Molecular Determinants for Recognition of Divergent SAMHD1 Proteins by the Lentiviral Accessory Protein Vpx

Graphical Abstract

Highlights
- The crystal structure of SIVmnd-2 Vpx complexed with SAMHD1 and DCAF1
- Comparisons to SIVsmm, Vpx reveal determinants for N- versus C-terminal SAMHD1 binding
- Vpx and Vpr use conserved determinants to bind to DCAF1

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In Brief
Vpx of HIV-2 and SIV target the restriction factor SAMHD1 for degradation using the host ubiquitin ligase substrate receptor DCAF1. Schwefel et al. present the ternary complex of Vpxmnd-2 with DCAF1 and the SAMHD1 N terminus and reveal how Vpx subverts host cell defenses by employing differing strategies to recruit SAMHD1 for degradation.

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Molecular Determinants for Recognition of Divergent SAMHD1 Proteins by the Lentiviral Accessory Protein Vpx

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INTRODUCTION

Host cell restriction factors are the first line of cellular defense against retrovirus infection. Successful viral evasion of these adaptive hurdles is essential for productive infection of new hosts and has occurred during multiple cross-species transmission events such as those that led to the emergence of pandemic human immunodeficiency virus-1 (HIV-1) (Hatzioannou and Bieniasz, 2011; Hatzioannou et al., 2014; Sharp and Hahn, 2011). Restriction factors are often interferon induced and inhibit distinct stages of the viral replication cycle (Towers and Noursadeghi, 2014; Wolf and Goff, 2008). Important examples are the TRIM5α, APOBEC3 family, and Tetherin proteins, which interfere with retroviral uncoating, reverse transcription, and budding processes, respectively (Neil et al., 2008; Sheehy et al., 2002; Stremlau et al., 2004). The sterile alpha motif domain and histidine-aspartate domain-containing protein 1 (SAMHD1) was first identified as a disease gene associated with the rare infantile encephalopathy Aicardi-Goutières syndrome (Rice et al., 2009). More recently, SAMHD1 was shown to be a factor that restricts HIV-1 replication in non-dividing myeloid-lineage cells and resting T cells (Baldauf et al., 2012; Berger et al., 2011; Descours et al., 2012; Hrecka et al., 2011; Laguette et al., 2011). SAMHD1 is a dGTP/GTP-activated deoxynucleotide (dNTP) triphosphohydrolase (Amie et al., 2013; Goldstone et al., 2011; Ji et al., 2013; Miazzi et al., 2014; Powell et al., 2011; Zhu et al., 2013) involved in balancing cellular dNTP pools (Franzolin et al., 2013) and regulated by Cyclin A2/CDK1-dependent phosphorylation (Cribier et al., 2013; Kretschmer et al., 2015; Pauls et al., 2014; White et al., 2013). It is proposed that SAMHD1 restriction results from this triphosphohydrolase activity by reducing the cellular dNTP concentration to a level insufficient for the viral reverse transcriptase to function (Kim et al., 2012; Lahouassa et al., 2012; Rehwinkel et al., 2013; St Gelais et al., 2012). An alternative mechanism of SAMHD1 HIV-1 restriction requiring a putative nuclease activity as also been reported, but the nature of the polynucleotide substrate is disputed (Beloglazova et al., 2013; Ryoo et al., 2014; Tüngerl et al., 2013).

A hallmark of most retrovirus restriction factors is the existence of viral antagonists in the form of accessory proteins (Malim and Bieniasz, 2012; Strebel, 2013). A common mechanism of action of accessory proteins is the subversion of host cell protein degradation pathways (Strebel, 2013). In particular, the host cell’s Cullin-RING-type E3 ubiquitin ligases are often engaged by viral accessory proteins to induce restriction factor poly-ubiquitylation to direct proteasomal degradation (Barry and Früh, 2006). Cullin-RING ubiquitin ligases consist of a central Cullin scaffold protein, a catalytic RING subunit, and varying substrate receptors (Zimmerman et al., 2010). Owing to their modular architecture, Cullin-RING ligases allow for specific placement of a large number of substrates in the ubiquitylation zone of the catalytic subunit for efficient poly-ubiquitylation (Fischer et al., 2011; Zimmerman et al., 2010). Simian immunodeficiency virus (SIV) and HIV accessory proteins exploit these...
characteristics by modifying the specificity of Cullin-RING sub-
strate receptors (e.g., HIV-1 Vif and Vpu redirect cullin-1 and
cullin-5 receptor specificity to induce APOBEC3 and Tetherin re-
striction factor downregulation, respectively) (Guo et al., 2014;
Mitchell et al., 2009). Vpr and Vpx from HIV and SIVs target the
cullin-4 ligase substrate receptor DDB1- and CUL4-associated
factor 1 (DCAF1, also known as VprBP) from their host (Berga-
maschi et al., 2009; Hrecka et al., 2007; Le Rouzic et al., 2007;
Srivastava et al., 2008).

Restriction factors and accessory proteins are engaged in an
evolutionary “molecular arms race” consisting of multiple rounds of host adaptation, virus counteraction, and host re-
adaptation, resulting in accumulation of amino acid changes in
restriction factor-accessory protein interaction interfaces
(Daugherty and Malik, 2012). In the lentiviral Vpx/Vpr accessory
proteins, analyses of positively selected residues and subse-
quent functional studies have demonstrated the occurrence of
several significant specificity changes during adaptation to pri-
mate hosts (Laguet et al., 2012; Lim et al., 2012). It is proposed
that originally a subset of Vpr proteins acquired the capability to
induce cullin-4/DCAF1-dependent proteasomal degradation of
SAMHD1. These SAMHD1-degrading Vpr proteins are found in the
SIVsyk (SIV that infects Sykes’ monkey), SIVdeb (De Brazza’s
monkey), and SIVagm (African green monkey) lineages. Later in its
evolutionary history, a Vpr gene duplication or recombination
event gave rise to Vpx, which completely took over the anti-
SAMHD1 functionality and produced viruses that contain both
Vpx and Vpr genes (Lim et al., 2012). In these adaptation events,
the mode of SAMHD1 binding seems to have toggled between
recognition sequences at the SAMHD1 N terminus and others
located at the C terminus. Vpx from HIV-2, SIVsrm (sooty man-
bagaiey) and SIVmac (macaque) all target the SAMHD1 C terminus,
whereas SIVmnd-2 (mandrill) and SIVrcm (red-capped mangabey)
Vpx induce SAMHD1 degradation dependent on the N-terminal
110 residues. The SAMHD1-degrading Vpr proteins from SIVsyk,
SIVdeb, and SIVagm have varying specificities for the SAMHD1 N
or C terminus or even target both (Fregoso et al., 2013; Spragg
and Emerman, 2013).

Here, we report the crystal structure of the ternary complex of
SIVmnd-2 Vpx bound to the N-terminal region and SAM domain of
mandrill SAMHD1 (SAMHD1mnd-NtD) and the WD40 domain of the
cullin-4 substrate receptor DCAF1 (DCAF1-CtD). This structure
reveals unexpected complexity in Vpx-mediated target-
ning of SAMHD1. Comparison to the recently determined
structure of a SIVsrm, Vpx/SAMHD1-CtD/DCAF1-CtD (Schwefel
et al., 2014) complex provides the molecular details of the struc-
tural determinants for N-terminal versus C-terminal SAMHD1
specificity and a plausible scenario for the events occurring dur-
ing evolutionary change.

