HDAC inhibition induces HIV-1 protein and enables immune-based clearance following latency reversal

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Introduction

HIV persistence in latently infected, resting CD4+ T cells is broadly recognized as a barrier to eradicating HIV (1, 2). To eliminate the latent reservoir, infected cells must be recognized as harboring virus and then effectively destroyed and cleared. Latency disruption is under intense investigation, and several strategies are being evaluated (3–6). Recent therapeutic approaches have focused on the use of latency-reversing agents (LRAs) to induce viral transcription, initiate viral translation, and subsequently elicit death through viral cytopathic effects or immune-mediated cell killing (“shock and kill”). A class of small molecules known as histone deacetylase inhibitors (HDACis) has been broadly examined as LRAs, and clinical studies have shown that single and multiple administration results in induced viral RNA transcription in CD4+ T cells from ART-suppressed HIV+ individuals, and we show viral protein induction following treatment with LRAs. Importantly, we demonstrate that clinical administration of histone deacetylase inhibitors (HDACis; vorinostat and panobinostat) induced HIV gag p24, and ex vivo stimulation produced sufficient viral antigen to elicit immune-mediated cell killing using anti-gp120/CD3 bispecific antibody. These findings extend beyond classical nucleic acid endpoints, which are confounded by the predominance of mutated, defective proviruses and, of paramount importance, enable assessment of cells making HIV protein that can now be targeted by immunological approaches.

Promising therapeutic approaches for eradicating HIV include transcriptional activation of provirus from latently infected cells using latency-reversing agents (LRAs) and immune-mediated clearance to purge reservoirs. Accurate detection of cells capable of producing viral antigens and virions, and the measurement of clearance of infected cells, is essential to assessing therapeutic efficacy. Here, we apply enhanced methodology extending the sensitivity limits for the rapid detection of subfemtomolar HIV gag p24 capsid protein in CD4+ T cells from ART-suppressed HIV+ individuals, and we show viral protein induction following treatment with LRAs. Importantly, we demonstrate that clinical administration of histone deacetylase inhibitors (HDACis; vorinostat and panobinostat) induced HIV gag p24, and ex vivo stimulation produced sufficient viral antigen to elicit immune-mediated cell killing using anti-gp120/CD3 bispecific antibody. These findings extend beyond classical nucleic acid endpoints, which are confounded by the predominance of mutated, defective proviruses and, of paramount importance, enable assessment of cells making HIV protein that can now be targeted by immunological approaches.
The rare incidence of latently infected cells for which robust phenotypic markers are still needed presents a significant challenge for identifying these cells and measuring changes in viral expression (3, 19, 20). In addition, latency reversal may induce proviral transcription in only a fraction of this population, and the observed increases in cellular HIV-1 RNA may represent defective proviruses and could also result from alternative splicing or read-through transcripts, which may not give rise to efficient splicing, translation, and protein expression (19–22). Quantifying virus production has been difficult, and traditional cell culture–based assays, such as viral outgrowth assays (VOA), which measure HIV gag p24 antigen in culture medium after expansion in culture for 2–3 weeks, require large sample volumes and have limited throughput and clinical application (23, 24). TILDA, as an example of many cell-associated HIV RNA assays described, may offer some advantages to quantitative VOA (qVOA) such as increased throughput and decreased sample requirements, but still only measures RNA transcripts (25). An alternative approach that directly measures protein production and also affords increased sensitivity as a component of an existing ex vivo assay for measuring HIV reservoir and/or can be leveraged to assess protein levels needed to induce viral cytopathicity or promote immune-mediated clearance is an important advancement for HIV cure research (26–28). Toward this goal, we report and apply enhanced methodology for rapid and sensitive detection of HIV gag p24 protein in latently infected CD4+ T cells from ART-suppressed HIV+ individuals and demonstrate that latency reversal of HIV can produce sufficient viral antigen to enable immune-targeted clearance.

