGGACATRTCTATTCCGCAAATGTGTA-3′ performed under the same conditions used for first-round PCR, but with a total of 45 cycles. Correctly sized amplicons were identified by agarose gel electrophoresis and directly sequenced with second-round PCR primers and nine SIV-specific primers using BigDye Terminator technology (Applied Biosystems). To confirm PCR amplification from a single template, chromatograms were manually examined for multiple peaks, indicative of the presence of amplicons that resulted from PCR-generated recombination events, Taq polymerase errors or multiple variant templates. Alignment and phylogenetic trees were implemented in Geneious Prime 2019 (Biomatters) using the Muscle algorithm and the neighbour-joining method with the Tamura–Nei genetic distance model, respectively.

Cell-associated SIV RNA and DNA quantification in NHPs
Cell-associated SIV RNA and DNA were measured simultaneously in resting CD4+ T cells isolated from peripheral blood, spleen, bone marrow and lymph nodes (52,279–500,000 cells) or in total CD4+ T cells isolated from peripheral blood or lymph nodes (100,000–500,000 cells), lysed in RTL Buffer (Qiagen), and stored at −80 °C. Nucleic acids were extracted using the AllPrep DNA/RNA mini kit (Qiagen) according to the manufacturer’s recommendations with an on-column DNase digestion step. Cell-associated DNA quantification of SIV\textsubscript{mac239} gag DNA was performed on the extracted cell-associated DNA by qPCR using a 5′ nuclease (TaqMan) assay with SIV gag primers and normalized to the rhesus macaque albumin gene, as previously described. For cell-associated RNA quantification, RNA was reverse-transcribed using the High Capacity cDNA Reverse Transcription kit (ThermoScientific) and random hexamers. SIV gag and the rhesus macaque CD4 gene were quantified by qPCR of the resultant cDNA using Taqman Universal Mastermix II (ThermoScientific). The CD4 primer and probe sequences were Rh-CD4-F 5′-ACATCGTGGTGCTAGCTTTCCAGA-3′, Rh-CD4-R 5′-AAGTGTTAAGGCGATTGGAGAGA-3′ and Rh-CD4-probe 5′-AGGCCCTCAGCAGCTTATAAGAAAAGGG-3′. The means of two replicate wells were used in all analyses. Samples with undetectable SIV DNA or RNA were assigned a level of 1 copy per million cells or million CD4 RNA copies, respectively, for display purposes.

SIV quantitative viral outgrowth assay
Replication-competent SIV reservoirs were measured by the Viral Reservoir Core Laboratory of the Emory Center for AIDS Research. Latently infected cells were quantified using a limiting dilution culture assay in which CD4+ T cells—enriched from lymph node or spleen cells using magnetic beads and column purification (Miltenyi Biotec)—were co-cultured with CEMx174 cells in fivefold serial dilutions ranging from 5 × 10^4 cells per well to 4 × 10^6 cells per well. The cells were cultured in RPMI containing 10% FBS and 100 U ml−1 IL-2 (Sigma). The ratio of target cells added was 4:1 for the two highest dilutions. A constant number of 1 × 10^5 CEMx174 cells was added to all other wells. The cultures were split every 7 days, and fresh medium was added. After 21 days, the growth of virus was detected by RT–qPCR. SIV RNA was isolated from 400 μl of culture supernatant using the Zymo viral RNA isolation kit (Zymo Research). DNase treatment was performed using a RQ1 RNase-free DNase kit (Promega). A one-step RT–qPCR targeting SIV gag was performed using an Applied Biosystems 7500 Real Time PCR System (Applied Biosystems) and the Taqman Fast Virus 1-step Master Mix (Thermo Scientific) for qRT–PCR with the following primers and probe: SIV gag Fwd 5′-GCAGAGAGGAAATCCACGCATTAC-3′, SIV gag Rev 5′-CAATTCCCGACCATTTATGTF-3′ and SIV gag probe 5′-6FAM-GTAGCCACCTGCCATTACGGCCA-3IBFQ-3′. The frequencies of infected cells were determined by the maximum-likelihood method and were expressed as infectious units per million CD4+ T cells.

