Cell Reports

Cellular IP $_6$ Levels Limit HIV Production while Viruses that Cannot Efficiently Package IP₆ Are Attenuated for Infection and Replication

Graphical Abstract

Authors

Donna L. Mallery, K.M. Rifat Faysal, Alex Kleinpeter, ..., Eric O. Freed, Adolfo Saiardi, Leo C. James

Correspondence

lcj@mrc-lmb.cam.ac.uk

In Brief

Mallery et al. demonstrate that HIV is crucially dependent upon the host metabolite $IP₆$ to produce infectious virions. Cells deficient in $IP₆$ produce fewer virions, while virions that fail to package sufficient numbers of IP₆ molecules are poorly infectious and fail to replicate in primary cells.

Highlights

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- \bullet HIV needs cellular IP₆ to efficiently produce and assemble virions
- \bullet HIV packages IP₆ using two rings of lysine residues in immature gag hexamers
- \bullet Viruses that fail to package IP₆ do not mature properly and have unstable capsids
- \bullet Deficient IP₆ packaging prevents HIV replicating in primary cells

Cell Reports Article

Cellular IP_6 Levels Limit HIV Production while Viruses that Cannot Efficiently Package IP_6 Are Attenuated for Infection and Replication

Donna L. Mallery,^{[1](#page-1-0)} K.M. Rifat Faysal,^{[2,](#page-1-1)[6](#page-1-2)} Alex Kleinpeter,^{[3,](#page-1-3)6} Miranda S.C. Wilson,^{[4,](#page-1-4)6} Marina Vaysburd,¹ Adam J. Fletcher,^{[5](#page-1-5)} Mariia Novikova,^{[3](#page-1-3)} Till Böcking,^{[2](#page-1-1)} Eric O. Freed,³ Adolfo Saiardi,^{[4](#page-1-4)} and Leo C. James^{[1,](#page-1-0)[7,](#page-1-6)[*](#page-1-7)}

1MRC Laboratory of Molecular Biology, Francis Crick Avenue, Cambridge, CB2 0QH, UK

2EMBL Australia Node in Single Molecule Science and ARC Centre of Excellence in Advanced Molecular Imaging, School of Medical Sciences, UNSW Sydney, Sydney NSW, Australia

3Virus-Cell Interaction Section, HIV Dynamics and Replication Program, Center for Cancer Research, National Cancer Institute, Frederick, MD 21702-1201, USA

4MRC Laboratory for Molecular Cell Biology, University College London, London, UK

5MRC Protein Phosphorylation and Ubiquitylation Unit, University of Dundee, Dundee, UK

6These authors contributed equally

7Lead Contact

*Correspondence: lcj@mrc-lmb.cam.ac.uk

<https://doi.org/10.1016/j.celrep.2019.11.050>

SUMMARY

HIV-1 hijacks host proteins to promote infection. Here we show that HIV is also dependent upon the host metabolite inositol hexakisphosphate (IP $_6$) for viral production and primary cell replication. HIV-1 recruits IP_6 into virions using two lysine rings in its immature hexamers. Mutation of either ring inhibits $IP₆$ packaging and reduces viral production. Loss of $IP₆$ also results in virions with highly unstable capsids, leading to a profound loss of reverse transcription and cell infection. Replacement of one ring with a hydrophobic isoleucine core restores viral production, but IP_6 incorporation and infection remain impaired, consistent with an independent role for $IP₆$ in stable capsid assembly. Genetic knockout of biosynthetic kinases IPMK and IPPK reveals that cellular IP_6 availability limits the production of diverse lentiviruses, but in the absence of IP_6 , HIV-1 packages $IP₅$ without loss of infectivity. Together, these data suggest that IP_6 is a critical cofactor for HIV-1 replication.

INTRODUCTION

The HIV-1 capsid undergoes a number of transformations during viral replication: assembly, maturation, host interaction, and uncoating. Assembly occurs at the cell membrane, where the viral protein Gag polymerizes into a hexagonal structure called the immature lattice [\(Bush and Vogt, 2014\)](#page-13-0). Once HIV virions bud from the cell, viral protease initiates maturation by cleaving Gag into multiple fragments and disrupting the immature lattice [\(Lee et al., 2012](#page-13-1)). The p24 capsid fragment CA then polymerizes inside virions to form a fullerene cone structure made up of a mature hexamer lattice and 12 pentamers. During post-entry infection, the capsid recruits multiple cytosolic and nuclear pore cofactors and protects HIV as it reverses transcribes its RNA into DNA ([Di Nunzio et al., 2012; Lee et al., 2010; Matreyek](#page-13-2) [and Engelman, 2011; Matreyek et al., 2013; Price et al., 2012,](#page-13-2) [2014; Schaller et al., 2011\)](#page-13-2). Finally, the capsid undergoes uncoating freeing the viral DNA to integrate into the host genome. The molecular mechanisms that drive assembly of the immature and mature lattices during production and that trigger capsid uncoating during infection remain unclear. Understanding these processes is complicated by the conflicting structural requirements imposed by capsid function: the capsid may need to be stable for many hours inside the cell [\(Holmes](#page-13-3) [et al., 2015](#page-13-3)) yet undergo rapid disassembly in the right place and at the right time. Definitions of even these fundamental parameters remain controversial, and there is no consensus about exactly what uncoating means and when and where it happens.

Recently, we reported that the metabolite inositol hexakisphosphate (IP_6) binds to an electropositive pore created by R18 in CA hexamers [\(Jacques et al., 2016; Mallery et al.,](#page-13-4) [2018\)](#page-13-4). We demonstrated that IP $_6$ binding increases HIV-1 capsid stability from minutes to hours and promotes the accumulation of DNA inside intact structures during reverse transcription [\(Mal](#page-13-5)lery et al., 2018; Márquez et al., 2018). Importantly, we also found that HIV-1 packages IP $_6$ in a Gag-dependent manner during viral production. As the mature R18 pore only forms after budding, we postulated that a similar electropositive pore in the immature hexamer provided by K158 and K227 might be similarly used to bind IP_6 , and this was demonstrated in a subsequent crystal structure ([Dick et al., 2018](#page-13-6)). On the basis of these results, we proposed that IP_6 might be an HIV "pocket factor" that regulates stable capsid assembly. Pocket factors are small-molecule ligands that are incorporated into picornavirus capsids and dissociate upon infection to allow uncoating ([Rossmann, 1994; Ross](#page-13-7)[mann et al., 2002](#page-13-7)). Here we have tested the hypothesis that by driving assembly of capsid lattices, IP_6 might promote viral production and, by mediating stable capsid maturation, promote infection. By manipulating IP_6 levels in both cells and viruses, we provide physiological evidence that IP_6 is a critical HIV cofactor for viral replication.

Figure 1. Depletion of IP₆ in IPMK-Knockout Cells Reduces HIV Production but Does Not Affect IP₆ Incorporation or Infectivity (A) Biosynthetic pathway of inositol phosphates, illustrating the central role of IPMK and IPPK in IP $_6$ production.

(B) Analysis of IP₆ levels in IPMK CRISPR/Cas9 knockout clones by TiO₂-PAGE and toluidine blue staining of cell extracts (left), [³H]inositol labeling with SAX-HPLC, and scintillation counting of fractions, normalized to background (right). Synthetic polyP was used as ladder for gel orientation. (C) p24 western blot of pelleted virions showing p24 levels in HIV virions produced from IPMK-KO clones.

(D) Measurement of virus production through quantification of RT in viral supernatants from IPMK-KO clones. Error bars depict mean ± SD of three independent experiments. Values are represented as fold WT virus for comparison. Reduction compared with WT is statistically significant (p < 0.0003 in all cases).

RESULTS

To determine whether IP_6 is a necessary cofactor in either HIV virus production or infection, we sought to alter the availability of $IP₆$ in producer and target cells and determine the capability of virus to package it. IP $_6$ biosynthesis is a complex metabolic pathway involving the conversion of IP $_3$ to IP $_5$ by inositol polyphosphate multikinase (IPMK or IPK2), followed by phosphorylation of IP₅ to IP₆ by inositol-pentakisphosphate 2-kinase (IP5-2K, IPPK, or IPK1) [\(Figure 1](#page-2-0)A). As previous work has shown that genetic deletion of IPMK in mouse embryonic stem cells (ESCs) leads to a marked reduction in IP_6 levels without altering cell viability ([Frederick et al., 2005\)](#page-13-8), we used CRISPR/Cas9 to remove IPMK from human 293T cells [\(Figure S1A](#page-12-0)). Because flux through the mammalian IP pathway is incompletely understood, the effects of knocking out a particular kinase on the levels of different IP species are difficult to predict. Therefore, we directly determined the levels of IP $_5$, IP $_6$, ATP, and GTP in extracts from knockout (KO) clones using titanium dioxide purification ([Wilson](#page-14-0) [et al., 2015](#page-14-0)) with PAGE (TiO₂-PAGE) [\(Losito et al., 2009](#page-13-9)) and comparison with known standards ([Figure S1B](#page-12-0)). We observed significant (\sim 90%) IP₆ reduction in all successful IPMK knockouts, although absolute levels varied among clones [\(Figure 1B](#page-2-0)). $IP₇$ has been implicated in regulating ATP levels [\(Szijgyarto et al.,](#page-14-1) [2011\)](#page-14-1), but although ATP and GTP varied among IPMK clones, there was no consistent trend, suggesting that differences were due to clonal variation. To quantify the relative levels of inositol phosphates in IPMK-knockout clones, we grew cells in inositol-free culture media supplemented with tritiated inositol ([³H]inositol) and used strong anion exchange high-performance liquid chromatography (SAX-HPLC) to isolate specific IP species. Scintillation counting of purified samples confirmed a reduction of both IP₅ and IP₆, with little change in IP₄ ([Figure 1B](#page-2-0)), consistent with the availability of an alternative metabolic pathway to produce IP_4 , albeit a different isomer ([Figure 1A](#page-2-0)).

