The Carboxyl-Terminus of Human Immunodeficiency Virus Type 2 Circulating Recombinant form 01_AB Capsid Protein Affects Sensitivity to Human TRIM5α

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Abstract

Human immunodeficiency virus (HIV) type 2 shows limited geographical distribution compared with HIV type 1. Although 8 genetic groups of HIV type 2 (HIV-2) have been described, recombinant viruses between these groups are rarely observed. Recently, three HIV-2 patients in Japan were described with rapidly progressive, acquired immunodeficiency. These patients were infected with an A/B inter-group recombinant designated CRF01_AB. Here, we characterize the capsid protein (CA) encoded by the viruses from these patients. HIV-2 CRF01_AB CA showed unique amino acid sequence almost equally distinct from group A and group B viruses. Notably, HIV-2 CRF01_AB CA showed potent resistance to human TRIM5α. In addition to the previously identified amino acid position 119 in the N-terminal domain of CA, we found that HIV-2 CRF01_AB-specific amino acid substitutions in the C-terminal domain also were necessary for resistance to human TRIM5α. These results indicate that retroviruses can evade TRIM5α by substitution at residues within the C-terminal domain of CA.

Citation: Miyamoto T, Nakayama EE, Yokoyama M, Ibe S, Takehara S, et al. (2012) The Carboxyl-Terminus of Human Immunodeficiency Virus Type 2 Circulating Recombinant form 01_AB Capsid Protein Affects Sensitivity to Human TRIM5α. PLoS ONE 7(10): e47757. doi:10.1371/journal.pone.0047757

Introduction

Human immunodeficiency virus type 2 (HIV-2) has been detected primarily in West Africa, in contrast to the global distribution of the type 1 epidemic virus (HIV-1). Based on molecular evidence, HIV-2 and HIV-1 are presumed to derive from simian immunodeficiency viruses that originated in sooty mangabeys (SIVSm) and chimpanzees (SIVcpz), respectively, as a result of zoonotic transfer between non-human primates and human. The HIV-1 and HIV-2 bear a considerable degree of homology in both gene organization and RNA sequence [30%–60%] [1–4]. It is generally believed that HIV-2 is less pathogenic than HIV-1. However, certain HIV-2 patients with high plasma HIV-2 loads develop acquired immune deficiency syndrome (AIDS) as rapidly as HIV-1 patients do [4]. To date, eight HIV-2 genomes have been distinguished based on the pattern of phylogenetic (sequence) analysis; each group is presumed to have originated from an independent zoonotic event [5].

TRIM5α was identified as a factor that restricts HIV-1 infection in rhesus monkey (Rh) cells [6]. TRIM5α is thought to degrade the core of the incoming virus [7,8]. TRIM5 proteins are members of the tripartite motif family containing RING, B-box, and coiled-coil domains. The alpha isoform of TRIM5 has an additional C-terminal PRYSPRY (B30.2) domain [9]. In cynomolgus monkey (CM), TRIM5α also has been demonstrated to restrict HIV-1 infection [6,10]. In contrast, the human TRIM5α exhibits minimal restriction of HIV-1 infection [11–14], but shows moderate levels of restriction for HIV-2 [15].

Capsid (CA) proteins are components of the viral core; the CAs of HIV-1 and HIV-2 have similar primary and three dimensional structures [16]. CA is composed of a surface-exposed N-terminal domain (NTD) and a C-terminal domain (CTD) that is required for oligomerization [17]. We previously identified a single amino acid of the HIV-2 capsid that determines the susceptibility of HIV-2 to CM TRIM5α. Viruses that encoded CAs with either alanine or glutamine at amino acid residue 119 (which corresponded to the 120th amino acid of the CA of the G1H123 viral strain) could grow in cells harboring the CM TRIM5α. In contrast, HIV-2 encoding CA with proline at the same position showed restricted growth in cells harboring the CM TRIM5α. Similar results, although to a lesser extent, were observed when the human TRIM5α was used [15]. Furthermore, an analysis of HIV-2 CA variation in a West African Caio cohort demonstrated that the presence of proline at CA positions 119, 159, and 178 was more frequent in individuals with lower viral loads (VLs); the presence of non-proline residues at all 3 residues was more frequent in
individuals with high VLs. The in vitro replication levels of viruses bearing changes at the 3 positions suggested that these 3 residues influence virus replication by altering susceptibility to TRIM5α [10]. These results also suggested that TRIM5α controls virus replication in HIV-2-infected individuals.