RNA-seq analysis of NHP cells
RNA-seq analysis was conducted at the Yerkes Non-human Primate Genomics Core Laboratory (http://www.yerkes.emory.edu/nhp_genomics_core/). RNA was purified from 50,000 peripheral-blood- or lymph-node-derived CD4+ T cells, which were purified by flow cytometry and lysed in 350 μl of RTL buffer at −80 °C, using Qiagen Micro RNAeasy columns, and RNA quality was assessed using an Agilent Bioanalyzer. Then, 2 ng of total RNA was used as input for mRNA amplification using 5′ template switch PCR with the Clontech SMART-seq v4 Ultra Low Input RNA kit according to the manufacturer’s instructions. Amplified mRNA was fragmented and appended with dual-indexed bar codes using Illumina NexteraXT DNA library preparation kits. Libraries were validated by capillary electrophoresis on an Agilent 4200 TapeStation, pooled and sequenced on an Illumina HiSeq 3000 using 100-bp single reads at an average depth of 25 million reads.

RNA-seq statistical analyses
RNA-seq data were mapped to the MacaM (v.7.8) assembly of the Indian rhesus macaque genome (available at https://www.unmc.edu/rhesusgenephip/index.htm) and alignment was performed using STAR (v.2.5.2) using the annotation as a splice junction reference. Transcripts were annotated using the MacaM (v.7.8.2) annotation. Transcript abundance was estimated within STAR using the htseq-count algorithm and differential-expression analyses were performed using DESeq2. For significance testing using DESeq2, all of the samples from three- and ten-dose rhesus macaques were grouped together, separately for peripheral blood and lymph nodes, and compared with their pre-treatment samples. Genes determined to be differentially expressed by DESeq2 (adjusted p < 0.05 and fold change > ±1.5) were tested for enrichment of molecular pathways using the DAVID database (https://david.ncifcrf.gov/) and by using GSEA with the desktop module (https://www.broadinstitute.org/gsea/). Gene sets for GSEA analysis were selected from the MSigDB database (http://software.broadinstitute.org/gsea/msigdb/index.jsp) and from the previously described ‘blood transcriptional modules. Heat maps, DAVID enrichment bar charts and GSEA enrichment plots were generated with the R (v.3.5.0) package ggplot2. Principal component analysis was performed using Partek Genomics Suite software (v.6.6.6) using a covariance matrix.

Immunophenotyping of NHPs by flow cytometry
Multicolour flow cytometry analysis was performed on whole blood and lymph-node mononuclear cells using predetermined optimal concentrations of the following fluorescently conjugated monoclonal antibodies: CD3 APC-Cy7 (clone SP34-2), Ki-67 AF700 (clone B56), HLA-DR (clone EPR18756; Abcam). Cells were washed twice and stained with a secondary chicken anti-rabbit Alexa Fluor 488 antibody (Invitrogen). Samples were acquired using a four-laser Fortessa flow cytometer (BD Biosciences) and analysed with FlowJo software (version 9.7.6, TreeStar).
PerCP-Cy5.5 (clone G46-6), CCR5 APC (clone 3A9), CCR7 FITC (clone A20), CD45RA PE-Cy7 (clone SH9) and CD62L PE (clone SK11) (all from BD Biosciences); CD8 BV711 (clone RPA-T8), CD4 BV650 (clone OKT4), CD95 BV605 (clone DX2) and PD-1 BV421 (clone EH12.2H7) (all from Biologend) and CD28 PE-Cy5.5 (clone CD28.2) (from Beckman Coulter). Flow cytometry acquisition and analysis of samples was performed on at least 100,000 events on an LSRII flow cytometer driven by the FACSDiva software package (BD Biosciences). Analyses of the acquired data were performed using FlowJo software (Tree Star, v.10.0.4).