Results

Direct measurement of HIV gag p24 in ART-suppressed HIV+ CD4+ T cells. Recent studies have reported the use of a digital immunoassay to measure HIV gag p24 protein from serum or plasma of acutely infected HIV+ individuals or cultured media of HIV-infected CD4+ T cells (29–31). Despite these advances, commercial methods encounter matrix issues and lack the same degree of sensitivity in quantifying cell-associated viral p24 protein from aviremic HIV+ individuals. To overcome this limitation, we optimized the digital immunoassay protocol, detailed in Methods, for the rapid and sensitive detection of HIV gag p24 protein in latently infected CD4+ T cell lysates prepared from ART-suppressed HIV+ individuals and applied this approach to assess proviral reactivation and immune-mediated cell kill. Robust linearity in detecting p24 in CD4+ T cell lysates from aviremic HIV+ individuals was demonstrated with an analytical limit of detection (LOD) of 5 fg/ml (Figure 1A), corresponding to 125,000 p24 copies/ml or 18,000 copies/144 μl assay input and, thus, about 9 viruses/144 μl input (29, 30). The optimized methods extended the sensitivity LODs for HIV p24 in cell lysates (Figure 1B). Additional studies in the HIV-1–infected MOLTIIIB cell line also suggested HIV gag protein quantification is feasible in 0.125 cells/ml lysate (~2% cell/144 μl) in this cell model (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.92901DS1).

To demonstrate assay specificity and breadth, we first evaluated p24 levels in BSA/PBS buffer alone and CD4+ T cells isolated from uninfected or ART-suppressed HIV+ donors and treated ex vivo with either PMA/Ionomycin or the HDACi vorinostat (VOR). Detectable p24 levels were observed only in HIV+ cell lysates (Figure 1C). Additionally, magnetic bead-based immunodepletion studies were performed using either p24 ELISA antibodies or mouse IgG incubated with CD4+ T cell lysates from HIV+ viremic donors. HIV gag p24 was absent in the flow-through (FT) of p24 antibody–depleted samples but measurable in the IgG control FT, demonstrating specificity of the assay for HIV gag protein (Supplemental Figure 1B). To further assess whether the assay also recognized unprocessed gag polyprotein, recombinant proteins p55 and p24 were diluted 2-fold from 20–0.3 fM in 3% BSA/PBS buffer and measured simultaneously. Linear dilution and low fg/ml detection of recombinant p55 was observed, suggesting recombinant HIV gag p55 polyprotein is also measurable in vitro (Supplemental Figure 1C). It is unclear, however, how efficiently the p24 antibody recognizes gag p55 in vivo. Assay breadth was demonstrated by evaluating viral lysates from 18 genetically diverse clinical isolates, including 8 HIV-1 subtypes spanning groups M and O, and 1 HIV-2 isolate. As shown in Figure 1D, viral HIV gag protein was detected in all 18 clinical isolates, and protein values correlated significantly with RNA copies/μl (r = 0.837; P < 0.001; data not shown).

Latency-reversing agents induce viral p24 expression. Viral protein production is critically important to investigational immunotherapies (5, 17, 18, 32, 33), and thus, it is equally important to assess viral protein production following latency reversal and, subsequently, to determine if the level of viral protein induction on a reactivated cell is sufficient to enable immune-mediated clearance. Classical measures of the HIV reservoir leverage nucleic acid readouts to estimate the burden of disease or viral transcription, but they often
overestimate reservoir size due to the presence of defective proviruses. Techniques that quantify the frequency or percentage of cells that become transcriptionally active following viral reactivation have recently been developed, including the Tat/Rev-induced msRNA limiting dilution assay (TILDA); however, assays like TILDA do not quantify translationally competent proviruses. Toward the initial goal of detecting viral protein upon latent cell reactivation, we assessed the relationship between HIV-1 \( \text{gag} \) protein induction and inducible reservoir size, as measured by TILDA, in 11 ART-suppressed subjects following 48-hour PMA/Ionomycin stimulation. TILDA measurements allowed the segregation of subjects into 3 categories of small, moderate, and large inducible reservoirs (Figure 2A). The majority of subjects displayed inducible p24 levels that trended positively with the TILDA readout; however, a statistically significant correlation between TILDA and p24 was lacking (\( r = 0.54, \, P < 0.1; \) Figure 2B). Notably, a significant correlation was observed between p24 measured in cell lysates and culture medium (\( r = 0.83, \, P = 0.005; \) Figure 2C), and extended studies are underway to assess virus-associated p24 in the culture medium versus contribution, if any, from cellular leakage. Next, we evaluated HIV reactivation in cells treated ex vivo with either 10 nM ingenol-3-angelate, 5 \( \mu \)M bromosporine, or 750 nM VOR and observed comparable trends between TILDA and p24 following 48-hour stimulation of CD4\(^+\) T cells from ART-suppressed HIV\(^+\) subjects (Figure 2, D).
D and E). Measurable induction was also observed at shorter incubation periods (6–24 hours; data not shown). Importantly, we demonstrate that HDACis and other LRAs can elicit both viral RNA and protein expression ex vivo.