To test whether a reduction in cellular $IP₆$ levels affects viral production, we transfected a range of IPMK clones with the HIV-1 Gag-Pol expression plasmid pCRV-1 and the HIV-GFP encoding plasmid pCSGW. Analysis of Gag expression revealed that some clones had slightly higher or lower levels than parental cells ([Figure S1C](#page-12-0)). In contrast, all IPMK-knockout clones produced substantially fewer viral particles, as determined by p24 blotting [\(Figure 1](#page-2-0)C) and quantitative ELISA for RT enzyme [\(Fig](#page-2-0)[ure 1](#page-2-0)D). As with viral gene expression, the ratio of RT to p24 showed some random variability between clones but no consistent difference compared with wild-type (WT) [\(Figure S1D](#page-12-0)). To confirm that reduced production in the IPMK knockouts was not the result of clonal variation, we reconstituted IPMK by stable transduction ([Figure S1E](#page-12-0)). Analysis of inositol phosphates in cell extracts by TiO₂-PAGE demonstrated successful restoration of IP₆ levels in IPMK-knockout clones reconstituted with IPMK but not empty vector constructs ([Figure S1](#page-12-0)F). Crucially, ectopic overexpression of IPMK was sufficient to partially rescue virus production in the tested IPMK clones, without altering protein expression levels ([Figure 1](#page-2-0)E; [Figure S1G](#page-12-0)).

We also investigated whether depletion of IP_6 had any impact on $IP₆$ incorporation by virions. We purified virus from two IPMK-KO clones grown in inositol-free media supplemented with $[{}^{3}H]$ inositol as described previously ([Mallery et al., 2018\)](#page-13-5). The presence of IP $_6$ in virions was determined by scintillation counting of fractions collected using SAX chromatography. IP $_6$ was detected in virus produced from both tested IPMK clones, despite the reduced number of viral particles. When normalized for viral particles (CPM/ng p24), we observed no significant difference in $IP₆$ levels between virus produced in parental cells and IPMK knockouts [\(Figure 1F](#page-2-0)).

The above data suggest that cellular IP_6 levels limit the number of HIV virions that are produced but that viruses retain the same capacity for IP_6 incorporation. On the basis of this finding, we hypothesized that once normalized for viral production, HIV virions from IP_6 -depleted cells should be as infectious as those produced in parental cells. Infection experiments carried out at a range of virus dilutions confirmed this prediction [\(Figure 1](#page-2-0)G). To test whether $IP₆$ levels are important in target cells, we challenged parental cells and two IPMK-knockout clones with viruses produced from each of these cell lines. Virus titration showed that all combinations were at least as infectious as parental cell-derived virus in parental cells, with 2_1 virus marginally more infectious ([Figure 1H](#page-2-0)).

Assembly of an immature lattice from recombinant Gag protein can be promoted *in vitro* by both IP_5 and IP_6 ([Dick et al.,](#page-13-6) [2018\)](#page-13-6). To test whether IP₅ can be used by HIV to promote viral production in the absence of available IP_6 , we created CRISPR/Cas9 knockouts of IPPK, the enzyme responsible for conversion of IP₅ to IP₆ ([Figure S2A](#page-12-0)). We used TiO₂-PAGE to show that our IPPK KOs have dramatically reduced levels of cellular IP $_6$ while IP $_5$ was unaffected ([Figure 2A](#page-4-0)). Quantification of IPs following growth in [³H]inositol-supplemented media further revealed that IP_5 levels in two of three IPPK clones were identical to parental cells, but IP_4 was unexpectedly increased in all clones ([Figure 2B](#page-4-0)). This is in contrast to the phenotype in IPK1 yeast knockouts, in which IP₅ but not IP₄ accumulates ([York et al., 1999](#page-14-2)). To determine how the different availability of IP species in IPPK knockouts alters HIV packaging and viral production, we quantified IP incorporation in virions produced from IPPK knockout clones [\(Figure 2C](#page-4-0)). In contrast to virus produced from IPMK KOs, IPPK viruses had little or no IP_6 . However, IP₅ was now detectable in virions produced from

⁽E) Levels of virus production from viral supernatants collected from parental clones and the same cell lines stably transduced with either EV or IPMK. Error bars depict mean ± SD of two independent experiments. Values normalized to the levels in 293T untransduced cells.

⁽F) Quantification of IP₆ packaging in virions produced in wild-type cells and IPMK-KO clones by [³H]inositol labeling, SAX-HPLC, and scintillation counting of fractions then normalized to background.

⁽G) Infectivity of viruses produced in selection of IPMK clones plotted against viral volume input (left) and plotted against quantity of RT in viral supernatants to normalize for differences in production (right).

⁽H) Infectivity of viruses purified from wild-type, 1_7, or 2_1 IPMK-KO producer cells and titrated onto the three different cell lines, normalized to RT levels. Error bars for infection data depict mean ± SD of three replicates from one experiment representative of three independent experiments.

Figure 2. HIV Incorporates IP₅ in the Absence of IP₆ without Loss of Production or Infectivity

(A) TiO₂-PAGE and toluidine blue staining of cell extracts showing IP₅ and IP₆ levels in IPPK CRISPR/Cas9 knockout clones.

(B) Inositol phosphate quantification in selected IPPK-KO clones using ³H-inositol labeling and inositol phosphate fractionation by SAX-HPLC.

(C) Quantification of IP₅ and IP₆ packaging in virions produced in wild-type and IPPK-KO cells through [3 H]inositol labeling, SAX-HPLC, and scintillation counting of fractions.

(D) p24 western blot of pelleted virions showing p24 levels in HIV virions produced from IPPK-KO clones.

(E) Measurement of virus production through quantification of RT in viral supernatants from IPMK-KO clones. Error bars depict mean ± SD of three independent experiments. Values are represented as fold WT virus, and reduction compared with WT is statistically significant (p < 0.0012 in all cases).

(F) Infectivity of viruses from (E), as a function of viral dose measured by RT levels. Error bars depict mean ± SD of three replicates from one experiment representative of three independent experiments.

all tested clones. Two discrete peaks for IP_5 were observed following SAX chromatography, which is likely the result of phosphate jumping between adjacent hydroxyl groups that occurs during IP extraction conditions ([Pisani et al., 2014](#page-13-10)). Summing the two $IP₅$ species reveals that HIV packages a similar number of IP₅ molecules per virion when it is produced in IPPK KOs as it does IP $_6$ when produced in parental cells ([Figure S2B](#page-12-0)). Viral protein expression in IPPK KOs was broadly similar to parental cells, except for 2_7, in which it was reduced ([Figure S2C](#page-12-0)). Importantly, there was a consistent and substantial decrease in virion production in all IPPK clones as assessed both by p24 blot [\(Fig](#page-4-0)[ure 2](#page-4-0)D) and RT incorporation [\(Figure 2](#page-4-0)E). Despite this defect in production, IPPK-derived viruses were as infectious as those from parental cells [\(Figure 2](#page-4-0)F), again consistent with the behavior of viruses produced from IPMK knockouts. These results suggest that HIV can substitute IP $_5$ for IP $_6$ during viral production when the latter is not available and that this does not substantially alter infectivity. The similar decrease in viral production observed in IPPK and IPMK KOs likely reflects that the IP molecule packaged by virus in these cells (IP $_5$ or IP $_6$, respectively) is at similar levels (5-10 uM).