**ELISPOT for SIV-specific IFNγ production in NHPs**

IFNγ production was evaluated after the stimulation of PBMCs with SIVmac239 Gag and SIVmac239 Env peptide pools. The following reagents were obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH: SIVmac239 Env Peptide Pool and the SIVmac239 Gag Peptide Pool. To analyse IFNγ expression, we used the Monkey Elispot kit from MABTECH. The manufacturer’s instructions were followed as provided with the exception that concanavalin A (final concentration of 2.5 μg ml⁻¹) was used as the positive control agent. ELISPOT plates were blocked for 30 min with 100 μl of RPMI 1640 supplemented with 10% FBS. PBMCs (4 × 10⁶ per well) were incubated with 1 μg ml⁻¹ of DMSO, SIVmac239 Gag Peptide Pool or SIVmac239 Env Peptide Pool for 18 h before running the assay. For each plate, concanavalin A was used for the positive control wells. Samples were run in duplicate.

**Intracellular cytokine staining**

PBMCs from five ART-suppressed SIV-infected rhesus macaques were thawed. Pre-treatment with AZD5582 was performed for 1 h at 100 nm. After two washes, phorbol 12-myristate 13-acetate (PMA) and ionomycin were added for 1 h at 500 ng ml⁻¹ and 10 μg ml⁻¹, respectively. DMSO treatment controls were prepared in parallel. After 1 h, brefeldin-A and Golgi stop solution were added following the manufacturer’s recommendations (BD GolgiStop Protein Transport Inhibitor, BD Biosciences). Cells were incubated at 37°C, 5% CO₂ in R10 for 6–8 h before staining with the following antibodies: CD3 APC-Cy7 (clone SP3-42), CD8 BV711 (clone RPA-T8), CD4 BV650 (clone OKT4) (all from Biologend), and IFNγ PE (clone B7), TNF AF700 (clone Mab21) and IL-2 BV605 (clone MQ1-17H12) (all from BD Biosciences). Flow cytometry acquisition and analysis of samples was performed on at least 100,000 events using an LSRII flow cytometer driven by the FACSDiva software package (BD Biosciences). Analyses of the acquired data were performed using FlowJo (Tree Star, v.10.0.4) and simplified presentation of incredibly complex evaluations (SPICE, v.6.0) software.

**Proliferation assay**

PBMCs from five ART-suppressed SIV-infected rhesus macaques were thawed and labelled with CellTrace Violet Proliferation Kit according to the manufacturer instructions (Molecular Probes). Cells were plated in R10 in 96-well round-bottom plates at 4 million cells per ml. Pre-treatment with AZD5582 was performed for 1 h at 100 nM. After two washes, PMA and ionomycin were added for 1 h at 500 ng ml⁻¹ and 10 μg ml⁻¹, respectively. DMSO treatment controls were prepared in parallel. Cells were incubated at 37°C, 5% CO₂ in R10 supplemented with 20 IU ml⁻¹ IL-2. After 5 days, cells were stained with CD3 APC-Cy7 (clone SP3-42), CD8 BV711 (clone RPA-T8), CD4 BV650 (clone OKT4) (all from Biologend) and analysed by flow cytometry on an LSRII instrument. Data were analysed using FlowJo software (Tree Star, v.10.0.4).

**Statistical analysis**

Statistical analyses were performed using GraphPad Prism Software (v.6 or v.7). P < 0.05 was considered statistically significant. At least three samples were used for each group, the minimum to achieve statistical significance. No statistical methods were used to predetermine sample size. Investigators were not blinded to group allocations or when assessing outcomes. In some instances, cells were pooled from individual humanized mice for each tissue and experimental group for the isolation of resting CD4⁺ T cells (Fig. 2d and Extended Data Fig. 2). To test the statistical significance of the differences that we observed in PCR data in BLT mouse brains in Fig. 2e and grouped tissue PCR data in Fig. 2d, unpaired two-sided Student’s t-tests were used. To test the statistical significance of the differences that we observed in PCR data in PBMCs and female reproductive tract in Fig. 2e, T cell activation markers in Supplementary Table 5 and plasma cytokines and chemokines in Supplementary Table 6, we used unpaired two-sided Mann–Whitney U-tests. To assess the statistical significance of the differences observed over time in serum enzymes in Supplementary Table 4, we used paired nonparametric Wilcoxon matched-pairs signed-rank tests. To test the statistical significance of the differences in activation marker levels expressed on T cells from rhesus macaques in Fig. 4d, e, as well as the ultrasensitive plasma viral load results in Extended Data Fig. 8b and the ex vivo proliferation assay in Extended Data Fig. 10g, we used Wilcoxon matched-pairs signed-rank tests. To assess the statistical significance of the differences observed between SIV DNA and SIV RNA levels in resting or total CD4⁺ T cells from rhesus macaques in Fig. 3e and Extended Data Fig. 6b, respectively, as well as for the quantitative viral outgrowth results in Extended Data Fig. 6c, we used unpaired two-sided Mann–Whitney U-tests. Unpaired two-sided Mann–Whitney U-tests were also used to compare rhesus macaques with stable compared with increased viraemia in Extended Data Fig. 8c.