VOR and panobinostat induce HIV gag p24 following clinical HDACi administration. HDACis are the most studied latency-reversing agents and have shown induction of cell-associated HIV RNA in vivo using single or multiple dosing paradigms (7–11, 34); however, no study to date, to our knowledge, has demonstrated the direct induction of viral protein after clinical administration of HDACis. To determine if in vivo dosing of HDACis can induce viral protein expression, we examined changes in cell-associated p24 from peripherally isolated CD4+ T cells following clinical administration of 2 HDACis, either VOR or panobinostat. A limited number of available samples were obtained from previously published or recently completed clinical trials (9, 10, 34). Briefly, aviremic HIV+ individuals received either VOR (400 mg) at 72-hour intervals or panobinostat (20 mg) 3 times per week every other week for 8 weeks while maintaining combination ART (9, 10, 34). We quantified changes in p24 from baseline following administration of 1, 2, or 10 doses of VOR (9, 34), as well as multiple doses (6–10) of panobinostat (Supplemental Table 1), for subjects in whom remaining samples were available for p24 assessment. In the VOR study, PBMCs were collected from HIV+ subjects by leukapheresis 3–6 hours after VOR, and total CD4+ T cells were subsequently isolated and cultured with 1 μM raltegravir to prevent further rounds of infection. Peripheral blood cells were obtained by venipuncture on panobinostat-treated subjects at baseline and 8 hours after dose. Similarly, CD4+ T cells were isolated and cultured with 1 μM raltegravir
for 48 hours. Cells were cultured to allow sufficient time for p24 translation to occur, since collection was performed within a few hours of drug administration to enable the primary clinical study endpoint measurements of RNA quantitation. Applying this approach, 3 of 6 VOR-treated subjects showed measurable increases in p24 protein above baseline, primarily after the second dose, and dilution of donor V-1 sample demonstrated linear detection of p24 within the analytical range of the assay (Figure 3A and Supplemental Figure 2). An additional sample was available for a patient receiving 10 doses of VOR and showed p24 induction above baseline in both total and resting memory CD4+ T cell populations (Figure 3B). Both VOR dosing paradigms resulted in increased cell-associated HIV-1 gag RNA in the subjects studied (Figure 3, C and E; refs. 9 and 34). For the participant receiving 10 doses of VOR, p24 was assessed immediately after thawing cells and demonstrates the ability to also detect protein without time in culture, although it is unknown whether or not dependencies exist between the numbers of clinically administered doses, the time of sample collection, and the detection of p24. In the panobinostat study,