The above data suggest that the availability of IP_5 and IP_6 in producer cells limits the production of HIV virions. This could be a result of impaired Gag recruitment to the plasma membrane or inefficient viral budding. To investigate this, we compared budding sites in WT, IPMK, and IPPK cells using a membrane flotation assay to measure Gag localization ([Kutluay and Bien](#page-13-11)[iasz, 2010; Ono and Freed, 1999\)](#page-13-11). We observed similar levels of Gag in the membrane fractions of each cell line, suggesting that $IP₆$ is not involved in recruitment ([Figure 2](#page-4-0)G). To determine viral release efficiency, we infected cells with VSV-G-pseudotyped env(-) pNL4-3 (pNL4-3/KFS) virus ([Freed et al., 1992\)](#page-13-12) and calculated the cell/virion Gag ratio. In both IPMK and IPPK KOs, there was an increase in this ratio consistent with a reduc-tion in release efficiency ([Figure 2H](#page-4-0)). These data indicate that IP_6 levels limit viral budding rather than recruitment of Gag to budding sites. Moreover, they suggest that IP_6 is limiting for viral production irrespective of whether producer cells are transfected or transduced.

To understand how HIV-1 can substitute IP₅ for IP₆ without loss of infection, we solved the crystal structure of the mature capsid hexamer in complex with the myo isomer of $IP₅$ (inositol 1,3,4,5,6-pentakisphosphate) [\(Table S1\)](#page-12-0) and compared it with our previously solved structure with hexakisphosphate ([Mallery](#page-13-5) [et al., 2018\)](#page-13-5). In mammals, myoIP₅ is the precursor for IP₆ synthesis by IPPK, which phosphorylates the axial 2-OH. As expected, we found that IP $_5$ was coordinated at the center of the HIV-1 hexamer by the ring of arginine residues at position 18 ([Figures 3A](#page-6-0) and 3B). The location of the ligand at the center of the six-fold axis leads to symmetry averaging of six equivalent binding positions. However, in contrast to IP_6 , which with its axial phosphate is significantly less planar, the density for $IP₅$ is much clearer, and the ligand can be unambiguously placed in a parallel stacking arrangement with the R18 ring [\(Figure 3C](#page-6-0)). This stacking conformation is reminiscent of that observed with the nonphysiological compound hexacarboxybenzene and allows multiple hydrogen bonds to form between all five equatorial phosphates and the arginine side chains in the ring ([Jacques et al., 2016](#page-13-4)). The additional axial phosphate present in IP_6 would be orientated away from the R18 ring. Thus the probable reason why $IP₅$ can substitute for IP_6 without loss of infectivity is because interaction is driven largely by the equatorial phosphates, at least when the ligand is in a planar conformation. The packing of IP_5 above the plane of the R18 ring also places it in the cavity bounded by the β -hairpin, meaning that, like IP₆, its dissociation from the hexamer could be regulated by opening and closing of the b-hairpin [\(Figure 3D](#page-6-0)).

The pandemic strain HIV-1 M uses multiple protein cofactors to promote its replication in human cells. However, cofactor use is not always conserved in other diverse lentiviruses. For instance, feline immunodeficiency virus (FIV) can replicate independently of nuclear import proteins that are required by HIV-1 M [\(Lee et al., 2010\)](#page-13-13). We therefore sought to determine whether IP $_6$ is an HIV-1 M specific cofactor or an evolutionary conserved component of lentiviral replication. To this end, we investigated the production of HIV-1 O, HIV-2, simian immunodeficiency virus (SIV), and FIV in cells depleted of either IPMK or IPPK [\(Figure S3\)](#page-12-0). In all cases, we observed a reduction in the viral titer of lentiviruses produced in cells lacking either kinase ([Figure 4](#page-7-0)A). This reduction in titer was accompanied by a decrease in the amount of virions produced in IPMK- or IPPK-KO cells, as assayed by RT ELISA of the supernatants ([Figure 4B](#page-7-0)). The data are in agreement with our findings that reducing the availability of $IP₆$ in producer cells reduces viral production rather than infectivity. Moreover, there was no substantial variation in protein expression (as measured by virally encoded GFP), consistent with a requirement for IP_6 during assembly ([Figure 4C](#page-7-0)). Together this suggests that $IP₆$ is a highly conserved lentiviral cofactor.

Next, we sought to complement our results in IP_6 -depleted cells by mutating residues within Gag that are implicated in $IP₆$ recruitment. Previously, we hypothesized that because the R18 pore only forms after budding, IP_6 may be recruited by two lysines (K158 and K227) that form a similar charged ring in immature hexamers and bind IP_6 *in vitro* [\(Figure 5](#page-8-0)A). Our prediction was that mutation of these lysine residues could potentially affect both production and infection by altering recruitment of IP_6 . Production might be limited because the ability of IP₆ to promote assembly of the immature lattice could be impaired, while infection might be reduced if insufficient IP_6 levels are available inside virions to promote maturation and stabilize the capsid. We mutated each lysine residue to alanine or isoleucine, with the latter residue chosen because K227I has previously been shown to be infectious ([Rihn et al., 2013\)](#page-13-14). As before, we measured Gag production in virus-producing cells and found

⁽G) Membrane flotation analysis of cell lysates from WT, IPMK-KO, and IPPK-KO cells. Western blotting of sucrose gradient fractions for Gag show that similar levels of Gag are associated with the membrane fractions. Gag precursor Pr55Gag (pr55), p41, and mature capsid protein (p24) are indicated. (H) Virus release assays showing levels of Gag in lysates and virions after transduction of WT and KO cells with virus produced in WT cells. Graph shows relative quantification of p24 from two independent experiments and western blots. Representative data from at least two independent experiments are shown, unless otherwise indicated. CPM data are normalized to background.

that Pr55Gag protein expression was at least as abundant as wild-type ([Figure S4A](#page-12-0)). In contrast, all mutants except K227I had reduced viral production as determined both by p24 blotting and RT enzyme quantification ([Figures 5](#page-8-0)B and 5C), with K227I being produced at wild-type levels by both measures.

To test whether the defects in production associated with the lysine mutants were accompanied by a loss of IP_6 recruitment, we measured IP $_6$ incorporation as described above. We were able to obtain data for both K227 mutants but not K158A, because too few virions were recovered during post-production processing for incorporation to be measured. Given that similar levels of K158A and K227A virions are initially produced, this suggests that K158A may be less stable. We observed a substantial reduction in IP_6 packaging per virion ([Figure 5](#page-8-0)D), suggesting that the K227 ring in the immature Gag lattice is involved in recruiting IP_6 from producer cells. The finding that K227A/I mutation reduces IP $_6$ incorporation allowed us to investigate whether IP $_6$ is important for HIV infectivity in addition to its effect on production. Challenging cells with each mutant virus normalized per nanogram RT revealed that removal of either K158 or K227 results in a profound loss of infectivity ([Figure 5](#page-8-0)E). Mutation of K227 reduced infection by up to 10-fold, while K158 mutants were \sim 100-fold less infectious than wild-type. The fact that K227A and K227I caused the same reduction in infectivity, despite the fact that only K227A is defective for production, suggests that mutation at this position affects both processes independently. Meanwhile, the finding that both mutants display similarly reduced infectivity correlates with their similarly impaired levels of incorporated IP $_6$. Despite having reduced IP $_6$ incorporation, production of K227I remained sensitive to cellular IP $_6$ levels and a similar reduction in production was observed in IPMK-

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Figure 3. Structure of the HIV-1 Capsid Hexamer: IP₅ Complex

(A and B) Secondary structure representation of the complex shown from above (outside-facing, A) and side (within the capsid lattice, B). The six equivalent binding positions for IP_5 are shown, together with the location of the R18 ring.

(C) The structure of IP_5 alone and interacting with the six arginine residues at position 18. Electron density ($2F_o - F_c$) centered around the ligand is shown as a mesh contoured at 1.4σ . Putative hydrogen bond interactions between $IP₅$ and R18 side chains are shown as black dashes.

(D) Molecular surface of the capsid hexamer sliced through the middle to reveal the internal cavity where IP_5 is bound. All surface residues are colored gray except R18 (blue), and the β -hairpin (residues 1–12) that forms one end of the cavity (green). IP_5 , positioned above the R18 ring, is also shown.

and IPPK-KO cells as with wild-type virus ([Figure S4B](#page-12-0)). Just as with wild-type virus, this production defect was not due to large changes in viral gene expression [\(Figure S4C](#page-12-0)). Also consis-

tent with wild-type data, knockout of IPMK or IPPK in target cells did not alter K227I infectivity [\(Figure 5](#page-8-0)F).

A defect in IP6 packaging could affect the assembly and function of the mature capsid, such as supporting encapsidated reverse transcription inside the cell. Consistent with this hypothesis, all lysine mutants produced significantly less viral DNA by 4 h post-infection [\(Figure 5](#page-8-0)G). Importantly, within each lysine pair the same magnitude of defect was observed, with K158 mutants giving a more profound reduction in reverse transcription than K227. Taken together, this suggests that a failure in DNA synthesis is behind the reduced infectivity of the lysine mutants. The more profound impact of mutating K158 versus K227 also agrees well with the relative importance of these residues in binding IP $_6$ in the immature hexamer structure: K158 is primarily responsible for binding the five equatorial phosphates, whereas K227 is oriented more toward the single axial phosphate [\(Dick](#page-13-6) [et al., 2018\)](#page-13-6). Finally, we tested whether defective DNA synthesis, and consequently infection, was a kinetic effect by following infection of K227I for 150 h. The fraction of infected cells reached a plateau that remains lower for the mutant than the WT, suggesting that K227I is not merely slower in its infection kinetics [\(Figure 5H](#page-8-0)).