**Reporting summary**

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

**Data availability**

Source Data for Figs. 1–4, Extended Data Figs. 1, 2, 4–6 and Supplementary Tables 4–6 are provided with the paper. Gene-expression data are available at the Gene Expression Omnibus (GEO) repository (accession number GSE141546 and GSE142774). Any other data are available from corresponding authors on reasonable request.

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**Author contributions**

J.V.G., A.C., G.S., R.M.D., A.W. and D.M.M. conceived and designed the studies. G.C.S., D.M.I., S.R.W., S.D.F., C.G., E.P.B. and R.G.F. assessed in vitro AZD5582 activity. M.K., Z.W., J.H.B., D.F. and C.D.J. performed the in vitro transcriptomic study. D.F. and J.-P.R. contributed clinical specimens. C.C.N., P.T.H., C.D., N.J.S. and B.L. performed mouse experiments. R.A.S. and C.G.C. performed the qPCR analysis of HIV RNA levels in mouse samples. R.A.C. performed the immunohistochemical analysis. N.M.A. isolated human resting CD4+ T cells. C.C.N. analysed mouse data. M.M., A.D.B. and S.J. performed the monkey experiments. G.C.S., J.D., S.M. and R.M.D. conducted pharmacokinetic studies in NHP. C.M. and N.S. isolated resting CD4+ T cells for macaques. T.H.V. supervised the qPCR analyses of SIV RNA and DNA levels in monkey samples. M.M. and A.D.B. performed flow cytometry and analysed monkey data. G.K.T., A.A.U. and H.W. performed the macaque RNA-seq analyses and S.E.B. analysed these data. C.M.F. and B.F.K. performed env sequencing analyses. J.D.L. supervised the ultrasensitive plasma viral load. G.C.S. and C.G. analysed AZD5582 target engagement in human, mouse and macaque samples and J.D. and R.M.D. analysed the data. C.G. performed the ELISPOTs. C.C.N., M.M., A.W., A.C. and J.V.G. wrote the manuscript with input from all authors.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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**Correspondence and requests for materials** should be addressed to R.M.D., A.C. or J.V.G.