p24 measurements were also made on available samples from 5 of 15 participants from the Rasmussen et al. study (10), as shown in Supplemental Table 1. Briefly, 4 participants had matched baseline and post-panobinostat samples, with the panobinostat samples collected the same day as drug administration (participants 6, 8, 17, and 18; Supplemental Table 1). To rule out selection bias, we have included data comparing the fold change in cell-associated HIV RNA at the relevant time points in the panobinostat study for the 5 individuals contributing samples for the present study versus the remaining 10 study participants (Supplemental Figure 3). Four of 4 subjects showed increased p24 relative to baseline following either the seventh panobinostat dose (participants 8, 17, 18) or ninth dose (participant 6; Figure 3D), with the increase being statistically significant ($P < 0.001$). Participant 5 was examined 5 days after the sixth dose of panobinostat (i.e., during the rest period; Supplemental Table 1) and showed no detectable change relative to baseline (Figure 3D). Of note, this patient displayed very low levels of cell-associated RNA during panobinostat treatment and, like only 1 of 15 participants in the original study, plasma HIV RNA from this individual remained continuously undetectable throughout the clinical study (10). Thus, it remains unclear if this represents a return to baseline 5 days after drug administration or if it represents very low p24 levels. Comparison of post-/pre-HDACi ratios of mean cell-associated HIV gag RNA and protein following clinical administration of HDACi showed statistical significant correlations (Figure 3E, $P < 0.01$). Collectively, these data demonstrate that clinical administration of HDACi increase viral RNA transcription and viral protein expression.
Latency reversal induces sufficient viral protein to enable immune-based clearance. As highlighted above, several current immunological strategies are aimed at engaging the immune system to clear HIV-expressing cells (5, 17, 18, 32, 33). We hypothesized that in vivo and ex vivo treatment with latency-reversing agents could result in sufficient viral protein expression to enable redirected CD8+ cell killing and that changes in viral levels could be assessed directly by measuring a reduction in HIV gag p24. To explore this hypothesis, autologous CD4+ and CD8+ T cells were isolated from ART-suppressed HIV+ subjects and coincubated at endogenous levels with either a low concentration of PMA/ionomycin (10 ng/ml PMA + 0.1 μg/ml ionomycin) or 380 nM VOR for 72 hours in the presence or absence of gp120/CD3 or RSV/CD3 bispecific antibodies (BsAbs; Figure 4A). As shown in Figure 4, statistically significant increases in p24 protein were observed over DMSO controls following ex vivo stimulation of CD4+ T cells with either latency-reversing agent (P < 0.01). The addition of gp120/CD3 BsAbs to the culture of HIV+ reactivated cells resulted in reduced p24 protein, comparable with DMSO levels, whereas the addition of the control RSV/CD3 BsAb had no appreciable effect and showed comparable p24 levels with PMA/ionomycin (Figure 4B) or VOR (Figure 4C) treatment alone. We also found that shorter incubations in p24 protein following latency reversal alone and significantly greater than respiratory syncytial virus (RSV) BsAb controls (P < 0.01), thereby suggesting efficient clearance of reactivated latent cells, as well as cells persistently expressing HIV antigen (Figure 4D). CD4+ T cell viability remained similar across treatment conditions (Supplemental Figure 4). These proof-of-concept studies illustrate HDACis, as well as other latency-reversing agents, can elicit viral protein production to facilitate redirected cell killing and support immune clearance approaches under evaluation (5, 17, 18, 35–39).