RESULTS

Identifying the Mandrill SAMHD1 Degron
To locate the Vpx-DCAF1 recognition sequence within the
SAMHD1mnd N terminus, a cell-based EGFP-degron reporter
degradation assay was employed (Schwefel et al., 2014). Degron fusion proteins (Figure 1A) were constructed comprising
two copies of EGFP with a nuclear localization signal (NLS)
fused to either the N-terminal 114 residues of SAMHD1mnd
(SAMHD1mnd-NtD) or a control C-terminal region of human
SAMHD1, residues 600–626, (SAMHD1CtD-CtD), which is tar-
ged for degradation by Vpx from SIVsrm (Schwefel et al.,
2014). Stable cell lines expressing each reporter construct
were produced and expression levels of degron fusion proteins

Figure 1. The Mandrill SAMHD1 Degron Comprises the N-Terminal and SAM Domains
(A) Schematic of the human and mandrill SAMHD1 and the tandem NLS-EGFP degron fusion protein construct employed in fluorescent degron assays. The C- and N-terminal degron regions in SAMHD1 are highlighted in green and magenta, respectively.
(B) Stable cell lines expressing either the mandrill (mnd) SAMHD1 1–114 degron (left) or human (hs) SAMHD1 600–626 degron (right) were transduced with increasing amounts of SIVmnd-2 Vpx; see also Figure S1. Quantification of degron reporter expression in stable cells lines expressing constructs containing the indicated N-terminal regions of SAMHD1mnd after transduction with increasing amounts of SIVmnd-2 Vpx; see also Figure S1.
were confirmed by western blot (Figure S1). Cells were then transduced with increasing amounts of a bicistronic IRES vector expressing Vpx from SIVmnd-2 or SIVsmm and YFP, and 48 hr later, the level of degron reporter (EGFP) and Vpx (YFP) expression was measured by flow cytometry. These data show that SIVmnd-2 Vpx induces degradation of the SAMHD1mnd-NtD reporter construct but not EGFP-SAMHD1hs-CtD (Figure 1B, left panel). In contrast, SIVsmm Vpx induces degradation of the SAMHD1hs-CtD reporter but not SAMHD1mnd-NtD (Figure 1B, right panel). To further delineate the SIVmnd-2 SAMHD1 binding determinants, reporter constructs containing residues 1–37 and 37–114 of SAMHD1mnd were prepared. Residues 1–37 comprise an N-terminal disordered sequence containing a NLS and 37–114 constitutes the SAM domain (cp. PDB: 2E8O). However, neither fragment alone was sufficient to induce degradation of the reporter construct (Figure 1C), demonstrating that both the N-terminal NLS region and SAM domain are required for SIVmnd-2 Vpx/DCAF1-mediated degradation.

**Structure of the SIVmnd-2 Vpx/DCAF1-CtD/SAMHD1mnd-NtD Complex**

(A) Schematic of components of the protein complex. (B) Analytical gel filtration of the in vitro-assembled ternary protein complex consisting of SIVmnd-2 Vpx, DCAF1-CtD and SAMHD1mnd-NtD (Peak-1). The inset shows SDS-PAGE analysis of the indicated peak fractions. (C) Structure of the ternary complex. The backbone for each protein is shown in cartoon representation, SIVmnd-2 Vpx (blue), DCAF1-CtD (gray), and SAMHD1mnd-NtD (magenta). See also Figure S2. The solvent-accessible surface is also shown for DCAF1-CtD, and a zinc ion co-ordinated by Vpx is displayed as a gray sphere, see also Figure S3. For SIVmnd-2 Vpx and SAMHD1mnd-NtD, residue numbers at chain termini are indicated and secondary structure elements labeled.
The DCAF1 Binding Mode Is Conserved in Vpx Proteins

The extensive Vpx-DCAF1 interface comprises four contact regions with a total surface area of 1,600 Å² (Figure 3; Table S1A). The first region comprises the C-terminal section of α2 together with first half of α3 of Vpx. On one face, apolar amino acid side chains from both α2 and α3 pack against a cluster of hydrophobic amino acids located in the “tall” loop that connects β2 to β3 in WD40 repeat 7 on the upper surface of DCAF1-CtD (Figure 3B, upper panel). On the other face, a network of hydrogen bonds and salt bridges connects Vpx residues Y62, Y65, R66, and K73 at the N terminus of α3 to acidic side chains E1191, D1092, and E1193 in the “acidic” loop of DCAF1 that intersperses WD40 repeats 7 and 1 (acidic loop, Figure 3B, lower panel). A second region of interaction involves the carboxy terminus of Vpx α3, which packs into a radial groove on the top side of DCAF1 lined by amino acids from WD40 repeats 1, 2, 4, 6, and 7. Interactions here include both the packing of hydrophobic residues as well as specific hydrogen bonding between the side-chains of Vpx Y76 and Q81 and the DCAF1 protein backbone (Figure 3C). The third binding interface involves residues at the extreme amino terminal of the Vpx N-terminal extended arm that inserts into a cavity between WD40 repeats 1 and 2 on the underside of DCAF1. Here, Vpx A2, E3 and E7 make main chain and side chain hydrogen-bonds to DCAF1 residues S1102, R1106, Y1131, and S1168 while the interposing Vpx residues A5 and P6 project into the hydrophobic cavity (Figure 3D). The remaining site of interaction involves contacts between residues in the WD40 repeat 2 of DCAF1 and residues in both Vpx α1 and α3. These include hydrophobic interactions between DCAF1 T1155 and W1156 and apolar side chains displayed on the underside of Vpx α1 and α3 and Vpx H72 on α3 that makes a hydrogen bond with mainchain carbonyl of N1135 of DCAF1 (Figure 3E).

Structural alignment of the DCAF1-CtD and Vpx molecules in the SIVmnd-2 and SIVsmm complexes (Schwefel et al., 2014) reveals that in both instances they adopt very similar conformations. The root-mean-square deviations (RMSDs) are 0.65 Å (305 aligned residues) for the DCAF1-CtD molecules and 1.24 Å for Vpx (80 aligned residues) (Figure S4A). Moreover, detailed comparison of SIVmnd-2 and SIVsmm Vpx amino acid side chains that contact DCAF1 shows that nearly all are at least type-conserved (Figures S4B–S4E). A notable exception is T47 in α2 that is exchanged to arginine in SIVsmm Vpx and is concomitant with a small movement of α2 and differences in conformation of the interacting DCAF1 tall loop between the two complexes (Figure S4B, upper panel). However, residues on the upper face of α2 also interact with the SAMHD1mnd SAM domain (see below) so the cause of the α2 shift is unclear. Therefore, given that the mode of the N-terminal arm - α2 - α3 Vpx association with DCAF1 is a common feature of SIVmnd-2 and SIVsmm complexes and that nearly all the DCAF1-interacting residues are type-conserved in HIV-2/SIV Vpx proteins (Figure S5), it is likely that the same principles of DCAF1 binding apply to both the SIVmnd-2 and SIVsmm Vpx lineages.

A Combined SIVmnd-2 Vpx/SAMHD1mnd(1-22)/DCAF1-CtD Ternary Interface

In the complex, the N-terminal NLS region (residues 1–22) of SAMHD1mnd comprises an extended chain that binds to the
edge of DCAF1 in an interface that buries 500 Å² of solvent accessible surface (Figure 4A; Table S1B). Here, residues D5, D7, and Q8 make extensive sidechain and mainchain hydrogen bonds to residues in the interspersing loops of DCAF1 WD40 repeat-1. In addition, SAMHD1 residues R12 and R14, which are part of the SAMHD1 NLS (Brandariz-Núñez et al., 2012; Hofmann et al., 2012), make salt bridges to E1091 and D1092 in the DCAF1 acidic loop. The ternary interaction is completed by a 750 Å² SAMHD1mnd-SIVmnd-2 Vpx interface that includes multiple interactions with Vpx residues from the N-terminal arm, α1, and the α2-α3 loop (Figure 4B; Table S1C). One feature of this interface is a “trapped” region where the Vpx N-terminal arm wraps over SAMHD1 packing it against DCAF1. Within the trapped region, the main chains of SAMHD1 and Vpx interact through hydrogen bonding between E10, G13, and E14 of Vpx and Q2 and S10 of SAMHD1. In addition, the main chain of Vpx G13, and E14 also make interactions with the SAMHD1 D7 and S10 side chains. C-terminal to the trapped region there are further interactions where Vpx V15, L17, and W20 create a hydrophobic pocket at the N terminus of α1 that SAMHD1 P9 inserts into. Additionally, SAMHD1 NLS residue R12 makes a hydrogen bond to Vpx Y65, SAMHD1 F15 stacks with Vpx R58, and SAMHD1 R20 makes an electrostatic interaction with Vpx D54 (Figure 4B).