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Extended Data Fig. 1 | Short-duration exposure to AZD5582 activates the ncNF-κB pathway. **a**–**d**, Isolated total human CD4⁺ T cells treated with 100 nM AZD5582 and then either not washed (**a**) or washed three times with PBS 8 h (**b**), 4 h (**c**) or 1 h (**d**) after treatment. Whole-cell lysates were then analysed by immunoblot for components of the ncNF-κB pathway. **e**, Densitometry analysis of the ratio of p52 to p100 from the pulse-wash assay immunoblots. Points represent values for the densitometric ratio from one western blot, representative of several independent experiments. **a**–**e**, Entire set representative of 1 experiment with 5 replicates of the 1-h wash condition conducted. **f**, Fold induction of ncNF-κB target gene expression in isolated CD4⁺ T cells from an uninfected donor treated with the indicated concentrations of AZD5582 and either washed after 1 h or not washed, and subsequently cultured for 24 h as measured by RT-qPCR. Points represent two technical replicates and lines represent the mean. The data presented are representative of three independent experiments. **g**, DMSO-normalized induction of luciferase activity from the Jurkat reporter model after exposure to AZD5582 (10, 100 or 1,000 nM) for 30 min (blue), 1 h (red), 2 h (green) or continued exposure (purple). Points represent three replicates in one assay run, representative of two independent experiments. Lines represent the mean of the three replicates. For gel source data, see Supplementary Fig. 1.
Extended Data Fig. 2 | AZD5582 induces HIV RNA expression in resting CD4\(^+\) T cells from tissues of HIV-infected, ART-suppressed BLT mice. HIV RNA levels in resting CD4\(^+\) T cells isolated from the bone marrow, lymph nodes, liver, lung and spleen of control (Cntrl, blue circles) or AZD5582-treated (AZD, red circles) mice (cells pooled from \(n = 6\) mice per group for each tissue) were analysed in triplicate. Statistical significance was determined with a two-sided Student’s \(t\)-test. The mean fold increase in viral RNA levels in resting CD4\(^+\) T cells from tissues in the experiment shown in Fig. 2 was 12.1 (±3.7), whereas in this figure it was 8.4 (±6.0). These values were not statistically different (\(P = 0.4286\), two-sided Mann–Whitney \(U\)-test). Data are mean ± s.e.m.
Extended Data Fig. 3 | AZD5582 ex vivo target engagement. a, Western blot analysis of p100, p52 and cIAP1 protein levels in cells isolated from the bone marrow, lymph node and spleen of BLT mice before and 20 h after ex vivo treatment with AZD5582. Loading control, β-actin. Representative of two experiments. b, cIAP expression in resting CD4+ T cells isolated from the thymus, spleen, lymph nodes, liver, lung and bone marrow. Loading control, β-actin. Representative of two experiments. c, cIAP expression in the thymic organoid of HIV-infected ART-suppressed BLT mice 48 h after the administration of vehicle control or AZD5582 (three control and one AZD5582-treated analysed). Positive cells, brown. Imaged at 4× and 40× magnifications; scale bars, 100 μm (4×) and 50 μm (40×). Boxes in the 4× images indicate regions corresponding to images at 40× magnification. For gel source data, see Supplementary Fig. 1.
Extended Data Fig. 4 | Pharmacokinetic and pharmacodynamic assessment of AZD5582 in rhesus macaques. a, AZD5582 (0.1 mg kg\(^{-1}\)) was administered to healthy rhesus macaques (\(n = 3\)) by intravenous infusion. Plasma concentrations of AZD5582 (left y axis) are shown for the indicated time points. Flow cytometry was used to measure intracellular p100 levels, shown as the geometric mean fluorescence intensity (gMFI) in CD4\(^{+}\) T cells and plotted as the percentage of baseline of p100 gMFI (right y axis).

b, Plasma concentrations of AZD5582 after one (dark red), three (red), six (pink) or ten (orange) doses in six SIV-infected ART-treated rhesus macaques (RM) and after one dose in three uninfected control rhesus macaques (grey). Individual values are shown as symbols.

c, Western blot analyses of inactive p100 and active p52 forms of NF-κB in lymph-node mononuclear cells collected 48 h after the third or tenth dose of AZD5582 in SIV-infected ART-suppressed rhesus macaques (red; \(n = 3\) for both the 3-dose and 10-dose groups) or at equivalent time points for placebo controls (blue; \(n = 2\) for both the 3-dose and 10-dose groups). Immunoblots are shown in the top panels and densitometry analyses of the p52:p100 ratios are shown in the bottom panels. The line represents the median.

d, Cryopreserved control rhesus macaque splenocytes were treated with the indicated concentrations of AZD5582 for 48 h, then p100/p52 levels were analysed by western blotting to measure engagement of the ncNF-κB pathway.

e, DMSO-normalized densitometric p52:p100 ratio versus the AZD5582 concentration. For d, e, the experiments were performed in duplicate.

f, Cryopreserved rhesus macaque splenocytes were exposed to DMSO alone (Untx), 100 nM AZD5582 washed off after 1 h and cultured for 47 h (Pulsed) or continuous 100 nM AZD5582 for 48 h (Cont.), after which the cells were studied by western blot for p100 and p52 levels.