Recently, five HIV-2-seropositive cases were identified in Japan. Three isolates (NMC307, NMC716, and NMC842) were recovered from these patients and were shown by full-length genomic analysis to represent a recombinant (designated HIV-2 CRF01_AB) of group A and B strains [19]. Although more than 75% of patients with HIV-2 have asymptomatic prognoses throughout their lifetimes [1,20], all 3 of the CRF01_AB patients were found to be at an advanced stage of AIDS with low CD4+ cell counts and high HIV-2 VLs [19]. All 3 patients were under 40 years of age when first diagnosed as HIV-2 positive [19]. Assessment of risk factors suggested that all three were infected via heterosexual contacts; no personal connection was confirmed among any of these cases [19]. In the present study, we characterized the HIV-2 CRF01_AB CA obtained from these patients and found several unique properties of HIV-2 CRF01_AB, including potent resistance to human TRIM5α-mediated restriction.

Results

HIV-2 CRF01_AB Strains Show Unique CA Sequences

Fig. 1 shows an alignment of the deduced amino acid sequences of the CAs of HIV-2 group A (ROD, UC12, GH123, and UC2), HIV-2 group B (UC14, D205, and UC1), SIVs (SIVmac239 and SIVsm PBj14), and HIV-2 CRF01_AB (NMC307, NMC716, NMC842, and 7312A). As we reported previously [15,18], the 119th amino acid position is a proline, glutamine, or alanine in the CAs of HIV-2 group A, HIV-2 group B, and SIVs. However, the CAs of the HIV-2 CRF01_AB strains uniquely possess a glycine at this position. Based on the genomic structure of HIV-2 CRF01_AB, A/B recombinant breakpoints within this isolate are located near or within the env gene, such that HIV-2 CRF01_AB can be considered to consist of a group B backbone that incorporates group A env fragments [19]. These presumed breakpoints could be taken to suggest that CRF01_AB CA should be encoded as a B-like sequence. However, phylogenetic analysis of these CA sequences (Fig. 2) reveals that the deduced HIV-2 CRF01_AB CA proteins constitute a distinct cluster, with the dendrogram exhibiting a long branch length compared to the CAs of HIV-2 group A, HIV-2 group B, and SIV.

HIV-2 CRF01_AB CA is Highly Resistant to Human TRIM5α

In a previous study, we reported that the amino acid at residue 119 of the HIV-2 CA affects susceptibility to the restriction of virus replication by CM and human TRIM5α [15], such that HIV-2 encoding CA(Pro119) was sensitive to CM and human TRIM5α, while HIV-2 encoding CA(Gln119) or CA(Ala119) was resistant [15]. We also reported that mutation of HIV-2 strain GH123 to encode glycine at the corresponding position (GH123/G) rendered GH123 resistant to CM TRIM5α [21]. To further test the role of the CA protein in TRIM5α resistance, we generated recombinant versions of the GH123 virus (716 or 842) in which the CA-encoding segment of gag was replaced with that of the A/B recombinants NMC716 or NMC842 (respectively). We used a recombinant Sendai virus (SeV) system to express CM, Rh, and human TRIM5α and CM TRIM5α lacking the PRYSPRY domain as a negative control (Fig. S1). In the presence of CM TRIM5α, infection by the parental GH123 virus was restricted, but infection by GH123/G was resistant to CM TRIM5α-mediated restriction (Fig. 3A). Infection by 716 or 842 was resistant to CM TRIM5α (Fig. 3B). In contrast, infection by any of the 4 variants (GH123, GH123/G, 716, and 842) was completely restricted by Rh TRIM5α (Fig. 3A, B). These results for cells producing CM or Rh TRIM5αx are consistent with our previous findings [22]. In cells producing human TRIM5α, the replication of parental GH123 and of the GH123/G mutant were partially restricted (Fig. 3A), while 716 and 842 replicated as efficiently as in negative control cells that did not produce a functional TRIM5αx (Fig. 3B). The mean ratios of the p25 levels at 6 days after infection in the cells producing human TRIM5αx to those in the negative control cells were 0.14 for GH123, 0.30 for GH123/G, but 0.81 for 716 and 1.02 for 842 in three independent experiments. The ratio of GH123/G was significantly higher than that of GH123 (P = 0.0086, t test) but lower than those of 716 (P = 0.0059, t test) and 842 (P = 0.0030, t test). Similar results were obtained when we calculated the mean ratios of the p25 levels at 3 days after infection (data not shown). These data indicate that the CA sequences of the CRF01_AB strains conferred higher potential to escape from human TRIM5α than those of GH123/G.