Capsid functions required for infection can be affected at the level of the virus population if maturation becomes less efficient and fewer mature capsids form or if the functional properties of all capsids are compromised. To investigate this question, we first determined whether lysine mutant viruses such as K227I can form capsids that function like wild-type and retain the ability to interact with capsid cofactors inside the cell. Nuclear pore protein Nup358 and the karyopherin TNPO3 have both been implicated as HIV cofactors whose depletion by short hairpin RNA (shRNA) reduces viral infection [\(Schaller et al., 2011\)](#page-14-3). Nup358 binds directly to the

Figure 4. The Requirement for IP₆ during Viral Production Is Conserved in Diverse Lentiviruses

(A) Titers of viral supernatants for lentiviruses produced in WT, IPMK-KO, and IPPK-KO 293T cells. Titers are determined by percentage GFP infection on WT 293T cells.

(B) Quantification of viral production as determined by RT levels in viral supernatants. Values shown are mean ± SD from three independent experiments. Statistically fewer virions are produced in kinase-knockout cells (p < 0.002 in all cases).

(C) Western blots for GFP expression during viral production show similar protein levels for each cell line. Bottom panel shows loading control Cox-IV.

capsid via interaction with the CypA binding loop, while TNPO3 binds indirectly via cargo protein CPSF6 that binds between two monomers of the mature hexamer [\(Price et al., 2012, 2014\)](#page-13-15). Depletion of both Nup358 and TNPO3 by shRNA reduced infection of K227I to a similar degree as wild-type virus [\(Figure 5](#page-8-0)I; [Fig](#page-12-0)[ure S4D](#page-12-0)). Nuclear pore protein Nup153 has also been identified as an important HIV cofactor [\(Matreyek and Engelman, 2011](#page-13-16)). Fusion of an FG-repeat component of Nup153 to TRIM5 generates a synthetic restriction factor TRIM-Nup153, which potently inhibits HIV infection [\(Matreyek et al., 2013\)](#page-13-17). Importantly, Nup153 can only bind intact mature hexamers, meaning that its binding is sensitive to capsid oligomerization [\(Price et al., 2014\)](#page-13-18). Infecting cells expressing TRIM-Nup153 revealed that mutant K227I is as sensitive to restriction as wild-type virus ([Figure 5J](#page-8-0)). Finally, we tested whether K227I virus remained sensitive to the anti-capsid drug PF74. PF74 has bimodal inhibition kinetics, in which at low

(A) View of five subunits of the immature hexamer (on the basis of PDB: 6BHR) showing the lysine side-chains responsible for coordinating IP₆ (blue sphere denotes the ε -amino group). Symmetrically equivalent molecules of IP₆ are shown with the carbon rings in green.

(B) Western blot of pelleted virions to show p24 levels in HIV wild-type and mutant virions.

(C) Quantification of mutant virus production 293T cells as determined by RT levels in viral supernatants. Error bars depict mean ± SD of three independent experiments. Values are expressed as fold change from levels of RT produced in WT virus. The reduction in virus production between WT and K158A, K158I, and K227A is statistically significant ($p = 0.0121$, $p < 0.0001$, and $p < 0.0001$, respectively).

concentrations it reduces infection by competing with cofactors Nup153 and CPSF6 while at high concentrations it causes irreversible and catastrophic uncoating that prevents reverse transcription ([Price et al., 2014\)](#page-13-18). Mutant K227I remained sensitive to PF74 and displayed similar bimodal inhibition kinetics as wildtype ([Figure 5](#page-8-0)K). Interestingly, we did observe some resistance of K227I to intermediate drug concentrations. This is reminiscent of the partial rescue to PF74 inhibition in cells treated with CypA ligand cyclosporin or depleted of CPSF6 [\(Saito et al., 2016\)](#page-13-19). These effects are thought to be due to viruses' becoming less dependent upon nuclear entry by uncoating earlier in the cytoplasm. A similar interpretation when applied to the K227I PF74 inhibition data suggests that a lack of IP_6 may reduce capsid stability. Taken together however, the data collected on capsid sensors suggest that K227I can form infectious particles that appear to be phenotypically normal with respect to host factor dependence and sensitivity to capsid-targeting inhibitors.

The above experiments do not reveal the basis for the infectivity impairment in K227I. We decided to investigate whether this may be due to K227I forming fewer stable virions. We performed single-molecule total internal reflection fluorescence (TIRF) imaging assays to compare the capsid stability of wildtype and K227I mutant virions [\(Figure 6](#page-10-0)A) using our CypA paint method (Márquez et al., 2018, 2019). Virions were immobilized onto coverslips attached to microfluidic flow cells and permeabilized using the membrane pore-forming protein perfringolysin O (PFO) [\(Figure 6A](#page-10-0), step 1). Upon permeabilization, capsids were visualized using fluorescently labeled CypA, whereby the signal of bound CypA is constant while the capsid remains intact [\(Fig](#page-10-0)[ure 6](#page-10-0)A, step 2) and then decays when the capsid uncoats [\(Fig](#page-10-0)[ure 6A](#page-10-0), step 3). Capsid stability was determined by measuring the fluorescence lifetimes of hundreds of individual virions. Example traces of individual wild-type and K227I virions are shown in [Figure 6](#page-10-0)B. Collation of the data into survival curves reveals two phases, ''fast'' and ''slow,'' corresponding to capsids with low and high intrinsic stability (half-lives of <1 min and 5–8 min, respectively) ([Figure 6](#page-10-0)C). Importantly, K227I virions have a higher proportion of highly unstable capsids. This observation suggests that successful K227I maturation occurs less frequently than for wild-type virions, potentially explaining why the mutant has substantially reduced infectivity even though the number of produced virions is the same. Moreover, the proportion of unstable K227I capsids is likely to be even higher than measured, because the CypA-paint does not detect capsids that collapse immediately upon membrane permeabilization.

Repeating our TIRF experiments in the presence of added IP₆ revealed that the population of unstable capsids could not be stabilized even at 100 μ M, providing further support for this hypothesis [\(Figure 6C](#page-10-0)). Importantly, the remaining K227I capsids could be stabilized by $IP₆$ to the same degree as wild-type, resulting in a dramatically increased lifetime. This suggests that infection by K227 is due to a fraction of properly formed and stabilizable capsids, also likely to be those that are sensitive to the capsid cofactors used above. To further correlate the stability of K227I virions on the single-molecule level with their infectivity, we carried out endogenous reverse transcription (ERT) experiments on purified capsid cores. In ERT assays, the level of DNA synthesis that occurs within intact and stable capsids can be measured by adding nuclease to degrade any exposed DNA. These assays use capsid cores extracted and purified from virions, and we typically obtained fewer K227I cores than wild-type, consistent with the fact that the mutant forms fewer stable capsids ([Figure 6](#page-10-0)D). However, upon titration of nucleotides into successfully isolated cores, we observed a similar increase in encapsided DNA synthesis for both viruses [\(Figure 6](#page-10-0)E). Moreover, the accumulation of DNA in both wildtype and K227I capsids could be similarly promoted by the addition of IP $_6$. Addition of IP₅ also promoted the accumulation of DNA, consistent with the infection data of viruses produced in IPPK-KO cells, which incorporate IP₅ instead of IP $_6$. Taken together, the data suggest that K227I can form stable mature capsids, but far less efficiently than wild-type. This is consistent with its decreased incorporation of $IP₆$ and is the likely explanation of its reduced infectivity.

The above experiments were carried out in cell lines in which a single infectious cycle was measured. To investigate the importance of IP $_6$ -recruiting residues K158 and K227 in the context of replicating virus in relevant primary cells, we performed experiments in peripheral blood mononuclear cells (PBMCs) from three independent donors. The replication of wild-type virus and mutants K158A, K158I, K227A, and K227I was determined by quantifying viral production over 20 days. All mutants displayed a profound loss of replication, with only K227I giving measurable production and then only in two of three donors

⁽D) Quantification of IP₆ packaging in mutants K227A and K227I after normalization for background and input virus (per nanogram p24). Both K227 mutants package similarly reduced levels of IP₆ with respect to wild-type virus. Representative data from two independent experiments are shown.

⁽E) Infectivity of lysine mutant viruses normalized to nanogram RT input. Each mutant pair gives similarly reduced levels of infection, with K158A/I largely impaired. Error bars depict mean \pm SD of three replicates from one experiment representative of three independent experiments.

⁽F) Infectious titer of WT and K227I mutant viruses produced in WT 293T, IPMK-KO, and IPPK-KO cells. Infectivity is expressed as infection units (IU) per quantity of input virus (nanogram RT).