Discussion

The investment in HIV cure research and the ability to monitor the effectiveness of therapeutic interventions will depend largely on accurate and sensitive quantitation of viral burden in different tissues and body fluids. New ELISA technologies support increased detection of HIV gag p24 in plasma, serum, and cultured cell medium; however, these methods show decreased sensitivity in cell lysates due to matrix interference and in plasma and serum of chronically infected HIV+ subjects due to immune complex formation after seroconversion (29, 30). Here, we report optimized methods that enable sensitive and rapid detection of subfemtomolar cell-associated HIV gag p24 protein in HIV-infected CD4+ T cells from ART-suppressed individuals. Quantifying very small amounts of cell-associated viral antigen produced by low numbers of cells from aviremic/low-reservoir HIV+ individuals is an important advancement, as viral protein expression is believed to contribute to tissue inflammation and persistent immune activation, and induction is a prerequisite for several investigational immune-mediated “kill” strategies (3–5, 40). The assay can be completed in approximately 1 hour, typically requires 1–2 million peripheral CD4+ T cells from ART-suppressed HIV+ individuals, and can be applied to quantify viral protein in cell lysates or culture medium.

It is well established that the majority of HIV proviruses harbor defective genomes and, thus, fail to produce replication-competent virus (20). Traditional assays aimed at reservoir characterization measure HIV DNA or RNA and are sensitive and scalable but have limitations in distinguishing defective from translation- or replication-competent proviruses. This inherently leads to an overestimation of the size of the reservoir (20). In our study, we compared the relationship between HIV-1 gag protein production and inducible reservoir size, as measured by tat/rev multiply spliced RNA (msRNA) (TILDA). Although a positive trend was observed, this did not reach statistical significance, which may be attributable, in part, to poor PCR probe recognition due to mismatches in HIV proviral sequences or to the inability of inducible RNA to make protein. Conversely, the presence of transcriptionally competent “defective” proviruses capable of producing aberrant gag protein that may be poorly recognized by the p24 antibody is also a possible explanation (40). Worth noting, significant correlations were observed between HDACi-induced cell-associated RNA and protein from blood CD4+ T cells in subjects treated clinically with HDAC inhibitors, and further work is ongoing to inform on the correlations between p24 immunoassay and other viral endpoints, including qVOA. Overall, assessment of HIV gag protein provides a tool for measuring protein production following latency reversal or protein levels needed to induce viral cytopathicity or promote immune-mediated clearance; it also affords increased sensitivity as a component of an existing ex vivo assay for measuring HIV reservoir (26–28).
In this study, we show the first direct evidence to our knowledge that clinical administration of HDA-Cis results in measurable increases in HIV-1 gag p24 protein in CD4+ T cells. Additionally, we show ex vivo HDACi treatment of autologous CD4+ and CD8+ T cells from ART-suppressed HIV+ individuals can induce reactivation of HIV and produce sufficient viral antigen to enable redirected immune cell killing, mediated by gp120/CD3 BsAbs. These observations have implications for evaluating HIV-1 eradication strategies, as they highlight application of an ultrasensitive, rapid tool for monitoring the rarer population of translation-competent virus that must be effectively reactivated and cleared. Future studies geared at assessing other viral antigens, sequencing viral genomes, and conducting viral outgrowth studies to assess relationships between defective proviruses producing RNA-competent protein, replication-competent virus, and HIV protein production will be informative to understanding how the generation of viral antigens also contributes to persistent immune activation. Furthermore, in light of the clinical studies using broadly neutralizing antibodies and antibody-based approaches to engage the immune system, this approach becomes increasingly important for assessing the reduction of HIV-expressing cells and the effectiveness of “shock and kill” strategies in preclinical or clinical settings.

**Methods**

**Reagents.** PMA, ionomycin, ingenol-3-angelate, and bromosporine were purchased from Sigma-Aldrich. Raltegravir and VOR were synthesized at Merck. Recombinant HIV gag p24 protein was obtained from US Biological.

**Isolation of T cells.** Leukapheresis was performed on VOR subjects over a period spanning 3–6 hours after dose, and venipuncture was performed on panobinostat subjects at 8 hours after dose. PBMCs were isolated either from leukopaks or from whole blood via Ficoll-gradient. Resting CD4+ T cells, total CD4+ T cells, and total CD3+ T cells were isolated from PBMCs using Stemcell Technologies kits, as previously described (7, 9, 10, 25).

**Ex vivo HIV reactivation.** Enriched CD4+ T cells (1 × 10^6 to 5 × 10^6 cells/ml) were cultured in complete RPMI 1640 media (cRPMI) containing 10% FBS, 2 mM glutamine, and the antibiotics penicillin and streptomycin. Cells were treated ex vivo for indicated times with either 0.1% DMSO, 100 ng/ml PMA/1 μg/ml ionomycin, 10 ng/ml PMA/0.1 μg/ml ionomycin, 10 nM ingenol-3-angelate, 5 μM bromosporine, 750 nM VOR, or 380 nM VOR. For TILDA studies, cells were collected and put through a 35-μm cell strainer (BD Falcon; Thermo Fisher Scientific) to remove cell clumps, and cell yield following titration was greater than 90%. For p24 studies, cells and culture medium were recovered after the treatment period by centrifugation at 1500 g for 5 minutes at room temperature. Cell density and viability were determined with a Vi-cell XR instrument (Beckman Coulter). Samples were processed for p24 digital ELISA and TILDA as described below.