g, Densitometric p52:p100 ratio. For f, g, data represent a single experiment. For gel source data, see Supplementary Fig. 1.
Extended Data Fig. 5 | Phylogenetic trees based on full env sequencing. Plasma virus was sequenced from four time points per macaque (n = 5): near peak viraemia (2 weeks after infection; red), immediately before ART (8 weeks after infection; orange) and two time points of on-ART viraemia during AZD5582 treatment (green and blue). All sequences per macaque were phylogenetically analysed and the resulting phylogenetic trees are shown for each macaque. The horizontal bar indicates the genetic distance. nt, nucleotide.
Extended Data Fig. 6 | SIV DNA levels in total CD4+ T cells and replication-competent reservoir size in ART-suppressed SIV-infected rhesus macaques.

a. Longitudinal assessment of cell-associated SIV DNA levels in total CD4+ T cells isolated from peripheral blood and lymph nodes of AZD5582-treated (n = 12, red) and control (n = 4, blue) ART-suppressed SIV-infected rhesus macaques. Grey shading represents the period of ART administration.

b. Comparison of cell-associated SIV DNA levels in total CD4+ T cells isolated from the lymph nodes and peripheral blood of AZD5582-treated and control ART-suppressed SIV-infected rhesus macaques. Total CD4+ T cells were analysed from AZD5582-treated rhesus macaques 48 h after receiving 3 doses (lymph nodes and peripheral blood, n = 3) or 10 doses (lymph nodes and peripheral blood, n = 9) of AZD5582. Total CD4+ T cells were analysed from placebo-control rhesus macaques (lymph nodes and peripheral blood, n = 4) at equivalent time points. Open symbols indicate AZD5582-treated rhesus macaques with on-ART viraemia above 60 copies per ml of plasma. Statistical significance was determined using a two-sided Mann–Whitney U-test.

c. Quantitative viral outgrowth assays were performed for AZD5582-treated rhesus macaques 48 h after receiving 3 doses (lymph nodes, n = 2; spleen, n = 3) or 10 doses (lymph nodes, n = 3; spleen, n = 2) of AZD5582. Quantitative viral outgrowth assays were performed from control rhesus macaques (lymph nodes and spleen, n = 4) at equivalent time points. Open symbols indicate AZD5582-treated rhesus macaques with on-ART viraemia. Statistical significance was determined with a two-sided Mann–Whitney U-test. Horizontal lines represent the median (b, c).
Extended Data Fig. 7 | AZD5582-induced gene pathways and genes in SIV-infected ART-suppressed rhesus macaques. a, DAVID analysis showing pathways significantly enriched in genes differentially expressed after AZD5582 treatment relative to baseline in CD4+ T cells isolated from lymph nodes (black bars) and peripheral blood (grey bars). \( n = 6 \) for both lymph nodes and peripheral blood; for each, \( n = 3 \) for 3 doses of AZD5582 and \( n = 3 \) for 10 doses of AZD5582. b, Leading edge genes from the 'hallmark TNF signalling via NF-κB' pathway from MSigDB. Genes were identified in the leading edge of CD4+ T cell samples from the lymph nodes before and after treatment with AZD5582 (shown in Fig. 4b). The contrast depicted is the fold change of each gene for each rhesus macaque’s post-treatment sample relative to the pre-treatment values for CD4+ T cells from the lymph nodes (top) and the peripheral blood (bottom). \( n = 6 \) for both lymph nodes and peripheral blood; for each, \( n = 3 \) for 3 doses of AZD5582 and \( n = 3 \) for 10 doses of AZD5582. c, Heat map of cNF-κB (top) and ncNF-κB (bottom) pathway gene expression in rhesus macaques with (left, \( n = 5 \)) or without (right, \( n = 6 \)) on-ART viraemia of >60 copies per ml plasma. One rhesus macaque without on-ART viraemia was excluded from this analysis because of technical issues (higher than expected unmapped and multi-mapped reads, and lower than expected unique identified reads compared to the means). Colour scale, log2-transformed fold changes of post-treatment compared with pre-treatment values.
Extended Data Fig. 8 | Relationship between virological status before intervention and virological response to AZD5582 treatment in SIV-infected ART-suppressed rhesus macaques. 