Viral Sensitivity to Human TRIM5α-mediated Restriction in a Single Round Infection Assay

TRIM5α restricts viral infection at a post-entry step [6,23,24]. To focus on early steps of virus replication, we performed a single-round infection assay, in which infection is detected as fluorescence generated by production of the green fluorescent protein (GFP). To construct mutant viruses encoding GFP, the fragment of GH123, 842, or GH123/G that encoded the matrix (MA) and CA proteins was transferred to the env-disrupted HIV-2 genomic clone pROD-env(-)-GFP, which directs the production of GFP after infection [25]. Vesicular stomatitis virus glycoprotein (VSV-G) pseudotyped wild-type and mutant HIV-2 GFP viruses were inoculated into feline CRFK cells producing TRIM5α, and GFP-positive cells were counted 2 days after infection. In this experiment, we used feline cells, since feline cells lack expression of endogenous TRIM5α. In the presence of CM TRIM5α, the numbers of GFP-positive cells were greater in cells infected with GFP-expressing viruses encoding the GH123/G or 842 CAs than in those infected with the GFP-expressing viruses encoding GH123 CA (Fig. 4), confirming that viruses encoding CA(G119) were resistant to CM TRIM5αx. Consistent with the results shown in Fig. 3B, the GFP-expressing virus encoding the 842 CA from a patient was more resistant to human TRIM5α-mediated restriction than viruses encoding the CAs from GH123 (P = 0.0010, t-test) or GH123/G (P = 0.0026, t-test) (Fig. 4).

Viral Growth in TRIM5α Knock-down Cells

We next investigated whether the different resistance to human TRIM5α restriction among recombinant HIV-2 strains still applied in cells producing physiological levels of human TRIM5α protein. For this purpose, we used TRIM5α “knock-down” Jurkat cells (TRIM5α-KD Jurkat) and the corresponding control Jurkat line (Luci-siRNA Jurkat) [26]. It was demonstrated that the level of TRIM5α mRNA in TRIM5α-KD Jurkat is five times lower than that of Luci-siRNA Jurkat by TaqMan quantitative PCR. Three days after infection, GH123 replicated better in TRIM5α-KD Jurkat than in Luci-siRNA Jurkat (Fig. 5A). On the other hand, GH123/G, 716, and 842 yielded comparable titers in both cell lines (Fig. 5B, 5C, and 5D). In this experiment, we found that GH123/G also was resistant to human TRIM5α. Nevertheless, the data presented in Fig. 5 indicated that GH123 was sensitive to human TRIM5α produced at physiologically relevant levels, while 716 and 842 possessed potent resistance against human TRIM5α.
| HIV-2A | ROD | PVQHVGG-NYTHIPLSPRITLNAWVKLVEEEKFGAEVPGFQALSEGCTPDINQMVLNCVG | 59 |
|-------|-----|-------------------------------------------------------------|----|
| UC1   | .   | .Q.A-.-.V---------------------------------------------------| 59 |
| GH12  | .   | .T.G-.-.I.V--------------------------------------------------| 60 |
| UC2   | .   | .Q.A-.-.V.V---------------------------------------------------| 59 |
| HIV-2B| UC14| .QIA-.-.S.L---------------------------------------------------| 59 |
| D205  | .   | .QLA-.-.V.L---------------------------------------------------| 59 |
| UC1   | .   | .QIA-.-.V.M---------------------------------------------------| 59 |
| SIV   | mac239| .QI-.-.V.L--------------------------------------------------| 59 |
| PB134  | .   | .QI-.-.L------------------------------------------------------| 59 |
| HIV-2AB| NMC307| .Q.A-.-.V.V--------------------------------------------------| 59 |
| NMC716| .   | .Q.A-.-.V.V---------------------------------------------------| 59 |
| NMC842| .   | .Q.A-.-.V.V---------------------------------------------------| 59 |
| 7312A | .   | .Q.A-.-.V.V---------------------------------------------------| 59 |

