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Multiplex single-cell visualization of nucleic acids and protein during HIV infection

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Technical limitations in simultaneous microscopic visualization of RNA, DNA, and proteins of HIV have curtailed progress in this field. To address this need we develop a microscopy approach, multiplex immunofluorescent cell-based detection of DNA, RNA and Protein (MICDDRP), which is based on branched DNA in situ hybridization technology. MICDDRP enables simultaneous single-cell visualization of HIV (a) spliced and unspliced RNA, (b) cytoplasmic and nuclear DNA, and (c) Gag. We use MICDDRP to visualize incoming capsid cores containing RNA and/or nascent DNA and follow reverse transcription kinetics. We also report transcriptional “bursts” of nascent RNA from integrated proviral DNA, and concomitant HIV-1, HIV-2 transcription in co-infected cells. MICDDRP can be used to simultaneously detect multiple viral nucleic acid intermediates, characterize the effects of host factors or drugs on steps of the HIV life cycle, or its reactivation from the latent state, thus facilitating the development of antivirals and latency reactivating agents.
Despite progress in nucleic acid visualization techniques, visualization of HIV transcription from individual integration sites has proven elusive. Moreover, there is a need for an integrated approach to simultaneously monitor changes in spliced and unspliced viral RNA (vRNA), viral DNA (vDNA), and proteins at a single-cell level, during the various steps of the HIV replication cycle.

Various approaches have been reported over the past few years, for the combined imaging of HIV nucleic acids and proteins. One of the first approaches to allow visualization of integrated HIV-1 proviruses exploited the recruitment of specific histones to sites of DNA damage, in combination with a reporter virus containing a rare restriction site. This single-cell imaging of HIV-1 provirus (SCIP) approach provided sensitive labeling of integrated provirus, but not unintegrated vDNA, in apparent contrast to later techniques. Others exploited 5-ethynyl-2-deoxyuridine (EdU), which can be incorporated into nascent DNA and then labeled with fluorescent azides by click chemistry. This approach can be used with native virus, rather than a reporter virus, and has been successfully employed in non-dividing cells. The use of EdU is challenging in dividing cells; however, as EdU is incorporated into the genome of the infected cell, generating high background. For nucleic acid labeling in dividing cells, several groups have applied variations of fluorescence in situ hybridization (FISH); either immuno-DNA FISH or branched DNA (bDNA)-FISH. These FISH approaches allowed investigators to examine the vDNA localization at various points during infection, and to identify the number and position of viral integration sites in the host genome. Each method brings strengths and shortcomings, such as being limited to either RNA or DNA labeling, or requiring treatment of the infected cell during reverse transcription to label the viral genome.

Here we describe multiplex immunofluorescent cell-based detection of DNA, RNA and protein (MICDDRP), a bDNA-FISH method with the ability to label the native nucleic acids of the HIV-1 replication cycle, and show how it can be used to track various intermediates of HIV replication, focusing on the kinetics with which various species appear following infection. We follow the appearance of vDNA, nuclear import of vDNA, vRNA transcription from integrated vDNA, splicing of vRNA and nuclear export of vRNA. The ability to visualize these nucleic acid intermediates in the context of viral or host proteins will advance efforts to elucidate mechanisms of antiviral inhibition by small molecules or host restriction factors, enhance our understanding of latency reactivation, and further efforts for novel drug development.

**Results**

**Specific visualization of HIV-1 RNA and DNA.** FISH techniques have been established for detection of nucleic acids in cells, but lack the sensitivity required for some applications, and are often incompatible with immunofluorescent labeling. More recently, bDNA-FISH techniques have been developed to enhance the sensitivity and specificity of RNA detection, (e.g., PrimeFlow, ViewRNA (Affymetrix) and RNAscope) and permit co-staining by immunofluorescence. bDNA-FISH approaches have also been adapted for imaging of HIV-1 nucleic acids.