The functional importance of the Vpx trapping of the SAMHD1 N-terminal region was revealed in an EGFP-degron fusion protein degradation assay. Figure 4C shows that SIVmnd-2 Vpx induces efficient degradation of constructs containing SAMHD1 amino acids 1–114 and 5–114. In contrast, EGFP-SAMHD1 (10–114), which lacks D7 and Q8 that make hydrogen bonds to...
Vpx and DCAF1 within the trapped region, is not degraded by Vpx indicating the requirement of the trapping interaction for degron activity.

The SIVmnd-2 Vpx/ SAMHD1mnd SAM Domain Interface

In addition to the ternary interface that “traps” the SAMHD1mnd N terminus within the complex a further DCAF1-independent interaction between the SAM domain of SAMHD1mnd-NtD and SIVmnd-2 Vpx is also observed. This interface has a buried surface area of 580 Å² and comprises residues that project upward from Vpx α2 that contact SAMHD1 residues N-terminal to the SAM domain helix 1 as well as those on the underside of the SAM domain in helices 1 and 2 (Figure 5A). The center of the interface comprises a hydrophobic core created by Vpx residues P38, L41, F42, W45, and V49 and SAMHD1 residues P34, V36, L38, and F52 (Figure 5A, upper panel). In addition to this hydrophobic patch, there are electrostatic interactions between SAMHD1 R55 and Vpx D54 (Figure 5A, upper panel), SAMHD1 R69 and Vpx E50 (Figure 5A, lower panel) as well as hydrogen bonding from Vpx E39 to the backbone of the first turn of SAM domain helix 1 (Figure 5B, lower panel). The contribution of residues in the SAM domain-Vpx interface to degron function was assessed by introduction of alanine mutations at SAMHD1 residues R55 and R69 in the EGFP-fusion reporter. Both residues make extensive electrostatic interactions at the interface, and the alanine substitutions result in a loss of reporter degradation (Figure 5B). R69 is also among a group of residues in the N-terminal region of SAMHD1 that have undergone positive evolutionary selection (Lim et al., 2012) indicative of an interaction hotspot between restriction factor and viral antagonist, further underlining the importance of Vpx-SAM domain contacts for induction of SAMHD1 proteasomal degradation.

Comparison of the sequences of human and mandrill SAMHD1-NtD regions reveals 13 amino-acid changes, nine in the SAM domain and four in the NLS region. Of these, only two make side chain contacts in the mandrill ternary complex, F15 in the NLS region and F52 in the SAM domain. In human SAMHD1, these residues are replaced by non-conservative cysteine and serine substitutions (Figure S6). In the mandrill complex, both phenylalanine residues make interactions, F15 stacks against the side chain of R58 in the Vpx α2-α3 loop (Figure 4B, left panel), and F52 is central to the hydrophobic SAM

Figure 4. The Combined SIVmnd-2 Vpx/DCAF1-CtD/SAMHD1mnd-NtD Ternary Interface

(A) Interface of DCAF1-CtD and SAMHD1mnd NLS region. β strands from WD40 repeat-1 and -2 of DCAF1-CtD are shown in gray cartoon representation. The N-terminal NLS region of SAMHD1mnd is shown in magenta. Residues making contacts at the interface are shown in stick representation with hydrogen bonding and salt bridge interactions displayed as dashed lines.

(B) The Vpx-SAMHD1mnd NLS region interface. The ternary complex is displayed (central) in the same orientation and representation as Figure 2. Details of SAMHD1-Vpx interactions within the boxed region are shown left and right. Residues in SAMHD1 (magenta) and Vpx (blue) making contacts at the interface are shown in stick representation with hydrogen bonding and salt bridge interactions displayed as dashed lines.

(C) Quantification of reporter expression in stable cell lines containing degron constructs with the indicated N-terminal truncations of SAMHD1mnd. Cells were transduced with increasing titers of particles expressing SIVmnd-2 Vpx together with YFP and degron reporter expression (EGFP) measured by flow cytometry.
SAMHD1 and their contribution tested in the degron assay (Figure 5C). These data show the F15C mutation renders SAMHD1\textsubscript{mnd} resistant to SIV\textsubscript{mnd-2} Vpx-induced proteasomal degradation but that surprisingly the F25S mutation only slightly reduces degradation of the degron reporter protein. Nevertheless, introduction of the reverse mutations C15F and S52F in human SAMHD1 has previously been shown to render human SAMHD1 sensitive to SIV\textsubscript{mnd-2} Vpx-mediated degradation (Wei et al., 2014). Therefore, taken together, these data support the idea that amino acid changes in primate SAM domain and NLS regions determine the lineage-specific formation of a Vpx-SAMHD1 interaction interface.

### Specificity of Vpx-SAMHD1-NtD and -CtD Interactions

Despite similar overall structure and an identical mode of DCAF1 interaction (Figure S4), the SIV\textsubscript{mnd-2} Vpx/DCAF1 assembly recognizes the SAMHD1\textsubscript{mnd}-NtD, while the HIV-2/SIV\textsubscript{mac}/SIV\textsubscript{smm}\textsuperscript{+} type Vpx/DCAF1 are specific for the SAMHD1 C terminus. Comparison of the SIV\textsubscript{mnd-2} system presented here with the previously determined SIV\textsubscript{smm}/DCAF1/SAMHD1-CtD assembly (Schwefel et al., 2014) reveals the structural determinants for these fundamentally different specificities. In both complexes, a Vpx N-terminal arm and residues at the N terminus of \(\alpha1\) make contacts with either the amino-terminal region in SAMHD1\textsubscript{mnd} or the CtD region of SAMHD1\textsubscript{hs}. The location of this contact on the DCAF1 disk is similar in both combined ternary interfaces (Figure 6A). However, the SAMHD1\textsubscript{mnd} N terminus makes additional contacts and occupies a larger surface area on Vpx and DCAF1, 750 Å\(^2\) and 500 Å\(^2\) versus 710 Å\(^2\) and 210 Å\(^2\) for the SAMHD1\textsubscript{hs} C terminus. In addition, in the mandrill complex, the Vpx arm folds over the first 20 residues of SAMHD1\textsubscript{mnd} trapping it against DCAF1, while the human SAMHD1 C-terminal peptide associates rather peripherally to the DCAF1/Vpx interface (Figure 6A).