**a**, Individual representations of plasma SIV RNA levels measured by ultrasensitive assay (limit of detection 3 copies per ml of plasma) before and during AZD5582 treatment in SIV-infected ART-suppressed rhesus macaques (n = 12). ‘Baseline’ shows two or three plasma viral loads before AZD5582 treatment and ‘AZD5582’ shows three or four plasma viral loads during AZD5582 treatment. Open symbols indicate the five AZD5582-treated rhesus macaques with on-ART viraemia of >60 copies of SIV RNA per ml of plasma using the standard viral load assay (confirmed with ultrasensitive assay). The orange box highlights an additional 3 rhesus macaques who did not experience on-ART viraemia of >60 copies per ml of plasma but had ≥2 SIV RNA measurements above baseline values using the ultrasensitive quantification method. 

**b**, Comparison of the medians of plasma SIV RNA levels at baseline and during AZD5582 treatment measured by ultrasensitive assay for ART-suppressed SIV-infected rhesus macaques (n = 12). Statistical significance was determined with a Wilcoxon matched-pairs signed-rank test. 

**c**, Comparison of the levels of plasma SIV RNA at peak, plasma SIV RNA before ART initiation, SIV DNA in peripheral-blood CD4+ T cells before AZD5582 treatment (pre-LRA), SIV DNA in lymph-node CD4+ T cells pre-LRA, SIV RNA in peripheral blood CD4+ T cells pre-LRA and SIV RNA in lymph-node CD4+ T cells pre-LRA in rhesus macaques with increased plasma viral loads (PVL) by standard and/or ultrasensitive assay during AZD5582 treatment (increased plasma viral loads, n = 8) compared with rhesus macaques that did not demonstrate increased viral loads (stable plasma viral loads, n = 4). Statistical significance was determined using a two-sided Mann–Whitney U test. Horizontal lines represent the median. (a, c).
Extended Data Fig. 9 | AZD5582 can be safely administered in SIV-infected ART-suppressed rhesus macaques. a–c, Longitudinal assessment of serum chemistries (a), complete blood counts (b) and weight (c) of SIV-infected ART-treated rhesus macaques treated with AZD5582 (red, $n = 12$) compared with controls (blue, $n = 9$). Grey shading represents the period of ART administration. AST, aspartate aminotransferase; ALT, alanine aminotransferase; BUN, blood urea nitrogen; GGT, γ-glutamyltransferase; HGB, haemoglobin; WBC, white blood cell.
Extended Data Fig. 10 | Immunological effect of AZD5582 on SIV-infected ART-suppressed rhesus macaques. a, b, Longitudinal assessment of the count (a) and the frequency and viability (b) of CD4+ T cells in SIV-infected ART-treated rhesus macaques during the period of AZD5582 treatment (n = 12). Lines represent median values. c, d, CD4+ T cell naïve and memory subset frequencies (c) and their expression of Ki-67 (d) in SIV-infected ART-treated rhesus macaques during the period of AZD5582 treatment (n = 12). Lines represent median values. e, f, CD8+ T cell naive and memory subset frequencies (e) and their expression of IFNγ (f) in SIV-infected ART-treated rhesus macaques during the period of AZD5582 treatment (n = 12). Lines represent median values. g, h, CD8+ T cell proliferative response to stimulation with PMA and ionomycin with or without AZD5582 pre-treatment. Top, representative flow cytometry dot plots gated on memory CD8+ T cells. Bottom, comparison between divided cells 5 days after stimulation in each group (n = 5 for each). Statistical significance was determined with a Wilcoxon matched-pairs signed-rank test. Horizontal lines represent the median. i, Longitudinal assessment of plasma levels of IP-10, MIP-1β, MCP-1, IL-6 and IL-10 by multiplex assay in SIV-infected ART-suppressed rhesus macaques treated with AZD5582 (red, n = 6) or control rhesus macaques (blue, n = 4). An additional four analytes (IFNγ, IL-8, IL-1β and IL-2) were undetectable in all macaques. a–h, Grey shading represents the period of ART administration. Dashed lines represent AZD5582 or placebo infusions. LLOD, lower limit of detection.