Based on the RNAscope method, bDNA-FISH protocols that enable visualization of HIV-1 vRNA and vDNA were developed and optimized. Protocols described in Methods section were used with probes that target the gag region of HIV-1 RNA, enabling confocal microscopy-based detection of spliced genomic vRNA in the cytoplasm of cells, shortly after infection with HIV-1 (Fig. 1a, top panel and Supplementary Movie 1). For specific detection of vDNA and not vRNA, probes that target the gag-pol region of negative-strand vDNA were used (to avoid labeling of the positive-strand RNA) and conditions were established to optimize denaturation of dsDNA and hybridization of probes (Fig. 1a, lower panels). Labeling of vRNA and vDNA was highly specific, with no fluorescence detected in uninfected cells (Fig. 1a, left panels). This approach permitted specific detection of vRNA or vDNA, as shown by susceptibility or resistance of the fluorescent signal to treatment with RNase A or DNase I (Fig. 1a). In addition, a protocol for simultaneous observation of both HIV vRNA and vDNA was developed. This protocol enabled monitoring of vDNA without very significant reduction of the vRNA signal. Using this method, co-localization of incoming viral genomic vRNA with nascent vDNA was demonstrated at 4 h post-infection (hpi) (Fig. 1b). Hence, this protocol allows single-cell imaging of vRNA and vDNA individually and multiplexed, with sufficient sensitivity to discern what we interpret to be individual virions. Such highly sensitive staining of viral nucleic acid during entry may be used to track the progress of infection.

**Time course of reverse transcription and nuclear entry.** The protocols for detection of vRNA and vDNA were subsequently used to study the kinetics of HIV infection in a representative cell line (T2M-bl cells). Semi-synchronized cells were infected, fixed, then processed at 2 h intervals for vRNA visualization (Fig. 2). Cells were infected at a relatively low multiplicity of infection (MOI; ~0.2) to prevent saturation of relevant cellular interactions, and foci of vRNA were quantified from multiple randomized images per time point, for more than 300 cells per time point. Under these conditions, automated quantitation produced an estimate of 12 vRNA foci per cell following infection, and 0.26 foci per cell in the absence of infection; the vRNA signal subsequently decreased to an average of 1–2 vRNA foci per cell by 10 hpi (Fig. 2), presumably due to RNase H activity during reverse transcription. In fact, addition of an RNase H inhibitor counteracted the loss of vRNA signal at 12 hpi (Supplementary Fig. 1). These data suggest that reverse transcription is completed by ~10 hpi, which is consistent with previous estimates based on other experimental techniques that follow uncoating and reverse transcription kinetics. Although not part of these analyzes, the signal may be expected to increase again at later time points, as cells begin transcribing vRNA. At this time, it is no longer possible to track incoming genomes; the density of synthesized vRNA may also presents problems for the enumeration of individual foci. At these later time points, measurement of the fluorescent intensity may more accurately represent the extent of vRNA production.

We additionally monitored the appearance and localization of vDNA, as reverse transcription progressed during HIV-1 infection; HIV-1 vDNA was detected mainly in the cytoplasm of TZM-bl cells, and to a lesser degree in nuclei, at 2 hpi (Fig. 3). Both signals increased over time. Total cellular vDNA reached a peak of 1.3 foci per cell, and nuclear vDNA a peak of 0.27 foci per cell (compared to 0.04 and 0.011 foci per cell in uninfected cells), at 12 hpi, presumably as reverse transcription and nuclear import were completed. The number of vDNA foci was then observed to drop between 12 and 24 h, likely reflecting the degradation of unintegrated vDNA; this effect was more pronounced in the cytoplasm of the cell, presumably because the nuclear foci include integrated proviral vDNA, which is stably maintained. Nuclear localization was determined when foci overlapped the nucleus (DAPI stained), in x and y dimensions. To verify that the vDNA signal at 2 hpi truly represented newly synthesized reverse transcription products, virions were attached to cells at 4°C, then stained for vRNA, vDNA and capsid (CA) (Supplementary
It would be expected that only CA and vRNA should be present in HIV-1 particles prior to fusion with a target cell. If a signal was detected for vDNA, that would suggest either that reverse transcription occurred prior to infection, or, more likely, that a significant portion of the vDNA signal is a false positive.

Consistent with the accepted model of HIV-1 replication, we observed signals only for the vRNA probe and CA antibody, not for vDNA, confirming that no vDNA is present prior to infection and that our vDNA probe is specific.

In principle, the probes should label both integrated and unintegrated vDNA. To assess labeling in the absence of integration, TZM-bl cells were infected in the presence of 1 µM integrase inhibitor raltegravir (RAL) followed by DNase I. Probe set PS-3 was used for vDNA (red). Enlarged image shows co-localization between vRNA and vDNA. Scale bars represent 10 µm.