In this context, it is important to note that Vpx amino acid residues at the N terminus of \(\alpha1\) that contribute to SAMHD1 binding differ significantly between SIV\textsubscript{smm} and SIV\textsubscript{mnd-2} type Vpx proteins constituting a variable region (VR1) (Figure S5). In the SIV\textsubscript{smm} complex, VR1 side chains of the conserved GEET-motif form salt bridges with basic residues in SAMHD1-CtD (Schwefel et al., 2014). However, in SIV\textsubscript{mnd-2} Vpx, VR1 comprises a proline (P\textsubscript{g}\textsubscript{mnd-2}) followed by two glycine residues interspersed with a small hydrophobic amino acid and then a conserved acidic and hydrophobic residue (\(\Phi\text{P}[\text{GAG[D/E]}V/A]\textsuperscript{15}\)). This configuration specifically accommodates the SAMHD1\textsubscript{mnd}-NtD and sandwiches it between Vpx and the DCAF1 disk. The importance of this VR1 interaction for Vpx-mediated proteasomal degradation of SAMHD1\textsubscript{mnd} is demonstrated in EGFP-degron fusion protein assays where substitution of valine by bulky tryptophan at position 15 of SIV\textsubscript{mnd-2} Vpx substantially reduces degron reporter protein degradation (Figure 6B). In addition, analysis of the clustering of VR1 sequences (Figure S5) shows that SIV\textsubscript{com} and SIV\textsubscript{vrv1} Vpx that are SAMHD1-NtD specific also contain a SIV\textsubscript{mnd-2} type VR1, supporting the idea that VR1 is a major determinant for discrimination between SAMHD1 N-terminal and C-terminal target sequences.

As well as VR1, which confers NtD and CtD degron specificity to Vpx, a second region, VR2, in the \(\alpha2\)-\(\alpha3\) loop and N terminus of \(\alpha3\) is conserved among Vpx but not in Vpr (Figure S5). Here, a domain-Vpx interaction (Figure 5A). In order to assess their role in the specificity of the Vpx-SAMHD1 interaction, F15 and F25 were mutated to cysteine and serine that are found in human

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**Figure 5. SAM Domain-SIV\textsubscript{mnd-2} Vpx Interactions**

(A) The SAMHD1\textsubscript{mnd} SAM domain-SIV\textsubscript{mnd-2} Vpx interface. The SAMHD1\textsubscript{mnd} SAM domain (magenta) and SIV\textsubscript{mnd-2} Vpx (blue) are shown in cartoon representation. Residues making interactions are shown in stick representation with hydrogen bonding and salt bridges displayed as dashed lines.

(B and C) Quantification of reporter expression in cells stably expressing SAMHD1\textsubscript{mnd} (1–114) degron reporter constructs with (B) SAM domain point mutations and (C) mandrill to human SAMHD1 amino acid substitutions. Cells were transduced with increasing titers of particles expressing SIV\textsubscript{mnd-2} Vpx, and the level of transduction (YFP) and degron reporter (EGFP) expression was measured by flow cytometry.
The current properties of accessory proteins Vpr and Vpx likely reflect a lengthy history of conflict between lentiviruses and their hosts. Eight different lineages of exogenous primate lentivirus reflect a lengthy history of conflict between lentiviruses and their hosts. Eight different lineages of exogenous primate lentivirus including HIV-1 can be identified. All encode a Vpr that induces host G2/M cell-cycle arrest (Berger et al., 2015), probably by targeting a still poorly defined host cell protein for degradation through the pro-teasomal pathway or through interaction with proteins of the DNA damage pathway (Laguette et al., 2014). Vpr from two viral clades, typified by SIVagm and SIVsyk, is also able to target SAMHD1 for degradation, facilitating infection of some types of non-dividing cells. In two further clades, both Vpr and Vpx are present with Vpr retaining only cell-cycle arrest activity and Vpx only inducing SAMHD1 degradation.

Phylogenetic reconstructions suggest that the Vprs from SIVagm, and SIVsyk, acquired anti-SAMHD1 activity prior to the molecular events leading to the generation of Vpx (Lim et al., 2012). Moreover, it appears that once Vpr acquires anti-SAMHD1 activity, it does not lose it. This would imply that viruses containing both Vpr and Vpx arose by the recombination of two viruses; the first encoding an ancestral Vpr with the dual ability to cause cell-cycle arrest and degrade SAMHD1 and the second with a Vpr that had not acquired the capacity to induce SAMHD1 degradation. This allowed the ancestral, dual function Vpr to convert to Vpx and maintain SAMHD1 degradation activity but lose the cell-cycle arrest function provided by the second Vpr. Evaluation of our crystal structures is most informative in understanding these evolutionary changes and specifically the interaction between (i) Vpr/x and DCAF1 and (ii) between the two classes of Vpx and their SAMHD1 targets.

Amino acid residues that bind to DCAF1 and those that coordinate the bound zinc are highly conserved in all lentiviral Vpx molecules (Figure S5), suggesting that the overall structure of the protein as well as mode of DCAF1 binding have been conserved during Vpr/Vpx evolution. This implies that the current properties of accessory proteins Vpr and Vpx likely reflect a lengthy history of conflict between lentiviruses and their hosts.

Figure 6. Comparison to the SIVsmm, Vpx/ DCAF1-CID/SAMHD1ntd-Cid Structure
(A) Superposition of the SIVmnd-2 Vpx/DCAF1-CID/ SAMHD1mnd-NtD and SIVsmm Vpx/DCAF1-CID/ SAMHD1mm-Cid ternary complexes. Cartoons are colored as Figure 2 with the addition of SIVsmm Vpx (orange) and SAMHD1mm-Cid (green). For clarity, only DCAF1-Cid from the mandrill complex is shown. (B and C) Quantification of reporter expression in cells stably expressing the SAMHD1mnd-1-114 (1–114) degron reporter construct after transduction with increasing titer of particles expressing SIVmnd-2 Vpx with mutations in (B) the N-terminal arm (Y19W) and in (C) the di-tyrosine motif (Y62A/Y65A). The level of Vpx transduction (YFP) and degron reporter (EGFP) expression was measured by flow cytometry. (D) Detailed view of the conserved VR2 Vpx di-tyrosine motif interaction with the DCAF1-Cid acidic loop and basic motifs of the SAMHD1mnd-NtD (left) and SAMHD1mm-Cid (right) degrons; see also Figure S5. Residues that contribute to the ternary interface are shown in stick representation with hydrogen bonding and salt bridge interactions displayed as dashed lines; see also Figure S6.
HIV-1 Vpr/DCAF1 association is likely to be similar to the one observed in the SIVmnd-2 and SIVsmm (Schwefel et al., 2014) Vpx/DCAF1 crystal structures (Figure S4). Therefore, these structures might provide suitable starting points for rational design of compounds interfering with HIV-1 Vpr function even in the absence of definitive information concerning its cellular target. A notable exception is Vpr from SIVdies that induces SAMHD1 degradation apparently without binding to DCAF1 (Berger et al., 2015), and it will be interesting to determine whether it utilizes an alternative pathway or different receptor protein to target SAMHD1.

Vpx from SIVsmm and SIVmnd-2 have been shown to interact with the C- and N-terminal regions of SAMHD1, respectively (Fregoso et al., 2013). Comparison of Vpx/SAMHD1/DCAF1 ternary complexes reveals that this discrimination of SAMHD1 N- and C-terminal degrons by the two classes of Vpx is determined both by polymorphisms in the SAMHD1 amino-terminal region (Figure S6) and by specific sequences in the VR1 region of Vpx (Figures 6 and S5). By contrast, in both classes of Vpx the VR2 region contains a conserved di-tyrosine pair that interacts with the DCAF1 acidic loop to provide a hydrophobic platform for N- and C-terminal SAMHD1 degrons. In this way, basic residues from both degron types are positioned for electrostatic interactions with the DCAF1 acidic loop. Interestingly, only the second tyrosine is present in SIVagm Vpr proteins, which are incapable of inducing human SAMHD1 degradation, while the position corresponding to the first tyrosine is occupied by a glutamate residue (Figure S5). Gain of the bi-tyrosine motif might therefore have been a critical step in the development of Vpx allowing multiple modes of binding SAMHD1 while at the same time resulting in loss of the Vpr capacity to cause cell-cycle arrest.