**Figure 1.** Alignments of amino acid sequences of CA proteins encoded by selected HIV-2 isolates and SIV from the Los Alamos databases. Dots denote amino acid identity with the ROD CA; dashes denote gaps introduced to optimize alignment. HIV-2 CRF01_AB-specific amino acid residues are in red. Arrows indicate key residues at 119, 159, and 178, and the position (in the corresponding DNA sequence) of the HindIII restriction site used in the constructs. HIV-2A, HIV-2B, and HIV-2AB denote HIV-2 group A, HIV-2 group B, and HIV-2 CRF01_AB, respectively. doi:10.1371/journal.pone.0047757.g001
Since TRIM5α-KD Jurkat always showed reduced proliferative properties compared to Luci-siRNA Jurkat (data not shown), presumably due to reduced TRIM5α levels [27], the p25 levels of all these viruses in Luci-siRNA Jurkat became higher than those in TRIM5α-KD Jurkat at 10 days after infection (data not shown).

HIV-2 CRF01_AB CA C-terminal Domain-specific Sequence also Affects Viral Sensitivity to Human TRIM5α

We previously reported that the presence of proline at CA positions 119, 159, and 178 is more frequent in individuals with lower VLs [18]. Viral isolates NMC307, NMC716, and NMC842 all encoded CAs with proline at the 159th position (Fig. 1). However, the 178th amino acid residue was encoded as a threonine (NMC307 and NMC842) or as a glutamic acid (NMC716) in these isolates (Fig. 1). To test whether a single residue at amino acid 178 of HIV-2 CRF01_AB CA affects the sensitivity to human TRIM5α, we generated recombinant 716 or 842 viruses (designated 716GPP or 842GPP, respectively) that encoded CA (Pro178) proteins. As shown in Fig. 3C, 716GPP and 842GPP escaped from human TRIM5α restriction as efficiently as 716 and 842 did. These data suggest the existence of viral determinants for human TRIM5α-resistance other than the previously identified 119th and 178th amino acid positions of CA.

To search for the viral determinants of human TRIM5α resistance other than the 119th and 178th amino acid positions of HIV-2 CA, we constructed a chimeric virus 842Hind by replacing the segment of the 842 genome that encodes CA C-terminal residues 170 to 231 with the corresponding region of GH123 (Fig. 6A). When tested in cells that produced human TRIM5α, 842 was strongly resistant to human TRIM5α as expected (Fig. 6B). However, the 842Hind construct, which encoded the NMC842 CA with the GH123 CA C-terminal short region, lost this resistance to human TRIM5α (Fig. 6C). The mean ratios of the p25 levels at 6 days after infection in the cells producing human TRIM5α to those in the negative control cells were 0.73 for 842 and 0.16 for 842Hind in three independent experiments. The ratio of 842Hind was significantly lower than that of 842 (P=0.0003, t test). Similar results were obtained when we calculated the mean ratios of the p25 levels at 3 days after infection (data not shown). These results suggest that one or more of the HIV-2 CRF01_AB-specific amino acid residues within the CA C-terminal short region (Fig. 1, shown in red) also are necessary to fully evade human TRIM5α.

Molecular Dynamics of N-terminal Domain (NTD) of HIV-2 CRF01_AB CA

Residue 120 of the GH123 CA, which corresponds to residue 119 of the CRF01_AB CA, is located in the loop between α-helices 6 and 7 (L6/7) of CA NTD. Our previous molecular dynamics simulation study of HIV-2 CA NTD revealed that mutations at this position affected conformation of the neighboring loop between α-helices 4 and 5 (L4/5), and TRIM5α-sensitive viruses were predicted to share a common L4/5 conformation. In addition, the shared L4/5 structures of the sensitive viruses were associated with a decreased probability of hydrogen bond formation between GH123 CA’s Asp97 (in L4/5) and Arg119 (corresponding to residue 118 in HIV-2 CRF01_AB CA; in L6/7) [21]. TRIM5α-resistant viruses exhibited a variable L4/5 conformation and a higher probability of hydrogen bond formation between L4/5 and L6/7 [21]. As noted above, HIV-2 CRF01_AB strains have a unique Gly119 (Fig. 1), which we had not previously modeled by molecular dynamics simulation. Therefore, three-dimensional (3-D) models of HIV-2 GH123/G and NMC842 CA NTD were constructed using homology modeling based on the crystal structures of the HIV-2 CA NTD, and the models were subjected to molecular dynamics simulation to compare the results with those derived from previously constructed 3-D structural models of TRIM5α-sensitive GH123 and TRIM5α-resistant GH123/Q and GH123/A [21]. GH123/Q and GH123/A encode CA (Glu120) and CA (Ala120), respectively [15]. Contrary to our expectation, the predicted L4/5 conformations of the NTDs of the NMC842 CA and GH123/G CA differed from those of...
TRIM5α-resistant GH123/Q and GH123/A, better resembling that predicted for the CA NTD encoded by TRIM5α-sensitive GH123 (Fig. 7). Indeed, the calculated probability of hydrogen bond formation between L4/5 and L6/7 was even lower for the CAs of GH123/G (20.80%) and NMC842 (30.58%) compared to that of GH123 (44.6%). These results suggest that Gly119 endows the CRF01_AB CA NTD with unique structural properties.