**Transcription from integrated vDNA.** Study of HIV transcription from individual proviruses has been hindered by the lack of an efficient method for simultaneous detection of both vRNA and vDNA in individual cells. By employing this protocol, it is now possible to observe HIV-1 vDNA that is often associated with sites of vRNA transcription (Fig. 4a). Intense accumulation of nascent unspliced vRNA transcripts was observed in ~15% of infected TZM-bl cells shortly after completion of reverse transcription, at 12 hpi, and strictly within the nucleus (Fig. 4b, Supplementary Movie 2); at later time points vRNA distribution matured, with the vRNA transcripts being present predominantly in the cytoplasm (Fig. 2). This phenotype is consistent with reports on transcription of host as well as HIV-1 RNAs occurring in bursts, and accumulation of HIV-1 RNA in the nucleus prior to bursts of vRNA export to the cytoplasm.

**RNA splicing and nuclear export.** Export of unspliced HIV vRNA is dependent on Rev, a protein coded by a multispliced vRNA. It should therefore be possible to detect spliced vRNA in the cytoplasm before the newly synthesized Rev exports the unspliced vRNA. Conditions were established for simultaneous detection of unspliced and multispliced vRNA in individual cells. Probe set 1 (PS-1) (Supplementary Table 1) targets the gag region of the HIV genome and is thus specific for unspliced vRNA.
targets a region common to spliced and unspliced RNA, and thus detects both; the probes bind distinct regions of the vRNA and there is not significant overlap in staining, potentially owing to the length of the molecule or the efficiency of hybridization.

Comparison of the fluorescent signals at various times post-infection illustrates that multispliced vRNA was present in the cytoplasm hours prior to unspliced vRNA (Fig. 4c, d). At later times, presumably following Rev expression, the majority of vRNA, regardless of splicing, localized to the cytoplasm (Fig. 4c, d). These timelines were consistent with the time course of infection, which used a probe for unspliced vRNA; at 12 hpi the vRNA was exclusively nuclear, but by 16 hpi, vRNA was predominantly cytoplasmic (Fig. 2).

**Multiplex imaging of nucleic acid and protein.** Incompatibility with optimal conditions for immunofluorescence staining is a major shortcoming of conventional FISH methods. In contrast,
bDNA-FISH is readily compatible with immunofluorescence\(^3,\ 8\), permitting the visualization of vRNA and viral proteins in virions during cell entry (Fig. 5a) and release (Fig. 5b). We combined the protocol for vDNA and vRNA labeling with immunofluorescence, to simultaneously label vDNA, vRNA, and viral proteins. We defined this technique as MICDDRP. MICDDRP allowed co-visualization of transcription from integrated proviruses and translation of viral protein, in this case Gag or CA, at the single-cell level (Fig. 5c, Supplementary Movie 3). Importantly, this development may enable investigation of selective transcription in cells with multiple nuclear vDNA sites, and the factors that affect viral transcription (Fig. 5d, Supplementary Movie 4).

**Imaging with primary cells and virus isolates.** To verify the potential utility of this method when handling primary tissue samples, a variety of physiologically relevant human cell types were stained, following infection with NL4-3 (Supplementary Fig. 4) or the primary HIV-1 isolate BaL (Supplementary Fig. 5).
These cell types included HEK293T, Jurkat (T lymphocyte cell line), primary peripheral blood mononuclear cells (PBMCs), and monocyte-derived macrophages (MDMs), demonstrating high flexibility for the study of transcription and translation from integrated proviruses in diverse relevant cell types.