**HIV p24 digital immunoassay.** The fully automated Quanterix HD-1 analyzer for single molecule detection has been reported previously (29–31). In this study, cell lysates were prepared by adding a final concentration of 1% Triton X-100/PBS to 2 × 10^6 cells. For culture medium supernatants, Triton X-100 was added to a final concentration of 1% (10% volume of 10% Triton x-100 prepared in PBS). Samples were stored frozen at –80°C for more than 1 hour or until analysis. Upon thaw, cell lysates were diluted 2- to 5-fold with dilution buffer (50% of blocker casein in PBS [Thermo Fisher Scientific, catalog 37528] and 50% of FBS (Thermo Fisher Scientific, catalog 10438018]). For culture medium supernatants, samples were diluted with an equal volume of dilution buffer and frozen at –80°C until analysis or minimally > 1 hour. Samples were then centrifuged for 5 minutes at 10,000 g at room temperature to remove insoluble material prior to p24 measurement on the Quanterix analyzer. All other assay reagents and assay reaction conditions followed the manufacturer’s p24 kit protocol (29, 30) (Quanterix). p24 concentration was calculated based on a p24 calibration curve with 4-parameter curve fitting.

**p24 assay specificity.** p24 was quantified in BSA/PBS buffer alone or in lysates from uninfected or HIV+ CD4+ T cells in the presence and absence of either PMA/ionomycin or VOR. Additionally, HIV+ CD4+ T cells from viremic donors were lysed (1 million cells/ml) in 1% Triton x-100/PBS and divided into 4 aliquots (1 ml each) in eppendorf tubes. Simoa capture antibody beads (100 μl), or beads with detection antibody or normal mouse IgG (biotin-labeled simoa detection antibody or mouse IgG) were incubated with Dynabeads with streptavidin (M-280) at 4°C for 2 hours. Unbound antibody was washed away, and magnets were added into each tube, mixed, and incubated at 4°C overnight. The supernatant FT after IP was collected by magnet, and p24 in FT was measured in Simoa (Quanterix). BSA/PBS (3%) was negative in the assay, and mock IP lysate was used as positive sample. Immunodepletion results are shown in Supplemental Figure 1B.
In a separate study, MoltIIB cell line (AIDS Research Program) with 100% integrated HIV was resuspended initially at 1 × 10^6 cells/ml culture medium and further diluted to 100 cells/ml. 40 μl (4 cells) was added to 960 μl of 1% Triton x-100 in PBS to become 4 cells/ml lysate. Lysate was further diluted in 3% BSA/PBS from 4–0.125 cells/ml, and p24 was quantified using digital ELISA. Results are shown in Supplemental Figure 1A.

To assess whether the assay also recognized unprocessed Gag polypeptide, recombinant protein p55 (Invitrogen, catalog RP4920) and p24 (US Biological, catalog H6003-25) were measured simultaneously in Quanterix p24 assay. Both proteins were diluted with 3% BSA/PBS starting at 20, 10, 5, 2.5, 1.25, 0.62, 0.3, and 0 fm. Both samples were measured in p24 digital ELISA, and the data were graphed based on the spiking in p24 or p55 concentration (fm) and the signal average enzyme per bead (AEB). Results are shown in Supplemental Figure 1C.