A third highly variable region (VR3) in Vpr/Vpx proteins is directly C-terminal to the fourth conserved zinc-coordinating residue (Figure S5) that in both SIVsmm and SIVmnd-2 type Vpx contains proline-rich stretches of variable length. In the corresponding SAMHD1/Vpx/DCAF1 crystal structures, VR3 is either disordered (this study) or involved in crystal contacts (Schwefel et al., 2014), strongly suggesting that the region is not involved in SAMHD1 degron recognition. In agreement, SIVmac Vpx lacking the poly-proline tail is still able to assemble into a ternary SAMHD1/Vpx/DCAF1 complex and to induce cullin-4-dependent SAMHD1 poly-ubiquitination (Ahn et al., 2012) and degradation (Belshan et al., 2012). Functional studies suggest involvement of VR3 in nuclear localization of Vpx and Vpr (Pancio et al., 2000; Zhou et al., 1998) with consequences for viral replication (Belshan et al., 2012), but notably, mutations in this region also influence protein expression levels significantly (Miyake et al., 2014).

Predictions regarding DCAF1/Vpr-mediated recruitment of target protein(s) that result in G2/M cell-cycle arrest remain speculative. In HIV-1, VR2 differs significantly from other Vpr and Vpx proteins, suggesting that in contrast to the Vpx di-tyrosine/DCAF1 acidic loop/SAMHD1 interaction, other principles may apply to HIV-1 Vpr target recognition. By contrast, VR1 of HIV-1 Vpr contains a 12REPF[Y/W/D/N]16 motif that is also conserved in SIVagm Vpr proteins (Figure S5), suggesting similarities in VR1-mediated SIVagm and HIV-1 Vpr target recruitment. Therefore, considering recent evidence for HIV-1 Vpr/DCAF1 association with components of the SLX4/MUS81/EME1 DNA-repair complex (Berger et al., 2015; Laguette et al., 2014), structural analysis of SIVagm Vpr/DCAF1 together with the cognate SAMHD1agm will be of interest to further delineate the degron and/or target sequence(s) recognized by HIV-1 Vpr.

**EXPERIMENTAL PROCEDURES**

**Degron Assay**

Degron reporter constructs were generated by replacing the human SAMHD1-Ctd degron sequence in pCMs28-RL-EGFP-SAMHD1-Ctd with N-terminal sequences from SAMHD1mnd. Point mutations were created by site-directed mutagenesis. Virus-like particles (VLPs) were generated by co-transfecting 293T cells with pSVSVG, pKB4 and pCMs28-RL-EGFP-SAMHD1. Stable cell lines were produced by transduction of Mus dunni cells followed by puromycin selection. Expression of degron constructs was assessed using western blotting with anti-EGFP antibodies.

**Protein Expression and Purification**

The nucleotide sequences coding for SIVmnd-2 Vpx isolate 5440 and amino acid residues 1–114 of SAMHD1mnd were inserted into pET-49b and pET-52s (Merck Millipore) expression plasmids respectively to generate N-terminal GST-tagged and Strep-II-tagged fusion proteins. SAMHD1 (1–114) was expressed in E. coli strain Rosetta 2 and purified using Strep-Tactin affinity followed by size-exclusion chromatography. SIVmnd-2 Vpx was captured from the bacterial lysate onto Glutathione Sepharose (GSH-Sepharose) (GE Healthcare) prior to an assembly reaction (see below). His-tagged DCAF1, residues 1,058–1,396 (DCAF1-Ctd), was expressed in insect cells and purified using Ni-NTA Sepharose and size-exclusion chromatography.

For assembly, the GST-SIVmnd-2 Vpx bound beads were resuspended and incubated with 1 mg of DCAF1-Ctd, an equimolar amount of SAMHD1mnd-NtD, and 1 mg of HRV-3C protease (GE Healthcare) overnight at 4°C. After removal of the beads by centrifugation, the eluted ternary complex was further purified by size-exclusion chromatography. Details of protein expression constructs, protein production, and storage are provided in Supplementary Experimental Procedures.

**Cristallization and Structure Solution**

Cryostals of the SIVmnd-2 Vpx/SAMHD1mnd-NtD/DCAF1Ctd complex were grown using the hanging drop vapor diffusion at 18°C in 2 µl droplets comprising 1 µl complex (6.34 mg/ml) and 1 µl of reservoir solution (0.16 M Trisodium Citrate-HCl [pH 5.2] and 4% PEG 6000). X-ray diffraction data were collected on beamline I04 at the Diamond Light Source, UK at a wavelength of 0.9796 Å. The structure was solved by molecular replacement using the previously determined DCAF1-Ctd structure, and a homology model constructed with the previously determined SIVmnd-2 Vpx as template (PDB: 4CC9). Details of crystallization, data collection, processing, and structure solution are provided in Supplementary Experimental Procedures.

**Multiple Sequence Alignment**

Amino acid sequences were aligned using the ClustalW server and adjusted manually. NCBI accession numbers for Vpr, Vpx, and SAMHD1 are provided in Supplementary Experimental Procedures.

**ACCESSION NUMBERS**

The coordinates and structure factors of the SIVmnd-2/Vpx/SAMHD1mnd-NtD/DCAF1-Ctd ternary complex have been deposited in the Protein Data Bank (PDB: 5a21).
SUPPLEMENTAL INFORMATION

Supplemental Information includes six figures, three table, and Supplemental Experimental Procedures can be found with this article online at http://dx.doi.org/10.1016/j.chom.2015.03.004.

AUTHOR CONTRIBUTIONS

D.S., V.C.B., E.C., and P.A.W. performed experiments. D.S., V.C.B., E.C., P.A.W., J.P.S., K.N.B., and I.A.T. contributed to experimental design, data analysis, and manuscript writing. The authors declare no competing financial interests.

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Supplemental Information

Molecular Determinants for Recognition of Divergent SAMHD1 Proteins by the Lentiviral Accessory Protein Vpx

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Molecular determinants for recognition of divergent SAMHD1 proteins by the lentiviral accessory protein Vpx.

INVENTORY OF SUPPLEMENTAL INFORMATION

ITEM 1) Figure S1. Western blot analysis of expression and primary FACS data for SAMHD1 degron assay. Panel A of this item shows the level of expression of each SAMHD1 degron used in the study. Panel B shows the primary FACS data from individual degron assays. Both A and B are associated with Figures 1, 4, 5 and 6 in the main article.

ITEM 2) Figure S2. Experimental electron density for each component of the SIV\textsubscript{mnd-2} Vpx/SAMHD\textsubscript{mnd-NtD/DCAF1-CtD} ternary complex. This item shows the quality of the maps used to build the structure presented in the paper and is associated with Table 1 and Figures 2-6 in the main article.

ITEM 3) Figure S3. Zinc ion co-ordination of Vpx SIV\textsubscript{mnd-2} and Vpx SIV\textsubscript{smm}. This item shows the details of the co-ordinated zinc ions bound in the Vpx SIV\textsubscript{mnd-2} and Vpx SIV\textsubscript{smm} structures. It is associated with Figure 2 in the main article.

ITEM 4) Figure S4. Superposition of Vpx SIV\textsubscript{mnd-2} and Vpx SIV\textsubscript{smm} ternary complexes. This item shows the structural overlap between Vpx SIV\textsubscript{mnd-2} and Vpx SIV\textsubscript{smm} at the DCAF1 interface. This alignment facilitates a comparison of the common interactions and is associated with Figure 3 of the main article.

ITEM 5) Figure S5. Multiple sequence alignment of Vpx and Vpr proteins. This alignment in combination with Vpx SIV\textsubscript{mnd-2} and Vpx SIV\textsubscript{smm} structures shows all the conserved and variable regions found in the different Vpx and Vpr lineages and is associated with Figure 3 and 6 of the main article.
ITEM 6) **Figure S6.** SAMHD1 multiple sequence alignment. This alignment shows the sequence conservation in SAMHD1 NtD and CtD degron regions and highlights the residues making contacts at the Vpx interface. It is associated with Figure 5 and 6 of the main article.