Steric Locations of HIV-2 CRF01_AB-specific Amino Acid Substitutions

As noted above, HIV-2 CRF01_AB strains have several specific amino acid substitutions at the C-terminal domain (CTD) of CA (Fig. 1, shown in red); these substitutions were necessary for the potent resistance of these isolates against human TRIM5α (Fig. 6). Previously, we suggested that magnitudes of the computationally calculated binding energies of the CA CTD dimer models tend to be significantly greater in the TRIM5α-less-sensitive HIV-2s in West Africa [18]. To examine if the HIV-2 CRF01_AB-specific amino acid substitutions in CA CTD could influence the CTD-CTD dimer stability, we constructed the CA CTD dimer model of HIV-2 CRF01_AB NMC842 by homology modeling and analyzed steric locations of the specific substitutions and binding energies of the CTD dimer model. In the CA CTD dimer model of NMC842, HIV-2 CRF01_AB-specific amino acid substitutions are located in helix 9 and in the loop between helices 10 and 11, and all appeared to be situated near but distinct from the CTD-CTD dimer interface (Fig. 8A). The predicted binding energy of the CTD-CTD dimer model of the NMC842 isolate (79.6 kcal/mole) was similar to that reported in TRIM5α-sensitive viruses [18]. The results may imply that the HIV-2 CRF01_AB-specific amino acid substitutions in CTD do not necessarily influence the CTD-CTD dimer stability of the TRIM5α sensitive virus.

To further obtain structural insights into the roles of these CRF01_AB-specific mutations, we analyzed their steric locations in the CA hexamer. In the hexamer model of GH123 CA that we previously constructed based on the HIV-1 CA hexamer [28], HIV-2 CRF01_AB-specific amino acid substitutions in CTD form clusters and are located at the outermost part of the hexamer (Fig. 8B and C). Notably, these substitutions exist directly under the L4/5 of neighboring CA (Fig. 8C), and most of them are clearly visible from right above (Fig. 8B). These results raise a possibility that HIV-2 CRF01_AB-specific amino acid substitutions in CA CTD may be exposed to and accessible from the outside of the viral core.

Discussion

In the present study, we have shown that the CA of HIV-2 CRF01_AB isolates have a unique feature distinct from that of other HIV-2 strains; CRF01_AB-specific sequences conferred strong resistance to human TRIM5α. In addition to the previously identified role of amino acid 119 of the CA NTD, CRF01_AB-specific amino acid substitutions in the CA CTD also were necessary for strong resistance to human TRIM5α. These amino
Acid substitutions in CA CTD may be exposed to and accessible from the outside of the viral core.

Retroviral CA is known to form hexamers [29]. The CTD domain of retroviral capsid protein participates in CA dimerization, where intermolecular CTD-CTD interactions are mediated by symmetric, parallel dimerization of helix 9 from the CTD domains of adjacent hexamers [30]. This dimerization process is prerequisite for assemblies of multiple hexamers [29]. Previously, we found that the computationally calculated binding energies of the CA CTD dimer models could have positive relations with the TRIM5α susceptibilities of HIV-2s in West Africa [18]. We therefore calculated here the binding energy of the CTD-CTD dimer model of the NMC842 using computational method. However, the predicted binding energy of the CTD-CTD dimer of the NMC842 isolate was rather similar to that reported in TRIM5α sensitive viruses [18]. Therefore, previously undescribed mechanisms may be involved in the TRIM5α resistance of the HIV-2 CRF01_AB.