⁽G) Levels of reverse transcription products strong-stop (RU5) and post-strand-transfer (GFP) 4 h post-infection. Error bars depict mean ± SD of three replicates from one experiment representative of three independent experiments.

⁽H) Infectivity of WT and K227I virus matched for RT input over 150 h to determine whether K227I could recover infectivity over time. Infectivity is measured using an IncuCyte and determined as proportion of cell area that is GFP positive.

⁽I) HeLa cells stably expressing shRNA control or shNUP358 or shTNPO3 were infected with wild-type or K227I virus. Infection is quantified as infection units (IU) per quantity of input virus (nanogram RT).

⁽J) WT or TRIM-Nup153 expressing HeLa cells were infected with wild-type or K227I virus. The percentage infected cells was determined by percentage GFP positive cells for a range of input virus (quantified by nanogram RT).

⁽K) HeLa cells were infected with wild-type or K227I virus in the presence of anti-capsid inhibitor PF-74. Data are normalized to infection in the absence of inhibitor. Error bars depict mean \pm SD of three replicates from one experiment representative of at least two independent experiments.

Figure 6. Addition of IP6 Is Able to Stabilize Both WT and K227I Mutant Virions

(A) Outline of single-molecule analysis for virus stability. Briefly, viral particles were immobilized, and fluorescence traces were obtained at the locations of individual virions while permeabilizing their membranes with PFO in the presence of fluorescently labeled CypA (1). CypA binds to the capsid, resulting in the appearance of a stable fluorescence signal (2). As the capsid uncoats, the CypA signal disappears (3), and the fluorescence trace returns to background levels. (B) Example traces for individual virions demonstrating the tracking of fluorescence over time. Addition of IP6 to the virions results in a delay in the fluorescence signal reduction signaling uncoating.

(C) Capsid survival curves collated from single-virion traces showing that K227I particles have a high fraction of short-lived capsids in the presence and absence of IP₆, while the remainder of capsids is strongly stabilized by IP₆ addition for both wild-type and K227I particles.

and at substantially reduced levels ([Figure 7\)](#page-12-1). These experiments demonstrate the dependence of HIV replication in primary cells on IP₆-recruiting Gag residues.

DISCUSSION

Here we have tested the hypothesis that IP_6 is a capsid cofactor that promotes HIV-1 production and infection. Reducing cellular $IP₆$ levels by removal of IPMK, which converts the precursor metabolite IP_3 to IP_5 , decreased viral production but did not reduce $IP₆$ incorporation into virions. Consistent with this, viruses produced in IP₆-depleted cells had no infectivity defect. Knockout of the downstream kinase IPPK, which converts IP₅ to IP₆, also decreased virion production without altering virion infectivity. Thus, the reduced infection previously observed using viral supernatant produced in IPPK-deficient cells [\(Dick et al., 2018\)](#page-13-6) was probably the result of a reduction in production rather than the ability of virions to infect, as demonstrated here for both IPPK and IPMK. Our data also show that reducing cellular IP₅ and IP₆ levels affects producer but not target cells. Membrane flotation experiments in kinase KOs indicate that decreased HIV production caused by reduced cellular IP_6 levels is due to a defect in the assembly of virions at budding sites rather than recruitment of Gag to the plasma membrane. However, future studies will be necessary to investigate this process in more depth.

To complement our manipulation of IPs in producer and target cells, we also mutated the HIV Gag protein to determine how altering the ability to sequester IP_6 during assembly affects viral replication. In this second approach, we mutated two lysines in Gag, K158 and K227, that form two charged rings in the immature hexamer and bind IP_6 . We hypothesized that these lysines recruit $IP₆$ to drive assembly of the immature lattice prior to budding and also promote stable capsid formation during maturation. In agreement with this hypothesis, we found that lysine mutation greatly reduced packaging of IP_6 into virions and significantly reduced both viral production and infection. Consistent with a role for IP_6 in assembly of the mature capsid, we also found that the infectivity loss of mutant K227I was accompanied by a dramatic reduction in the frequency of stable mature capsids.

An important difference between our cellular (IPMK and IPPK) and viral (K158 and K227) datasets is that only the latter manipulation results in reduced IP $_6$ incorporation per virion. This is likely because the lysine mutants have a reduced affinity for IPs and do not efficiently recruit them. Such a conclusion is supported by *in vitro* data showing that although K158 and K227 mutants form immature VLPs with near wild-type efficiency, they are unresponsive to the hugely assembly-enhancing effect of IP_6 [\(Dick et al., 2018](#page-13-6)). Interestingly, we observed that K227I did not have a virus production defect, in contrast to K227A. We speculate that this may be because the substitution of an isoleucine allows hydrophobic packing in the core of immature hexamers to compensate for partial occupancy and weaker binding of $IP₆$ within the forming lattice. The failure of isoleucine to restore production in K158I may reflect the fact that this lysine ring is dominant in IP $_6$ coordination and essential for IP $_6$ promotion of assembly. Consistent with the greater importance of K158 in $IP₆$ engagement, mutation of this residue gave the most profound infection and DNA synthesis defect.

The data here suggest that the previous report of abolished infectivity in K158A and K227A is actually the result of combined production and infectivity defects [\(Dick et al., 2018](#page-13-6)). There may be several reasons why mutation of the immature lysine rings and diminished virion incorporation of IP_6 reduces infectivity. One explanation, supported by our data, is that IP_6 is required to stabilize the mature lattice and promote capsid formation during maturation of budded virions. TIRF imaging of K227I capsids reveals that many are highly unstable and collapse with a half-life of <1 min. A reduction in the frequency with which stable K227I capsids form correlates with their reduced IP_6 packaging and explains their lower per virion infectivity. This explanation is also supported by previous *in vitro* experiments, in which IP₆ was shown to promote mature lattice assembly ([Dick et al.,](#page-13-6) [2018\)](#page-13-6). It would nevertheless be useful to further investigate how *in vitro* virion properties predict in-cell stability and infectivity. This could be accomplished in future experiments by quantifying the structural defect associated with K227I using quantitative electron cryotomography. Interestingly, although cellular concentrations of $IP₆$ were sufficient to promote immature lattice assembly (10 μ M), significantly higher IP₆ concentrations than those in the cytosol were required to efficiently promote formation of a mature lattice $(>250 \mu M)$ [\(Dick et al.,](#page-13-6) [2018\)](#page-13-6). This is consistent with the finding that IP_6 is enriched into budded virions, creating an artificially higher $IP₆$ concentra-tion during maturation ([Mallery et al., 2018\)](#page-13-5). This IP $_6$ concentration dependence may help explain why formation of the immature lattice is favored during assembly at the plasma membrane.

Although the data here support the role of IP_6 post-assembly, they do not demonstrate a requirement post-entry. That there was no infectivity loss for any virus, WT or mutant, in IPMK or IPPK cells suggests either that IPs are no longer needed, virally packaged ligands are sufficient or that viruses can obtain sufficient quantities during infection despite the reduced levels. Previously we have shown that IP_6 greatly stabilizes the mature HIV capsid after it has formed ([Mallery et al., 2018\)](#page-13-5). Without IP $_6$, capsids fall apart on a timescale (minutes) that would not support DNA synthesis or cellular infection. Our results here also show that although K227I has fewer intact capsids, those that form can be stabilized by IP_6 in both TIRF microscopy and encapsidated RT assays. These findings provide a mechanistic basis for the mutants reduced but measurable infectivity. They also support the notion that a charged polyanion is required post-entry to maintain capsid stability, albeit that this may be an abundant molecule, like ATP, obtained from the target cell. Capsid stability may be important not just to promote infection but also to evade immune sensing. A failure to incorporate sufficient $IP₆$ could reveal HIV to pathogen receptors in target cells by

⁽D) Capsid core yields determined by RT quantification (representative of at least two independent experiments).

⁽E) ERT assay using cores isolated from WT and K227I virions, showing a dNTP titration in the absence of IPs or in the presence of either IP₅ or IP₆. Both WT and K227I cores are competent for ERT and are stabilized by both IP₅ and IP₆ to the same extent. Data are average of three biological replicas and have been normalized to the copies of RU5 measured in the absence of dNTPs.

Figure 7. Capsid Mutants Prevent Efficient Replication in PBMCs

PBMCs from three donors were infected with viruses carrying capsid mutations K158A, K158I, K227A, or K227I for 2 h. Cells were cultured over 20 days and supernatants sampled every other day and assayed for RT activity.

reducing capsid stability or increasing the number of incomplete cores. For instance, this could lead to exposure of the viral genome or synthesized DNA to the STING/cGAS pathway.