ITEM 7) **Supplemental Figure legends.** Detailed legends for Supplemental Figures S1-S6.

ITEM 8) **Supplemental Table S1A-C.** These tables provide a list of all the interactions observed in the SIV\textsubscript{mnd-2} Vpx/SAMHD\textsubscript{mnd-NtD/DCAF1-CtD ternary complex and are associated Figures 3-6 of the main article.

ITEM 9) **SUPPLEMENTAL EXPERIMENTAL PROCEDURES**
This item provides detailed methods to supplement the experimental procedures reported in the paper.

ITEM 10) **SUPPLEMENTAL REFERENCES**
This item contains the full references that are cited Supplemental Experimental Procedures
Figure S1

A

| Degron          | α-Hsp90 | Hsp90 | GFP-Depron fusion | GFP |
|-----------------|---------|-------|-------------------|-----|
| mnd SAMHD1 1-114| +       |       |                   |     |
| mnd SAMHD1 1-37 | -       |       |                   |     |
| mnd SAMHD1 1-37 | +       |       |                   |     |
| mnd SAMHD1 1-37 | -       |       |                   |     |
| mnd SAMHD1 1-114| +       |       |                   |     |
| mnd SAMHD1 1-114| -       |       |                   |     |
| mnd SAMHD1 1-114| +       |       |                   |     |
| mnd SAMHD1 1-114| -       |       |                   |     |
| mnd SAMHD1 1-114| +       |       |                   |     |
| mnd SAMHD1 1-114| -       |       |                   |     |

B

Degron: mnd SAMHD1 1-114

Degron: mnd SAMHD1 1-37

YFP vs. GFP Scatter Plots
Figure S2

DCAF1-CtD

SIV<sub>mmnd-2</sub> Vpx

SAMHD1<sub>mmnd 1-114</sub>
Figure S3

A

B

SIV_{mind-2} Vpx

H_{2}O

SIV_{smm} Vpx
SUPPLEMENTAL FIGURE LEGENDS

Figure S1, related to Figure 1. Degron fusion proteins. (A) Expression of (NLS)-GFP-degron fusion proteins in was assessed *M. dunni* cells by Western blotting. For each construct the top panel shows an anti-Hsp90 blot loading control. In the lower panels an anti-GFP antibody was used to detect the expression level of each degron fusion indicated. (B) Primary FACS data for degron assays. Stable cell lines expressing degrons containing SAMHD1<sub>mnd</sub> residues 1-114 (upper panels, magenta) or 1-37 (lower panels, purple) were transduced with increasing SIV<sub>mnd-2</sub> Vpx (left to right) and analysed by flow cytometry.

Figure S2, related to Figure 2. Electron density. Stereo image of (2F<sub>obs</sub> – F<sub>c</sub>) refined electron density for DCAF1 (top), Vpx (middle) and SAMHD1 (bottom) contoured at 1 σ. The density is shown as light blue wireframe and the backbone Cα traces of the final refined models as ribbon representation.

Figure S3, related to Figure 2. Vpx conservation of zinc coordination. The protein backbones of (A) Vpx from SIV<sub>mnd-2</sub> (blue) and (B) SIV<sub>smm</sub> (orange) are shown in cartoon representation, α-helices are labelled and zinc ions are shown as grey spheres. Residues that co-ordinate zinc ions are shown as sticks, the coordinating water molecule in SIV<sub>mnd-2</sub> Vpx as a red sphere and co-ordinating bonds as dashed lines.

Figure S4, related to Figure 3. Superposition of DCAF1-CtD-Vpx complexes. (A) Overview of structurally aligned Vpx/DCAF1-CtD complexes. The orientation is as in Figure 3. Bound Vpx molecules are shown in cartoon representation, SIV<sub>mnd-2</sub> Vpx (blue) and SIV<sub>smm</sub> Vpx.
(orange). (B–E) Details of Vpx/DCAF1-CtD interactions in the regions boxed in a. Residues that make interactions are shown in stick representation, the DCAF1-CtD cartoon is coloured white, in the SIV<sub>mnd-2</sub> and grey in the SIV<sub>smm</sub> complexes.

**Figure S5**, related to Figure 3 and Figure 6. Multiple sequence alignment of HIV/SIV Vpx and Vpr proteins. 90% type-conserved amino acid residues are highlighted in red, 60% type-conserved in cyan, variable regions are boxed and the position of secondary structure elements is displayed between the Vpr and Vpx groupings. Red stars above the alignment indicate zinc-binding side chains. Residues that interact with SAMHD1 degrons are indicated with blue (SIV<sub>mnd-2</sub>) or orange (SIV<sub>smm</sub>) stars (side chain) or dots (backbone). Grey stars indicate Vpx residues with side chains that interact with DCAF1. Darker shading is applied to those that are also type-conserved in Vpr. Numbering below is for SIV<sub>mnd-2</sub> 5440 Vpx.

**Figure S6**, related to Figure 5 and Figure 6. Multiple sequence alignment of primate SAMHD1 N- and C-terminal regions. 100% type-conserved amino acid residues are highlighted in red, 60% type-conserved in cyan. SAMHD1 residues that interact with the respective Vpx are indicated with blue (SIV<sub>mnd-2</sub>) or orange (SIV<sub>smm</sub>) stars (side chain) or dots (backbone). Flashes indicate the highly divergent residues 32 and 60.
| Vpx residue | Region | Interaction | DCAF1 residue | Region |
|-------------|--------|-------------|---------------|--------|
| A2<sub>MC</sub> | Nt     | HB          | R1106         | WD40 1 |
| E3<sub>MC</sub> | Nt     | HB          | R1106         | WD40 1 |
| E3         | Nt     | HB          | R1106<sub>MC</sub> | WD40 1 |
| E3         | Nt     | HB          | S1102         | WD40 1 |
| E3         | Nt     | HB          | R1106, F1107  | WD40 1 |
| A5         | Nt     | HI          | F1107, L1119  | WD40 1 |
| P6         | Nt     | HB          | Y1131         | WD40 1 |
| P6         | Nt     | HI          | Y1131         | WD40 1 |
| P6         | Nt     | HI          | M1166, F1170  | WD40 2 |
| E7<sub>MC</sub> | Nt   | HB          | S1168<sub>MC</sub> | WD40 2 |
| E7<sub>MC</sub> | Nt   | HB          | F1170<sub>MC</sub> | WD40 2 |
| I28        | a1     | HI          | W1156         | WD40 2 |
| E31        | a1     | HI          | W1156         | WD40 2 |
| L44        | a2     | HI          | A1377         | WD40 7 |
| T47        | a2     | HI          | L1378         | WD40 7 |
| C48        | a2     | HI          | L1378         | WD40 7 |
| H51        | a2     | HI          | L1378         | WD40 7 |
| C52        | a2     | HI          | L1378         | WD40 7 |
| Y62        | a3     | HB          | D1092         | WD40 7/1 |
| Y65        | a3     | HB          | E1091         | WD40 7/1 |
| R66        | a3     | SB          | E1093         | WD40 7/1 |
| L68        | a3     | HI          | T1114         | WD40 1 |
| L69        | a3     | HI          | C1113, T1114  | WD40 1 |
| L70        | a3     | HI          | A1377, L1378, M1380 | WD40 7 |
| H72        | a3     | HB          | N1135<sub>MC</sub> | WD40 1/2 |
| H72        | a3     | HI          | C1113, T1114  | WD40 1 |
| K73        | a3     | SB          | E1093         | WD40 7/1 |
| K73        | a3     | HB          | S1094<sub>MC</sub> | WD40 7/1 |
| K73        | a3     | HI          | C1113         | WD40 1 |
| M75        | a3     | HI          | W1156         | WD40 2 |
| Y76        | a3     | HB          | T1097<sub>MC</sub> | WD40 1 |
| Y76        | a3     | HB          | F1355<sub>MC</sub> | WD40 7 |
| Y76        | a3     | HI          | A1137, T1139  | WD40 1/2 |
| Y76        | a3     | HI          | F1330         | WD40 6 |
| T77        | a3     | HI          | P1329, F1330  | WD40 6 |
| M79        | a3     | HI          | T1155, W1156  | WD40 2 |
| Q81        | a3     | HB          | Q1314<sub>MC</sub> | WD40 6 |
| Q81<sub>MC</sub> | a3 | HB          | R1225         | WD40 4 |
| Q81        | a3     | HI          | L1313, P1329  | WD40 6 |