A time course comparable to that described in Figs. 2, 3 was also performed using CD4+ T-cells infected with NL4–3, simultaneously stained for vRNA and vDNA (Fig. 6a, b). Similar to the observations with TZM-bl cells, the levels of vRNA were found to be relatively high at early time points, between 4 and 7 foci per cell at 4 hpi, compared to 0.01 foci per cell in uninfected cells. The number of vRNA foci then dropped as reverse transcription proceeded, while vDNA increased over time in measurements both of the whole cell and the nucleus, peaking at 1.3 and 0.6 foci per cell, respectively, following infection, and 0.075 and 0.05 foci per cell in the cell and nucleus, respectively, in the absence of infection. These data demonstrate that the signal to noise ratio is higher for vRNA staining than vDNA (in this low MOI experiment, ~500 to 1 and 10 to 1, respectively), and that vDNA non-specific staining is mainly observed in the nucleus. Treatment of a separate set of samples with RNase A slightly reduced staining in infected and uninfected cells, but did not significantly change the vDNA signal (Fig. 6c). Finally, we compared the peak signals for number of vRNA, vDNA, and nuclear vDNA foci, as well as transcriptionally active cells (Fig. 6d). The signal reduces at each step of the transition from vRNA to actively transcribing cell, as no process is 100% efficient. In this case, we observed a number of vRNA foci per cell that was orders of magnitude greater than the MOI; ~5 vRNA foci per cell yielding an MOI of only 0.05; even at this low MOI, the signal from infection was far higher than the non-specific staining. The conversion rate of cytoplasmic vRNA to an actively infected cell will almost certainly vary depending on cell type used, and potentially between HIV isolates.

**HIV-1 and HIV-2 co-infection**. More than one million people are dually infected with HIV-1 and HIV-2, and co-infection of cells has been described\(^1\). However, co-infections have been challenging to study owing to significant cross-reactivity of reagents. Here, we combined labeling of vRNA and Gag protein to demonstrate that these reagents selectively label HIV-1 or HIV-2 vRNA in infected cells (Supplementary Fig. 6). These approaches were then applied with either epifluorescent or stimulated emission depletion (STED)\(^2\) super-resolution microscopy to detect HIV-1/HIV-2 co-infections in cell culture using virus-specific probes. The data demonstrate that it is possible to identify cells infected by HIV-1, by HIV-2, or by both viruses (Fig. 7). In these experiments, 31% of cells were infected with HIV-1, 17% with HIV-2, and 11% were co-infected, a slightly higher figure than would be predicted by random distribution. Future studies will attempt to determine whether this difference reflects interaction between viruses in co-infected cells. These results demonstrate that individual genomes from HIV-1 and HIV-2 can be observed in the same cell, providing the proof of principle for studies of the interplay between the two viruses in co-infected cells.

**Discussion**

Here we introduce MICDDRP, a microscopy method that is based on bDNA-FISH\(^3\), and that can be used to visualize nucleic acid intermediates during HIV replication. This method has been applied to label the viral molecules at various stages of the replication cycle, including the incoming genomic vRNA at early time points, and the various transcribed vRNAs at late time points. Probe sequences have been selected for specific labeling of
either the unspliced vRNA that codes for Gag and GagPol and is packaged into particles as the genomic vRNA, or that is transcribed from integrated vDNA and exported in a Rev-dependent manner, or for labeling of total vRNA (spliced and unspliced).

Similarly, various vDNA species may be labeled, including the nascent cytoplasmic vDNA (resulting from reverse transcription of the genomic vRNA), integrated proviral vDNA (following integration into the host chromosomes) and non-integrated vDNA (including circular and linear species). Labeling the various species of vRNA and vDNA by this approach is highly sensitive, specific, and compatible with protein immunofluorescence, and can be used to study a wide variety of processes, including reverse transcription of vRNA to vDNA, RNase H activity to degrade vRNA, nuclear import of vDNA, transcription of vRNA from integrated proviral vDNA, splicing of vRNA, export of vRNA from the nucleus, and translation of viral proteins. This approach could also be used for specific questions that we did not address, such as tracking the completion of reverse transcription. Our current vDNA probe labels the first strand of synthesized vDNA, showing that reverse transcription begins in the cytoplasm, consistent with current models of the HIV-1 replication cycle. Labeling of the vDNA requires denaturation of the double-stranded vDNA. If a denaturing protocol were combined with RNase A treatment (to prevent labeling of the positive-stranded vRNA), it should be possible to stain the second (positive) strand of the vDNA, and visualize the site where reverse transcription is completed.