A curious difference between IP_6 use in assembly versus maturation is that whereas immature hexamers use two lysine rings to engage IP $_6$, mature hexamers appear to use a single arginine ring. An attractive hypothesis is that the two lysine rings help preferentially recruit IP $_6$. The location of K227 immediately below K158 in the immature hexamer is favorable for coordinating the axial phosphate in IP_6 . IP₅ lacks an axial phosphate, as it is IPPK that modifies the inositol 2-OH when it converts the molecule into IP $_6$ (Gonzá[lez et al., 2010\)](#page-13-21). This means that IP₅ could not make the equivalent interaction with the K227 lysine ring as IP $_6$, thereby thermodynamically discriminating against $IP₅$ binding. Nevertheless, our data show that HIV can use IP₅ when IP₆ is not available. Moreover, this substitution does not seem to drastically decrease either viral production or infection. We did observe decreased virus production in IPPK KOs, where IP₅ rather than IP₆ was specifically incorporated into virions, but this was comparable with the decrease in IPMK KOs, in which IP $_6$ levels were reduced to a similar level. The cellular concentration of IP $_6$ is normally 5fold higher than IP_5 , suggesting it is IP availability that is the primary determinant of incorporation and the reason why HIV-1 is normally highly selective for IP_6 in wild-type cells ([Mallery](#page-13-5) [et al., 2018\)](#page-13-5).

In conclusion, our data support a model in which IP $_6$ is recruited from producer cells during assembly of HIV at the plasma membrane ([Dick et al., 2018; Mallery et al., 2018\)](#page-13-6). IP $_6$ is recruited by two lysines (K158 and K227) in the immature Gag hexamer, where it helps drive formation of the immature lattice. During maturation, coordination of IP $_6$ is disrupted when the immature lattice is cleaved by viral protease. IP $_6$ is then available to promote assembly of the mature capsid by coordinating the R18 pore in mature hexamers. Once inside the cell, IP $_6$ or another polyanion, such as ATP, may help stabilize the capsid as it transits the cytosol and undergoes DNA synthesis. Finally, and by analogy with picornavirus pocket factors, it is possible that engagement of cellular cofactors disrupts polyanion binding, allowing rapid uncoating to take place. The immature lattice lysine rings K158 and K227 and a mature capsid charged ring (e.g., R18) are highly conserved across diverse lentiviruses [\(Figure S5\)](#page-12-0), suggesting that IP $_6$ is an essential component of lentiviral replication.

STAR+METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.celrep.2019.11.050) [celrep.2019.11.050.](https://doi.org/10.1016/j.celrep.2019.11.050)

ACKNOWLEDGMENTS

This work was supported by the MRC (UK; U105181010), a Wellcome Trust Investigator Award (200594/Z/16/Z), a Wellcome Trust Collaborator Award (214344/A/18/Z), and the NHMRC (Australia; APP1100771 and APP1158338). A.S. was supported by the MRC (MC_UU_00012/4).

AUTHOR CONTRIBUTIONS

Conceptualization and Writing, L.C.J. and D.L.M.; Formal Analysis, D.L.M., K.M.R.F., A.K., M.S.C.W., M.V., A.J.F., T.B., E.O.F., A.S., and L.C.J.; Resources, M.N. and E.O.F; Editing, all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: April 23, 2019 Revised: October 11, 2019 Accepted: November 12, 2019 Published: December 17, 2019

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STAR+METHODS

KEY RESOURCES TABLE

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents, which may require a completed Materials Transfer Agreement, should be directed to and will be fulfilled by the Lead Contact, Leo C James [\(lcj@mrc-lmb.cam.ac.uk](mailto:lcj@mrc-lmb.cam.ac.uk)).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cells & Plasmids

293T CRL-3216 cells were purchased from ATCC and authenticated by the supplier. All cells are regularly tested and are mycoplasma free. Gag/Pol and GFP vectors were, respectively, for HIV-1 pCRV-1([Zennou et al., 2004](#page-14-4)) and CSGW([Naldini et al., 1996](#page-13-28)); for HIV-2, HIV-2 ROD Gag/Pol and HIV-2 GFP[\(Griffin et al., 2001\)](#page-13-22); for HIV-1 O p8.91 MVP([Ikeda et al., 2004](#page-13-23)); for SIVmac, SIV3⁺ and SIV-eGFP([Poeschla et al., 1998](#page-13-24)) and for FIV FP93([Poeschla et al., 1998](#page-13-24)). Lentiviral packaging plasmid pMDG2, which encodes VSV-G envelope, was used to pseudotype infectious virions (Addgene plasmid # 12259). Cells for depletion of TNPO3 and NUP358 were produced by expression of shRNAs as previously described[\(Schaller et al., 2011\)](#page-14-3). HEK293T and HeLa cell lines were maintained in Dulbecco's modified Eagle's medium (DMEM) with 10% FBS, 2 mM L-glutamine, 100 U/ml penicillin, and 100 μ g/ml streptomycin (GIBCO) at 37°C with 5% CO₂. PBMCs were stimulated with 2 µg/ml PHA-P for 4 days before infection, then cultured in RPMI 1640 with 10% FBS, 2mM L-glutamine, 100 U/ml penicillin, 100 µg/ml streptomycin, and 50 U/ml IL-2.

METHOD DETAILS

Infection Experiments

Infections of HeLa cells were performed in the presence of 5 mg ml-1 polybrene. GFP expressing cells were enumerated on a BD LSRII flow cytometer (BD Biosciences) 2 days post-infection after fixation of cells in 4% paraformaldehyde. Values are the mean \pm standard deviation. For PF74 inhibition experiments, the compound was dissolved in DMSO or DMSO and diluted in complete DMEM supplemented with polybrene as above and added to cells shortly before infection. Infections were carried out with sufficient virus to result in 10%–30% infection. Stable TNPO3 and Nup358 depletion experiments were performed by transducing HeLa cells (1 \times 10⁵) with retroviral vectors (pSIREN RetroQ) expressing specific short hairpin RNA (shRNA). Cells were selected with 10 µg/ml puromycin and stable cell-lines used in infection experiments. For further details see [Schaller et al. \(2011\)](#page-14-3). For TRIM-Nup153 restriction experiments, a construct containing Nup153 residues 896-1475 fused to the C terminus of the tripartite domains of TRIM5 (as described in [Matreyek et al., 2013](#page-13-17)) was transduced into HeLa cells and selected with puromycin. In all cases, infection was carried out in 4 \times 10⁴ HeLa cells at a multiplicity of infection (m.o.i.) between 0.1-0.3. Where viruses were quantified for levels of RT enzyme, a colorimetric RT assay kit (Roche) was used according to manufacturer's instructions.

Production of [³H]inositol-Labeled Virus and Cells

[³H]Inositol labeled viruses were prepared as previously described[\(Mallery et al., 2018](#page-13-5)). Briefly, 1 \times 10⁶ 293T cells were seeded into 2×10 cm dishes in inositol-free DMEM and left to adhere overnight. The media was replaced with 5ml inositol-free DMEM supplemented with 5 µCi/ml [³H]inositol (Perkin Elmer). After 3 days incubation, an additional 5ml inositol-free media containing 5 μCi/ml [3 H]inositol was added onto cells, which were then transfected with 2μg each pCRV GagPol and CSGW. For determining IP levels in cells, the same procedure was used except without transfection of viral plasmids. Cells were left for a further 3 days to produce HIV-1 viral particles. After 3 days, either the cells were pelleted for IP determination or viral supernatants were collected. Viral supernatants were topped up to 30ml and pelleted over a 5ml 20% sucrose cushion) in a SW28 rotor (Beckman) at 28,000 rpm at 4° C. Pellets were resuspended in inositol free media and pelleted as previously. After the second spin, pellets were resuspended in 1ml PBS and spun at 13,000rpm at 4°C in a bench top microfuge for 60 min. Pellets were frozen at -20° C until processing. Cells were washed with PBS, then harvested by scraping, counted, and pelleted for quantification of cellular IP $_6$ labeling. Pellets were frozen at -20°C until processing. For comparison of virion and purified capsid core samples, p24 levels were determined by ELISA for p24 (Perkin Elmer).

Purification and Analysis of Inositol Phosphates

Analysis of unlabeled inositol phosphates was performed following previously described protocols([Pisani et al., 2014, Wilson and](#page-13-10) [Saiardi, 2018](#page-13-10)). Cells were extracted using 1M perchloric acid and inositol phosphates purified using titanium dioxide (TiO₂) beads. Extracts were subsequently separated using polyacrylamide gel electrophoresis (PAGE) and visualized using Toluidine blue staining. Inorganic polyphosphate (polyP; Sigma Aldrich S6128) was used as a ladder. The relative IP₆ levels in kinase knockout cells were compared to 293T wild-type cells.