HB – hydrogen bond, HI – hydrophobic interaction, SB – salt bridge MC – mainchain
Table S1B, related to Figure 3. DCAF1-SAMHD1 interface residues (499 Å$^2$)

| DCAF1 residue | Region | Type of contact | SAMHD1 residue | Region |
|---------------|--------|----------------|----------------|--------|
| A1089<sub>MC</sub> | WD40 7/1 | HB | R14 | Nt |
| N1090 | WD40 7/1 | HI | R12 | Nt |
| E1091 | WD40 7/1 | SB | R12 | Nt |
| D1092 | WD40 7/1 | SB | R14 | Nt |
| T1114, T1114<sub>MC</sub> | WD40 1 | HB | Q8 | Nt |
| Q1116 | WD40 1 | HI | Q8, R12 | Nt |
| N1132 | WD40 1/2 | HB | D7, Q8<sub>MC</sub> | Nt |
| H1134<sub>MC</sub> | WD40 1/2 | HB | Q8 | Nt |
| N1135 | WD40 1/2 | HB | D5<sub>MC</sub> | Nt |

HB – hydrogen bond, HI – hydrophobic interaction, SB – salt bridge MC – mainchain
| Vpx residue | Region | Interaction | SAMHD1 residue | Region |
|-------------|--------|-------------|----------------|--------|
| P9          | Nt     | HI          | M1, Q2         | Nt     |
| Q10<sub>MC</sub> | Nt | HB          | M1<sub>MC</sub>, Q2<sub>MC</sub> | Nt     |
| G11         | Nt     | HI          | Q2             | Nt     |
| A12         | Nt     | HI          | Q2             | Nt     |
| G13         | Nt     | HB          | D7, S10, S10<sub>MC</sub> | Nt     |
| E14<sub>MC</sub> | Nt | HB          | D7             | Nt     |
| V15<sub>MC</sub> | Nt | HB          | D7             | Nt     |
| V15         | Nt     | HI          | D7, P9         | Nt     |
| L17         | α1     | HI          | P9, P13        | Nt     |
| W20         | α1     | HI          | Q8, P9         | Nt     |
| N29         | α1     | HI          | V36            | Nt     |
| E39         | α2     | HB          | P47<sub>MC</sub>, E48<sub>MC</sub>, Q49<sub>MC</sub> | SAM domain |
| L41         | α2     | HI          | V36            | Nt     |
| F42         | α2     | HI          | L38, Q49, F52  | SAM domain |
| W45         | α2     | HI          | P34            | Nt     |
| N46         | α2     | HB          | E48, E48<sub>MC</sub> | SAM domain |
| V49         | α2     | HI          | R55            | SAM domain |
| E50         | α2     | SB          | R69            | SAM domain |
| H53         | α2     | HI          | R20            | Nt     |
| D54         | α2     | SB          | R20            | Nt     |
| D54         | α2     | SB          | R55            | SAM domain |
| H56         | α2     | HI          | F15            | Nt     |
| Q57         | α2/ α3 loop | HB  | R20            | Nt     |
| Q57<sub>MC</sub> | α2/ α3 loop | HB  | R20<sub>MC</sub> | Nt     |
| R58         | α2/ α3 loop | HI  | F15            | Nt     |
| R58         | α2/ α3 loop | HB  | F15<sub>MC</sub> | Nt     |
| S59<sub>MC</sub> | α2/ α3 loop | HB  | S18<sub>MC</sub> | Nt     |
| Y62         | α3     | HI          | F15, S18      | Nt     |
| Y65         | α3     | HI          | P13            | Nt     |
| Y65         | α3     | HB          | R12            | Nt     |

HB – hydrogen bond, HI – hydrophobic interaction, SB – salt bridge MC – mainchain
SUPPLEMENTAL EXPERIMENTAL PROCEDURES

Degron assay

Degron reporter constructs comprising two copies of Nuclear Localisation Signal (NLS)-EGFP fused to N-terminal sequences from SAMHD1_{mnd} (NLS-EGFP-SAMHD1_{mnd}-NtD) were generated by replacing the human SAMHD1-CtD degron sequence in pCMS28-NLS-EGFP-SAMHD1-CtD (Schwefel et al., 2014) with sequences from the N-terminal region of SAMHD1_{mnd}. DNA coding for residues 1-114, 1-37, 37-114, 5-114 or 10-114 was amplified by PCR and inserted into the reporter construct using XhoI/EcoRI restriction sites. Point mutations were created by PCR-based site directed mutagenesis. Virus-like particles (VLPs) were generated by co-transfecting 293T cells with pVSVG, pKB4 and pCMS28-NLS-EGFP-SAMHD1 human-CtD or mandrill-NtD, wildtype or mutant (Schwefel et al., 2014). Stable cell lines were produced by transduction of *Mus dunni* cells followed by puromycin selection. Expression of degron constructs was assessed using Western blotting with anti-EGFP antibodies.

The SIV_{smm} Vpx sequence was amplified by PCR from pIRES2-EGFP-Vpx (a gift from Mario Stevenson), and SIV_{mnd.2} Vpx was amplified from the PET49 plasmid used for *E. coli* expression. Sequences were inserted into pENTR/D/TOPO (Invitrogen) and transferred into pLgatewayIeYFP (Gateway LR clonase™ II, Invitrogen) to create bicistronic Vpx-IRES-YFP expression constructs. Point mutations were created by PCR-based site directed mutagenesis. VLPs expressing Vpx-IRES-YFP were generated by co-transfecting 293T cells with pVSV-G, pKB4 and pLgatewayIeYFP-Vpx (wildtype or mutants) (Schwefel et al., 2014). Approximately 18 hours after transfection, cells were washed and sodium butyrate medium (0.02 M sodium butyrate, 10% FCS and 1% penicillin/streptomycin in DMEM) was added for 6 hours before

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*Schwefel et al., 2014*
replacing with fresh media. After a further 15 hours VLPs were harvested from the media by filtration.

Parental *Mus dunni* or stable cell lines expressing degron reporters were seeded at $5 \times 10^4$ cells per well in a 24-well plate one day prior to infection. Cells were infected with 2-fold serial dilutions of Vpx-YFP VLPs in the presence of $1 \mu$g/mL Polybrene. After 48 hours, cells were harvested and the percentage of EGFP-positive and YFP-positive cells was determined by flow cytometry using a FACSVerse analyser (BD Biosciences).

**Protein expression and purification**

The nucleotide sequences coding for SIV<sub>mnd-2</sub> Vpx isolate 5440 (Hu et al., 2003) and amino acid residues 1-114 of SAMHD1<sub>mnd</sub> (Uniprot ID H6WEA4) were synthesised codon-optimised for *E. coli* (Life Technologies). The open reading frames were inserted into pET-49b and pET-52b (Merck Millipore) expression plasmids respectively using flanking XmaI/NotI restriction sites to generate N-terminally GST-tagged and N-terminally Strep-II-tagged fusion proteins. DCAF1-CtD was cloned and expressed as described previously (Schwefel et al., 2014).