The high sensitivity achieved by bDNA-FISH also lends itself well to diagnostic applications; the high signal to noise ratio results in no “infected” cells being scored in the uninfected controls in our experiments. The RNAscope technology has been applied to identify infected cells in tissue samples, with a sensitivity and specificity comparable to the “gold-standard” method of radiolabeled ISH. The investigators were also able to adapt the

**Fig. 6** Tracking the early stages of infection of Primary CD4+ T cells. Primary CD4+ T cells were isolated from PBMCs, activated, and infected with HIV-1 NL4-3 at an MOI of 0.05. Cells were then fixed and stained at the indicated times. **a** Cells were stained for vRNA (PS-4; green), vDNA (PS-3; red) and nuclei (blue). Foci of vRNA and vDNA were counted in 1500–3500 cells per time point (4–24 h) or 500–1000 cells (no virus control and 72 h). Foci were counted in the complete field of view (whole cell), or within a nuclear mask (nuclear). **b** Number of vRNA and vDNA foci per cell of two independent experiments were calculated. Number of total vDNA foci or vDNA foci within nuclei (nuclear) are also shown. **c** Staining of vDNA was repeated with RNase A treatment. **d** Peak values for foci of vRNA, vDNA, nuclear vDNA, and transcriptionally active cells were calculated, to allow determination of the efficiency of each stage in the infection process. Scale bars represent 10 µm.
were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco), supplemented with 10% fetal bovine serum (FBS) and 2 mM L-glutamine (Gibco), in a humidified incubator at 37 °C with 5% CO2. HEK293-F cells are derived from human embryonic kidney cells, transformed with the SV40 large T antigen (ThermoFisher Scientific; catalog #R70007). They were cultured in the same medium and conditions as described for TZM-bl. Jurkats (NIH AIDS Reagent Program catalog #177) are a human, CD4 and CXCR4 positive, T lymphocyte cell line. They were cultured in Roswell Park Memorial Institute 1640 medium (RPMI; Gibco) supplemented with 10% heat-inactivated FBS and 2 mM L-glutamine, in a humidified incubator at 37 °C with 5% CO2. THP-1 cells (NIH AIDS Reagent Program catalog #9942) are a monocytic cell line that expresses CD14, CCR2, and (following activation) CCR5. They were cultured and used for Jurkats. Monocyte-derived macrophages (MDM) and lymphocytes were purified from peripheral blood mononuclear cells (PBMC; STEMCELL Technologies catalog #7002S) as previously described34, 35. Briefly, MDMs were isolated by adherence to the culture vessel and non-adherent cells were removed and stimulated with 60 U/mL IL-2 (PeproTech) and 4 μg/mL phorbolesterglutamin (Pepco) for 3–4 days prior to infection. Primary lymphocytes were cultured in RPMI supplemented with 10% heat-inactivated FBS and 60 U/mL IL-2, in a humidified incubator at 37 °C with 5% CO2. For time course experiments and clinical isolate infection, primary CD4+ T cells were isolated from PBMCs using EasySep Human CD4+ T Cell Enrichment Kit (STEMCELL Technologies), following the manufacturer’s instructions. T cells were cultured and stimulated in RPMI supplemented with 10% heat-inactivated FBS, 2 mM L-glutamine, 60 U/mL IL-2, and 25 μM/L ImmunoCult Human CD3/CD28/CD2 T-Cell Activator, for 8 days prior to infection. Cells were maintained a humidified incubator at 37 °C with 5% CO2, and every 2–3 days fresh media was added.

**Antibodies and compounds.** Capsid antibody, mouse monoclonal anti-p24 (NIH AIDS reagent Project mab-24-2-2 catalog #6457)36, 37, was used at a 1:2000 dilution and followed by goat anti-mouse secondary antibodies at a 1:2000 dilution, conjugated to either Alexa Fluor 647 (Invitrogen catalog A-21235) or Alexa Fluor 568 (Invitrogen catalog #11004). For HIV-1/HIV-2 co-infection experiments, mouse monoclonal anti-p24 (NIH AIDS reagent Program AG3.0 catalog #4121) was used at a 1:250 dilution to stain both HIV-1 and HIV-2 Gag38. RNase H-inhibitory naphthyridine [4-[(4-aminomethyl-1,1'-biphenyl)-methyl]-1-hydroxy-1,8-naphthyridin-2-one (NAPHRH)] is derived from 1-hydroxy-1,8-naphthyridin-2(1H)-one. It was prepared as described by Williams et al.34. Briefly, the benzyl ester-protected scaffold was saponified, decarboxylated, and the resulting phenol converted to triflate. NAPHRH was formed by Negishi coupling between benzyl glycine or benzyl benzylreduced glycine obtained by treatment with sodium bis(2-methyl diethyl)zinc, followed by Suzuki coupling of the bromide with 4-(N-Boc-aminomethyl)phenylboronic acid and by deprotection with HBr33 (U.S. patent application 20100056516). Raltegravir34 (ISENTRESS/ MK-0518) was obtained from the AIDS Reagent Program.