Inositol phosphates extraction and analysis by HPLC was performed modifying a previously described protocol([Azevedo and](#page-13-29) [Saiardi, 2006](#page-13-29)). Cells labeled with [³H]inositol as described above were resuspended in 200 µL of extraction solution (1M Perchloric acid, 5mM EDTA) and incubated on ice for 10 min. The samples were spun out at 13,000rpm at 4°C for 5 min and the supernatant recovered. Viral or cell pellets were extracted for 10 min at 100°C using 200 μ L of extraction buffer and spun out as before. Supernatants from acid extractions were neutralized to pH6-8 using 1M Potassium carbonate, 5mM EDTA (approx. 100 µl) and incubated on ice with the lids open for 1-2hrs. Samples were spun at 13,000rpm at 4°C for 5 min and supernatant containing inositol phosphates was loaded onto HPLC or stored at 4° C. Extraction of IP at low pH and high temperature can induce phosphate jumping; the movement of a phosphate group to a free hydroxyl, resulting in IP isomerization. Therefore, $I(1,3,4,5,6)$ P5, the single IP₅ isomer detectable in IPPK KOs, becomes two IP₅ species when preparing samples of viral particles.

Inositol phosphates were resolved by strong anion exchange chromatography Sax-HPLC on a Partisphere SAX 4.6° - 125 mm column (Hichrom). The column was eluted with a gradient generated by mixing buffer A (1mM EDTA) and buffer B (1mM EDTA; 1.3 M (NH4)2HPO4, pH 4.0) as follows: 0–5 min, 0% B; 5–10 min, 0%–30% B; 10–85 min, 30%–100% B; 85–95 min, 100% B. Fractions (1 ml) were collected and analyzed by scintillation counting after adding 4 mL of Ultima-Flo AP LCS537 cocktail (Perkin Elmer).

Preparation of HIV-1 Virions

Replication deficient VSV-G pseudotyped HIV-1 virions were produced in HEK293T cells using pMDG2, pCRV GagPol and CSGW as described previously([Price et al., 2014](#page-13-18)). pNL4-3 Virus Stocks were produced in HEK293T cells by transfection with HIV-1 proviral DNA (pNL4-3) using polyethylenimine (PEI) [\(Brissault et al., 2003\)](#page-13-30). VSV-G-pseudotyped virus stocks were generated from cells co-transfected with HIV-1 proviral DNA harboring the KFS mutation in *env* ([Freed et al., 1992](#page-13-12))) and the VSV-G expression vector pHCMV-G [\(Yee et al., 1994\)](#page-14-5) at a DNA ratio of 10:1. Viral supernatants were filtered through a 0.45- μ m membrane at 48 hours post-transfection and virus was quantified by measuring RT activity.

Virus Release

293T WT and IPMK/IPPK KO cells were transduced with VSV-G-pseudotyped HIV-1 KFS (*env*-) at 1 RT(cpm)/cell. At 48 hours posttransduction, viral supernatants were ultra-centrifuged and virions were pelleted through a 20% sucrose cushion at 4° C. Virus pellets and remaining cells were lysed and western blotted for Gag. Imaging and band quantification were performed using the Sapphire Biomolecular Imager and Azure Spot analysis software (Azure Biosystems). Virus release was calculated using the following formula.

> *virus p*24 *virus p*24 + *cell p*24 + *cell Pr*55

Replication Experiments

Peripheral blood mononuclear cells (PBMCs) from three donors were infected for 2 hours with RT-normalized virus stocks prepared by transfection of 293T cells with PEI. Cells were washed and cultured in RPMI 1640 medium with 10% FBS with penicillin (100 U/ml) and streptomycin (100 μ g/ml) in a 24-well plate. Supernatants were collected every other day and virus replication was quantified by RT activity.

Membrane Flotation Analysis

Membrane flotation analysis was based on published protocols[\(Kiernan et al., 1998, Kutluay and Bieniasz, 2010, Ono et al., 2000\)](#page-13-31). Briefly two 10cm dishes of 293T cells were seeded and transfected as for virus production. 48hrs post-transfection cells were washed twice with ice cold buffer (10mM Tris pH7.4, 100mM NaCl, 1mM EDTA) and harvested into 1ml of the same buffer. Cells were pelleted and resuspended in 500µl Hypotonic buffer (10mM Tris pH7.4, 1mM EDTA) with protease inhibitors. Cells were lysed by passing through a 25G needle multiple times, adjusted to 150mM NaCl, 1mM MgCl2 and spun at 1000 g for 10min at 4° C for 10min to remove nuclei and intact cells. Lysates were mixed to 80% sucrose and layered with 65% and 10% sucrose and centrifuged overnight at 35K rpm. 1ml fractions were removed from gradients from the top and TCA precipitated. Pelleted protein was resuspended in LDS Sample Buffer and western blotted for Gag.

CRISPR/Cas9 Knockouts

Knockout cell lines were largely produced following the IDT protocol. Briefly, 2.5µl each of crRNA and tracrRNA at 100µM were annealed. This was then mixed with 80-100 pmol Cas9 protein and incubated at 37°C for 10 min to form the Cas9/RNP mix which was kept on ice until use. 2µl of the Cas9/RNP mix was transfected into 8×10^5 cells using the Neon Transfection system, according to manufacturer's protocol (Thermo Fisher). Cells and RNP were immediately transferred into a tube containing 1ml pre-warmed antibiotic free media and subsequently into a flask containing 4ml complete media. Cells were allowed to recover for 24-72hr until being sorted to single cells. Single cell clones were grown up and DNA extracted and sequenced to analyze for CRISPR targeting. Sequences were analyzed by CRISP-ID software[\(Dehairs et al., 2016](#page-13-32)) and manual decoding.

Western Blotting

Samples were run on 4%–12% Bis Tris gels and transferred onto nitrocellulose membranes using iBlot (Life Technologies) and detected by ECL or by Li-COR for quantification. Anti-HIV p24 (183-H12-5C) was obtained from the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH: Anti-HIV-1 p24 Monoclonal (183-H12-5C) (Cat# 3537) from Dr. Bruce Chesebro and Kathy Wehrly [\(Toohey et al., 1995, Wehrly and Chesebro, 1997](#page-14-6)), anti-IPMK (NC12) from Genetex, anti-Nup358 (A301-797A-T) from Bethyl Laboratories, and anti-GFP (ab6556), anti-TNPO3 (ab109386) both from Abcam. For virus release, samples were subjected to SDS-PAGE (4%–20%), then transferred to a polyvinylidene fluoride (PVDF) membrane (Immobilon, Millipore) via semi-dry transfer (Bio-Rad Trans-Blot Turbo). The membrane was blocked for 1 hour with 5% nonfat milk in Tris-buffered saline + 0.05% Tween 20 detergent (TBST) and incubated overnight at 4C with anti-HIV IgG. The membrane was then washed with TBST and incubated for 2 hours with anti-human horseradish peroxidase-conjugated secondary antibody and washed again. SuperSignal West Pico PLUS (Thermo Scientific) was used to reveal protein bands.

Crystallization, Structure Solution, and Analysis

CA hexamer protein was prepared exactly as described previously([Mallery et al., 2018](#page-13-5)). Crystals were grown at 17°C by sitting-drop vapor diffusion in which 100 nL protein was mixed with 100 nL precipitant and suspended above 80 µl precipitant. The structure was obtained from 12 mg/ml hexamer mixed with 14% PEG550 MME, KSCN (0.15 M), Tris (0.1 M, pH 8.5) containing 1mM of myo-IP₅ and cryoprotected with precipitant supplemented with 20% MPD. Crystals were flash-cooled in liquid nitrogen and data collected at beamline I24 at Diamond Light Source. The datasets were processed using the CCP4 Program suite([Winn, 2003\)](#page-14-7). Data were indexed and integrated with iMOSFLM and scaled and merged with AIMLESS([Evans and Murshudov, 2013](#page-13-25)). Structures were solved by molecular replacement using the model 6ES8 in PHASER([McCoy, 2007\)](#page-13-26) and refined using REFMAC5([Murshudov et al., 1997](#page-13-27)). Between rounds of refinement, the model was manually checked and corrected against the corresponding electron-density maps in COOT([Emsley and Cowtan, 2004](#page-13-33)). Final figures were rendered in The PyMOL Molecular Graphics System, Version 1.5.0.4 Schrödinger, LLC. The model and data were deposited in the PDB database with code 6R8C.

Virus Production for TIRF Microscopy

Replication deficient HIV-1 virions without envelope protein were produced in HEK293T cells using pCRV-1 GagPol and CSGW, biotinylated using EZ-Link Sulfo-NHS-LC-LC-Biotin (Thermo Scientific, 21338) and purified as described (Márquez et al., 2018, 2019).

CypA Expression and Purification

CypA was expressed in *E. coli* using a pET-21 vector and purified as previously described(Márquez et al., 2018). For CypA paint, CypA was labeled with Alexa Fluor 568-C2-maleimide (AF568) and binding of conjugated CypA verified by surface plasmon resonance.