SIV<sub>mnd-2</sub> Vpx and SAMHD1<sub>mnd-NtD</sub> were expressed in the *E. coli* strain Rosetta 2 (DE3) (Merck Millipore). Bacterial cultures were grown in terrific broth medium in a shaking incubator at 37 °C. Protein expression was induced by the addition of 0.1 mM IPTG at $A_{600} = 0.5$, then further incubated at 18 °C for 20 hours to express recombinant proteins. Cells were harvested by centrifugation for 20 min at 4,500 xg and 4 °C, the cell pellets resuspended in 30 mL lysis buffer (50 mM Tris-HCl pH 7.8, 500 mM NaCl, 4 mM MgCl$_2$, 0.5 mM TCEP, 1x EDTA-free mini complete protease inhibitors (Roche), 0.1 U/ml Benzonase (Novagen) per pellet of 1 L bacteria culture and stored at -20 °C.
SAMHD1\textsubscript{mnd}-NtD cell suspensions were lysed by disruption using an EmulsiFlex-C5 homogeniser (Avestin). The lysate was cleared by centrifugation for 1 hour at 48,000 xg at 4 °C. All further purification steps were performed at 4 °C or on ice. Lysates were applied to 10 mL StrepTactin column (IBA). The column was washed with 600 mL of Wash buffer (50 mM Tris-HCl pH 7.8, 500 mM NaCl, 4 mM MgCl\textsubscript{2}, 0.5 mM TCEP) and bound proteins were eluted in wash buffer containing 2.5 mM d-desthiobiotin. Eluted fractions were concentrated to 5 mL and further purified on a Superdex75 gel filtration column (GE Healthcare) equilibrated in Gel filtration buffer (10 mM Tris-HCl pH 7.8, 150 mM NaCl, 4 mM MgCl\textsubscript{2}, 0.5 mM TCEP). Peak fractions containing SAMHD1\textsubscript{mnd}-NtD were pooled, concentrated to 30 mg/mL, snap-frozen in liquid nitrogen in small aliquots and stored at -80 °C.

\textit{Protein complex assembly}

Cell suspension from 1 L of GST-SIV\textsubscript{mnd-2} Vpx was lysed by homogenisation in an EmulsiFlex-C5 (Avestin). The lysate was cleared by centrifugation for 1 h at 48,000 xg at 4 °C. All further purification steps were performed at 4 °C or on ice. 1 mL of glutathione Sepharose (GSH-Sepharose) beads (GE Healthcare) were added to the lysate and incubated for 1 hour on a roller agitator. Beads were pelleted by centrifugation at 4000 xg for 10 min, the supernatant was discarded and the beads washed four times with 50 mL of Wash buffer. For assembly, the GST-SIV\textsubscript{mnd-2} Vpx bound beads were resuspended in 10 mL wash buffer in a 15 mL Falcon tube. 1 mg of DCAF1-CtD together with an equimolar amount of SAMHD1\textsubscript{mnd}-NtD and 1 mg of HRV-3C protease (GE Healthcare) were added and the tube was incubated overnight on a rolling agitator. Beads were then removed by centrifugation at 4000 xg for 10 min. The supernatant was concentrated to 5 mL and applied to a Superdex200 size exclusion column equilibrated in Gel
filtration buffer. Peak fractions containing the ternary complex were pooled, concentrated to 12 mg/mL, snap-frozen in liquid nitrogen in small aliquots and stored at -80 °C.

**Crystallization and data collection**

Crystals of the SIV_{mnd-2} Vpx/SAMHD1_{mnd}-NtD/DCAF1CtD complex were grown using the hanging drop vapour diffusion method by mixing 1 µL complex at a concentration of 6.34 mg/mL with 1 µL of reservoir solution containing 0.16 M Trisodium Citrate-HCl pH 5.2 and 4% PEG 6000. Drops were equilibrated over a 450 µL reservoir solution at 18 °C. Crystals were adjusted to 25 % glycerol and cryo-cooled in liquid nitrogen. A data set from a single crystal was collected on station I04 at the Diamond synchrotron light source, UK at a wavelength of 0.97965 Å.

**Structure solution**

Diffraction data were reduced with the program XDS (Kabsch, 2010). The high resolution cut-off was based on the CC_1/2 criteria (Karplus and Diederichs, 2012). The structure was solved by molecular replacement with the program Molrep (Vagin and Teplyakov, 2010) using the previously determined DCAF1-CtD structure and a homology model constructed with the previously determined SIV_{smm} Vpx as template (PDB code 4CC9 (Schwefel et al., 2014)). The SAM domain was placed manually into density using the NMR structure of the human SAMHD1 SAM domain as guidance (PDB code 2E8O). Iterative model adjustment using the program Coot (Emsley et al., 2010) combined with positional, real-space, individual b-factor and TLS refinement with the program phenix.refine (Adams et al., 2010) produced a final model for DCAF1-CtD residues 1073-1315, 1327-1392 (chain A), SIV_{mnd-2} Vpx residues 2-86 (chain B)
and SAMHD1_{mnd-NtD} residues 1-22, 34-88, 93-109 (chain C) with R(R_{free})-factors of 17.5\% (23.1\%). 95.8\% of all residues fall in the favoured region of the Ramachandran plot with 0.42\% outliers. Data collection and refinement statistics are shown in Table 1.

**Multiple sequence alignment**

Amino acid sequences were aligned using the ClustalW server and adjusted manually.

NCBI accession numbers for Vpr sequences: HIV-1 ETH220 - U46016, HIV-1 WEAU - U21135, HIV-1 pNL4-3 - AF324493, HIV-1 pNL432 - M28355, SIV_{srm} - AF077017, SIV_{cprCAM13} -AY169968, SIVcpz_{Lb7}- DQ373064, SIV_{mac239} - M33262, SIV_{mac251} - M76764, SIV_{rcm} - HM803689, SIV_{mnd2} - AF367411, SIV_{agmSAB1} - U04005, SIV_{agmGRi} - M66437, SIV_{agmVER} - KF741091, SIV_{agmTAN1} - U58991; for Vpx sequences: HIV-2A - M30502, HIV-2B - U27200, SIV_{srm} - AF077017, SIV_{mac251} - M76764, SIV_{mac239} - M33262, SIV_{rcmNG} - AF349680, SIV_{rcmCAM} - HM803689, SIV_{rcmGAB1} - AF382829, SIV_{dr1} - AY159321, SIV_{mnd2CM16} - AF367411, SIV_{mnd25440} - AY159322, SIV_{mnd2M14} - AF328295; for SAMHD1 sequences: Cebus paella - JN936910, Callithrix jacchus - JN936906, Ateles geoffroyi - JN936911, Alouatta palliata - JN936912, Hylobates lar - JN936889, Pongo pygmaeus - JN936888, Pan troglodytes - JN936887, Homo sapiens - BC036450, Colobus angolensis - JN936905, Chlorocebus tantalus 1 - JN936891, Chlorocebus tantalus 2 - JN936892, Chlorocebus pygerythrus - JQ231137, Cercopithecus Diana - JN936902, Cercopithecus neglectus - JQ231141, Macaca mulatta 1 - JN936894, Macaca mulatta 2 - JN936895, Macaca fascicularis - JN936893, Papio hamadryas - JN936890, Mandrillus sphinx - JN936897, Cerocebus atys - JQ231132, Cerocebus torquatus - JQ231133, Cerocebus chrysogaster 1 - JN936898, Cerocebus chrysogaster 2 - JN936899.
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