**Nucleic acid probes.** All probes were purchased from Advanced Cell Diagnostics. To detect HIV-1 vRNA two anti-sense probes were used; the first probe targets the Gag region of HIV-1 (HIV vRNA anti-sense probe 1, PS-1 [probe channel C1 or C2])39, and the second probe targets non-Gag Pol regions coding envelope and accessory proteins (HIV vRNA anti-sense probe 3, PS-3, [probe channel C3]). For HIV-1 DNA detection, a sense probe targeting the Gag-Pol coding region was used, to avoid binding to the vRNA (HIV vDNA sense probe 2, PS-2, [probe channel ID C1]). For detection of HIV-2 vRNA, a probe channel C3 anti-sense probe set targeting the gag region (HIV-2 vRNA anti-sense probe 5, PS-5, analogous to HIV-1 PS-1), was directly diluted into HIV-1 PS-1 probe at a ratio of 1:50. The mixture was added to coverslips as recommended by the manufacturer. Supplementary Table 1 contains a complete list of the probe sets used in this study.

**HIV-1 infection.** Infectious HIV-1 particles were prepared by transfection of 2 × 10⁸ 293FT cells (ThermoFisher Scientific) with 10 μg NL4-3 proviral plasmid40 in a 10 cm dish, using 30 μL Fugene 6 (Promega) in 500 μL Opti-MEM (Gibco). Media were changed 16 h post-transfection. Supernatant was collected 48 h after transfection, cleared by centrifugation for 5 min at 700 g followed by filtration through a 0.45 μm polycarbonate filter, and stored in aliquots at −80 °C. Virus particles were titrated in TZM-bl cells using fivefold serial dilutions in the presence of 20 µg/mL DEAE-Dextran. After 48 h, cells were fixed and the number of blue forming units (bfu) was determined. This titer was used to determine multiplicity of infection (MOI) for future experiments, where MOI of 1 is 1 bfu per cell. It should be noted that differences in experimental conditions (e.g., reduced incubation time, or infecting different cell types) may alter the observed MOI of the experiments, relative to that determined in titration. In these cases, we have quoted the apparent MOI for the experiment, rather than the MOI we would have expected to observe under the titration conditions.

Semi-synchronous infections of TZM-bl cells were performed using 2 × 10⁴ cells plated on 1 mm collagen-coated glass coverslips (GC-12-Collagen, Neuvitro)39. Cells and virus were pre-chilled separately at 4 °C for 10 min, then virus was added to cells and incubated at 4 °C for 20–30 min in the presence of 20 µg/mL DEAE-
In situ vRNA detection. HIV-1 vRNA in cells was probed using RNAscope reagents (Advanced Cell Diagnostics). The manufacturer’s protocol was used with some modifications. Following fixation, cells were dehydrated by removal of DPBS and sequential replacement with 50%, 70% then 100% ethanol, incubating the samples for 5 min at room temperature in each solution. Ethanol solutions were prepared as volume-to-volume ratios in ultrapure water (Synergy; Millipore). The samples were then treated as described for vRNA staining, up to the point of incubation with the probes. Following hybridization, coverslips were immobilized on glass slides; a small drop of nail polish was placed on a glass slide and the coverslip edge was placed on the nail polish drop. Using an ImmEdge hydrophilic barrier pen (Vector Laboratories), a circle was drawn around the coverslip and PBS was added to prevent sample evaporation. The samples were washed twice with DPBS, then incubated for 24 h at 37 °C. Cells were then collected for staining as described for HIV-1 infections above.

HIV-1 and HIV-2 coinfections. HIV-2 infectious particles were produced by transfection of 3 × 10^5 293FT cells with 10 µg HIV-2 ST proviral plasmid in a 10 cm dish, using 30 µL Fugene 6 in Opti-MEM. HIV-1 and HIV-2 particles were added simultaneously to TZM-BL cells, at MOIs of 0.5 and 1, respectively, then incubated for 24 h at 37 °C. Cells were then collected for staining as described for HIV-1 infections above.