TIRF Imaging of Capsids in Permeabilized Viral Particles

TIRF microscopy was carried out following the published method of Márquez et al. (2018, 2019). Briefly, biotinylated viral particles were captured onto coverslips attached to microfluidic flow cells and imaged using a custom built TIRF microscope with an ASI-RAMM frame (Applied Scientific Instrumentation), a Nikon 100 x CFI Apochromat TIRF (1.49 NA) oil immersion objective and NicoLase laser system. Immobilized virions were treated with imaging buffer containing 200 nM PFO, to permeabilize the lipid envelope, and labeled CypA (0.5 - 1 µM), to detect the capsid. TIRF images were then acquired with a frequency of 1 frame/6 s using a 561 nm laser with a 20 ms exposure time for excitation and an Andor iXon 888 EMCCD camera for detection. Single-virion fluorescence traces were extracted from the TIRF image stacks using the JIM Immobilized Microscopy analysis package ([https://github.](https://github.com/lilbutsa/JIM-Immobilized-Microscopy-Suite) [com/lilbutsa/JIM-Immobilized-Microscopy-Suite](https://github.com/lilbutsa/JIM-Immobilized-Microscopy-Suite)) and further analyzed in MATLAB (The MathWorks, Inc) using software adapted from previous work(Bö[cking et al., 2011\)](#page-13-34). Briefly, the duration of the CypA signal was extracted from fluorescence traces by stepfitting using change point analysis. Capsid stability was quantified as the time difference between acquisition of Alexa Fluor 568-CypA upon permeabilization and loss of fluorescence upon capsid uncoating.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical Analysis

Unless otherwise indicated, statistical analyses were performed with the Student's t test using GraphPad Prism 7 software (GraphPad). The number of experiments or (biological) replicates (n) used for the statistical evaluation of each experiment is indicated in the corresponding figure legends. The data are plotted as a mean \pm SD or SEM as indicated. Where the mean is calculated from technical replicates, it is representative of multiple experiments that have been repeated at least twice as described within the figure legend ([Figures 1F](#page-2-0), [2C](#page-4-0), and [5D](#page-8-0)). Data from infection experiments are plotted as the mean \pm SD of three replicates from one experiment representative of three independent experiments [\(Figures 1G](#page-2-0) and 1H, [2](#page-4-0)F, [4](#page-7-0)A, and [5](#page-8-0)E). Where the data are given with biological errors, this has been normalized to allow multiple independent datasets to be directly compared and is described within the figure legend [\(Figures 1](#page-2-0)E, [2](#page-4-0)E, and [5](#page-8-0)C). The single molecule TIRF data in [Figure 6](#page-10-0) was analyzed MATLAB and full details are described in the Methods section above.

DATA AND CODE AVAILABILITY

The published article includes an X-ray structure, which has been deposited in the PDB database with the code 6R8C. [Table S1](#page-12-0) gives the data collection and refinement statistics. A full PDB validation report is also provided.

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Supplemental Information

Cellular IP₆ Levels Limit HIV Production

while Viruses that Cannot Efficiently Package IP $_6$

Are Attenuated for Infection and Replication

Donna L. Mallery, K.M. Rifat Faysal, Alex Kleinpeter, Miranda S.C. Wilson, Marina Vaysburd, Adam J. Fletcher, Mariia Novikova, Till Böcking, Eric O. Freed, Adolfo Saiardi, and Leo C. James

Supplementary Figure 1 related to Figure 1: (A) Analysis of CRISPR knockout clones. Chromatograms and alignments for sequencing of CRISPR/Cas9 knockout clones of IPMK. Indels were identified using the program outlined in Dehairs, J. *et al.*(Dehairs et al., 2016), as well as manual decoding. (B) PAGE and toluidine blue staining of standard cell extracts including standard controls to demonstrate migration of IP₅, IP₆, ATP and GTP. Standards were run in the presence of cell extract to allow equivalent migration. (C) Western blot for Gag expression levels in transfected IPMK KO clones. Gag precursor Pr55Gag (Pr55), p41 and mature capsid protein (p24) are indicated. Lower panel shows loading control COX-IV. (D) Graph showing pg RT per ng p24 in virions from viral supernatants obtained from transfection of IPMK KO clones. Error bars depict mean ± SD of three independent experiments with no statistical difference to WT. (E) Western blots showing overexpression of IPMK in CRISPR knockout clones. Clones were blotted for IPMK and lower panel shows Cox-IV loading control. (F) TiO2-PAGE and toluidine blue staining of cell extracts from IPMK KO clones stably transduced with empty vector control (EV) or a CMV-driven IPMK gene. Synthetic polyP was used for gel orientation. Gels show knockout clones are successfully reconstituted and IP $_6$ levels restored. (G) Western blot for Gag expression levels in IPMK KO parental clones and cells stably transduced with either EV or IPMK. Gag precursor Pr55Gag (Pr55), p41 and mature capsid protein (p24) are indicated. Lower panel shows loading control COX-IV.

Supplementary Figure 2 related to Figure 2: (A) Analysis of CRISPR knockout clones. Chromatograms and alignments for sequencing of CRISPR/Cas9 knockout clones of IPPK. Indels were identified using the program outlined in Dehairs, J. *et al.*(Dehairs et al., 2016), as well as manual decoding. (B) Comparison of total IP₅ and IP₆ incorporation in virions from IPPK KO clones shown in Figure 2C. (C) Western blot for Gag expression levels in transfected IPPK KO clones. Gag precursor Pr55Gag (pr55), p41 and mature capsid protein (p24) are indicated. Lower panel shows loading control COX-IV.

Supplementary Figure 3 related to Figure 4: Alignment of lentiviral gag sequences. Residues are highlighted by conservation, with the mature and immature charged ring residues boxed.

Supplementary Figure 4 related to Figure 5: (A) Western blot showing Gag expression levels of lysine mutants in producer cells. Gag precursor Pr55Gag (pr55), p41 and mature capsid protein (p24) are indicated. (B) Quantification of viral production from 293T, IPMK KO and IPPK KO cells as determined by RT levels in viral supernatants. Error bars depict mean ± SD of two independent experiments. Values are expressed as fold change from levels of RT produced in WT virus. (C) Western blot for Gag expression levels in cell lysates from WT and IPMK and IPPK KO cells during WT and K227I virus production. Gag precursor Pr55Gag (pr55), p41 and mature capsid protein (p24) are indicated. (D) Western blots of cell extracts showing cell extracts from stably transduced cells, showing depletion of TNPO3 or NUP153 by expression of short hairpin RNAs.

Supplementary Table 1 related to Figure 3: Data collection and refinement statistics. Crystallographic statistics for the HIV-1 hexamer structure complexed with IP5.

Supplementary Figure 1 related to Figure 1

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0

1_7 1_8 1_10 1_14 2_1 2_19

IPMK 1_8

IPMK 1_14

IPMK 2_19

IPMK GGCTGCGTGCCCCTCTCG-CATCAGGTGGCCGGGCACAT
2 19(1) GGCTGCGTGCCCCTCTCGGCATCAGGTGGCCGGGCACAT

2_19(2) GGCTGCGTGCCCCTCTCG---TCAGGTGGCCGGGCACAT

2_19(1) GGCTGCGTGCCCCTCTCGGCATCAGGTGGCCGGGCACAT
2 19(2) GGCTGCGTGCCCCTCTCG---TCAGGTGGCCGGGCACAT

Supplementary Figure 2 related to Figure 2

A

IPPK B2_2

 $\begin{tabular}{ll} IPFK & CGTGGTACCCCCAT-TCATTCTCGTCCATCTTCCCCTCT \\ 2_2(1) & CGTGGTACCCCCATTTCATTCTCGTCCATCTTCCCTCT \\ 2_2(2) & CGTGGTACCCCCATCTT-TTCTCGTCCATCTTCCCCTCT \end{tabular}$ 2_2(1) CGTGGTACCCCCATTTCATTCTCGTCCATCTTCCCCTCT 2_2(2) CGTGGTACCCCCATCT--TTCTCGTCCATCTTCCCCTCT

IPPK GCCCTCTCCGTGGTACCCCCA-TTCATTCTCGTCCATCT 2_7 GCCCTCTCCGTGGTACCCCCATTTCATTCTCGTCCATCT

IPPK GCTCTTATTGCCCTCTCCGTGGTACCCCCATTCATTCTCGTCC 1_8 GCTCTTA-----------------------TTCATTCTCGTCC

IPPK B1_13

IPPK B2_3

IPPK B3_7

IPPK CGCGTGGGCCACCACCAGG-CTCTTATTGCCCTCTCCGT $3\begin{array}{lll} 3\!-\!7\,(1) \qquad \ \ \texttt{CGGTGGCACCACCACCGTCTTATTGCCCTTCCGT} \\ 3\!-\!7\,(2) \qquad \ \ \texttt{CGGTGGCACCACGGGTCTCTTATTGCCCTTCCGT} \end{array}$ 3_7(2) CGCGTGGGCCACCACCAGGTCTCTTATTGCCCTCTCCGT

B C

Supplementary Figure 3 related to Figure 4

Supplementary Figure 4 related to Figure 5

Supplementary Table 1 related to Figure 3: Data collection and refinement statistics

*Values in parentheses are for highest-resolution shell.