**HIV virus cultures.** HIV isolates representing HIV-1 group M (subtypes A, B, C, D, AE, G) and O and HIV-2 were obtained from the NIH AIDS Reference and Reagent Repository (Germantown, Maryland, USA) or isolated from CD8-depleted PBMCs by coculture with PHA-activated T cells from healthy donors. Virus isolates evaluated included 93RW034, JrFl, Bal, QZ4589, ASM57, 92TH954, 93BR029, BRH84155, BRH95436, ZA/’97/003, 93MW959, 92UG024, THA/’92/006, RU570, JV1083, 301342, 301340, and BCF03. Working stocks were amplified by infecting PHA-activated PBMCs or MT-4 cells stably expressing the CCR5 coreceptor and GFP virus. Viral isolates were harvested from the culture supernatant without additional purification. Diluted virus was inactivated with addition of final 1% Triton x-100 in PBS and incubated at room temperature for 30 minutes and then frozen at –80°C until analysis. The viral lysate was diluted at 1:100, 1:1,000, 1:10,000, and 1:100,000 with 3% BSA/PBS before assay.

**TILDA.** TILDA analysis was performed as previously described (25) with modifications in quantitative PCR (qPCR) in which TaqMan Fast Advanced Master Mix instead of LightCycler buffer was used and performed on a Quantstudio 12K flex instrument (Thermo Fisher Scientific).

**Measurement of p24 from clinical samples.** Cryopreserved cells were thawed, and CD4+ T cells were isolated as described above. 2 × 10^6/ml CD4+ T cells per sample were cultured in the presence of 1 μM raltegravir for 48 hours in cRPMI media and then harvested and prepared for p24 quantification, as described above.

**BsAbs and ex vivo shock and kill.** Antibodies with dual specificity for HIV gp120 and CD3 were generated using published sequences. Antibodies containing a RSV-specific antibody (Synagis) arm and anti-CD3 were also constructed and used as nonspecific control. CD3-enriched cells from HIV -infected ART-suppressed donors were prepared as described above and cultured in cRPMI. Cells were seeded at a density of 2 × 10^6 cells/ml and were incubated with either 0.1% DMSO, 10 ng/ml PMA and 0.1 μg/ml ionomycin, or 380 nM VOR in the presence or absence of 100 ng/ml BsAbs. Raltegravir was added at a final concentration of 1 μM to prevent secondary rounds of replication. Following a 72-hour incubation period, cell lysates were prepared and p24 was quantified, as described above.

**Statistics.** HIV gag protein concentration was calculated based on a calibration curve with linear regression 4-parameter curve fitting. A Spearman correlation test was used for analysis of p24 in cell lysates versus culture medium as well as comparison with TILDA, and P < 0.05 was considered significant. ANOVA or two-parameter student’s t tests were used for statistical significance for all other statistical tests, and the method used is noted in the figure legends.

**Study approval.** HIV-infected subjects receiving stable, standard-of-care ART with plasma HIV-1 RNA fewer than 50 copies per ml and a CD4 count of more than 300 μl^1 for at least 6 months were enrolled in the VOR clinical studies following informed consent (7, 9, 34); HIV-infected adults receiving ART with virological suppression (<50 copies per ml, at least 2 measurements per year) for at least 2 years and CD4 counts above 500 cells per μl were enrolled in the panobinostat study following informed consent (10). VOR studies were approved by the UNC institutional biomedical review board and the FDA (7, 9, 34); panobinostat clinical study was conducted at Aarhus University Hospital, Denmark, and approved by the ethics committee in accordance with the principles of the Declaration of Helsinki (10). For ex vivo studies, additional samples were obtained from HIV+ donors with suppressed viremia (<50 copies per ml) from Philadelphia Fight and Sanguine Biosciences under IRB approval and patient informed consent.

**Author contributions**

GW, MS, D. Graham, JS, NMA, and BJH performed the experiments and analyzed the data. GW, D. Gorman, RJOB, WB, DJH, and BJH provided input on research design. GW, MS, and BJH wrote the manuscript.
TAR, OSS, MT, LØ, DMM, NMA, and RPS oversaw clinical studies, provided clinical samples, and provided cell-associated gag RNA measurements. GW and AT performed statistical analyses. All authors reviewed and provided comments on the manuscript.

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