In situ vDNA detection. HIV-1 vDNA foci were imaged in cells after vDNA probe hybridization for 2 h at 40 °C. Following vRNA probe hybridization, samples were washed twice in the manufacturer’s wash buffer, then incubated with vDNA probes for 2 h at 40 °C in hybridization buffer, but without heating, to avoid loss of the vRNA signal. The hybridization of pre-amplifiers to the vDNA foci was performed as described for vRNA. Addition of the vRNA probe prior to denaturation of the DNA prevents hybridization of the probe to the coding strand of the vDNA.

Immunostaining. Staining for protein was performed after staining for nucleic acids. Coverslips were blocked with 1% bovine serum albumin (BSA) and 10% FBS in PBS containing 0.1% Tween-20 (PBST) at room temperature for 1 h. Capped protein (CA), or the capped domain of Gag protein, was then probed with anti-p24 diluted 1:2000 in PBST supplemented with 1% BSA and incubated at room temperature for 1 h. Samples were washed twice in PBST at room temperature for 10 min with rocking. Fluorescently labeled secondary antibodies were used at 1:2000 and incubated at room temperature for 1 h, then the samples washed once with PBST with room temperature for 10 min with rocking. Nuclei were stained either using the DAPI provided in the RNAscope kit, or with 0.5 µg/mL Hoechst 33258 in PBS at room temperature for 10 min. Coverslips were washed twice in PBST at room temperature for 10 min with rocking and washed once with PBS for 1 min. Finally, coverslips were mounted on slides using Prolong Gold Antifade, then sealed with nail polish.

Imaging and imaging quantification. Unless otherwise stated, images were taken with a Leica TCS SP8 inverted confocal microscope equipped with a ×63/1.4 oil-immersion objective, and a tunable supercontinuum white light laser. The excitation/emission bandpass wavelengths used to detect DAPI, Alexa 488, ATTO 550, Alexa 568, and Alexa 647 were set to 405/420–408, 488/505–500, 550/560–610, 568/580–630, and 647/655–705 nm, respectively. Confocal data sets were deconvolved using the Huygens Professional software (Version 16.10, SVI). Some images were captured using a Zeiss LSM 880 Airyscan confocal microscope. When images were captured with the LSM 880, a Plan-Apochromat ×63/1.4 oil objective was used in the Airyscan super-resolution mode. Following full 3D capture with a Z-step size of 0.16 microns, raw images were subjected to the standardized Airyscan process (which includes photon reassignment and linear parameter 3D deconvolution). Movies were produced based on volume-rendered Z-stacks of confocal images using LAS X (Leica Microsystems) or ZEN (Zeiss) imaging software, and edited using Fiji.

The numbers of nuclei and vRNA or vDNA foci were quantified using Gen5 software (BioTek), and the number of foci for each image was divided by the number of nuclei in the image to determine the average number of foci per cell. To determine the number of nuclear vDNA foci, a mask was applied the image based on the vDNA stain; vDNA foci that fell within the DAPI mask were counted as nuclear vDNA foci.

HIV-1/HIV-2 coinfection images were acquired with an Olympus IX81 microscope, using a ×40 objective. Super-resolution images of HIV-1/HIV-2 co-infected cells were acquired with a Leica TCS SP8 STED inverted confocal microscope. For the excitation/emission bandpass of the 488/560 nm wavelengths/depletion laser used to detect DAPI, Alexa 488, and ATTO 550 were set to 405/420–480, 488/495–550/592, and 550/560–610/660 nm, respectively.
Data availability. All data supporting the findings in this paper are included in the main text and Supplementary Information. All relevant data are available from the authors upon request.

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Author contributions
S.G.S. conceived the research; S.G.S., M.P.-C, V.Y., and P.R.T. designed experiments; M. P.-C, V.Y., D.L., O.B.U., J.J., J.1W. performed experiments; M.P.-C, A.D.H., O.B.U., A.N. E., P.R.T., and S.G.S. analyzed data; P.R.T., M.P., V.Y., and S.G.S. wrote the paper. All authors reviewed the manuscript prior to submission.

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