A possible mechanism for the findings may be that the CRF01_AB-specific substitutions influence directly or indirectly the structural properties of an interaction surface for the TRIM5α mediated inhibition. In this regard, we previously suggested with SIV that not only the NTD but also the CTD might constitute an intermolecular interaction surface [31]. Similarly, HIV-2 may have such interaction surface in CTD domain, and the surface may be used for the TRIM5α-mediated inhibition. Results on the steric locations of the CRF01_AB-specific substitutions in the hexamer model support this possibility (Fig. 7B and C). A preliminary modeling study of the assemblies of the CA hexamers also have supported this possibility; the NTDs are apart from each other among the hexamers, which allows to form accessible surface on the CTDs (data not shown), as suggested with Rous sarcoma virus CA [32]. Therefore, it would be interesting to examine whether HIV-2 CRF01_AB-specific amino acid substitutions in CTD could constitute a binding cleft for the TRIM5α itself or others involved in TRIM5α mediated inhibition in the assemblies of multiple CA hexamers in the viral core. Further study is necessary to address this issue.

Previously, we showed that the amino acid replacements at CA residue 119 affected the conformation of the neighboring L4/5, and that TRIM5α-sensitive viruses had a shared L4/5 conformation that was associated with a decreased probability of hydrogen bonding between L4/5 and L6/7 [21]. Although GH123/G and 842 showed resistance to TRIM5α, the calculated probability of hydrogen bond formation between L4/5 and L6/7 was lower than that calculated for the CAs of other TRIM5α-resistant viruses, including that from GH123/Q (55.15%) and GH123/A (64.47%) [21]. The conformations of L4/5 in the CAs of GH123/G and 842 also were similar to those of TRIM5α-sensitive viruses, and were distinct from those of the CAs of TRIM5α-resistant viruses. These characteristics of GH123/G and 842 were similar to those of GH123/E and GH123/D, mutant GH123 clones encoding glutamic acid and aspartic acid (respectively) at the residue corresponding to the residue 119.

**Figure 5. Viral growth in TRIM5α knock-down cells.** (A), (B), (C) and (D) TRIM5α-KD Jurkat (“knock-down”) or Luci-siRNA Jurkat (control) cells were infected with derivatives of GH123 virus. Culture supernatants were periodically assayed for levels of virus capsid. Error bars show actual fluctuations of duplicate samples from one of two independent experiments. Black and white bars denote TRIM5α-KD Jurkat and Luci-siRNA Jurkat cells, respectively.

doi:10.1371/journal.pone.0047757.g005
position 119 of HIV-2 CRF01_AB strains [21]. Although glutamic acid and aspartic acid have not been observed at this CA residue in HIV-2 isolated clinically, both GH123/E and GH123/D showed resistance against CM TRIM5α. In contrast to the CAs of GH123/Q and GH123/A, the CAs of both GH123/E and GH123/D show reduced likelihoods of hydrogen bond formation between the L4/5 and L6/7, and the L4/5 conformations were predicted to be similar to those of the CAs of TRIM5α-sensitive viruses. Therefore, our present results extend our previous observations, and additionally imply that the Gly119 of HIV-2 CRF01_AB CA prevents binding by TRIM5α, probably due to the small size of the glycine side chain. It is possible that the shared conformation of L4/5 might have some advantages in utilizing certain cellular factor(s) that bind CA. Our structural data suggests that HIV-2 CRF01_AB strains are highly adapted, since these strains have acquired potent resistance against TRIM5α without losing the shared L4/5 conformation.

In the case of GH123/E, disruption of the hydrogen bond between L4/5 and L6/7 by substitution of alanine for aspartic acid at position 97 (D97A) did not alter the resistant phenotype of HIV-2 CRF01_AB CA prevents binding by TRIM5α, probably due to the small size of the glycine side chain. It is possible that the shared conformation of L4/5 might have some advantages in utilizing certain cellular factor(s) that bind CA. Our structural data suggests that HIV-2 CRF01_AB strains are highly adapted, since these strains have acquired potent resistance against TRIM5α without losing the shared L4/5 conformation.

In the Los Alamos databases, almost all SIV isolates encode glutamine at the position corresponding to residue 119 of the HIV-2 CRF01_AB CA. It is likely that the sequential mutation from glutamine (encoded as CAA or CAG) to proline (C\textsubscript{GP}A or C\textsubscript{GG}; underlines denote single nucleotide changes) and then to alanine (G\textsubscript{GA} or G\textsubscript{GG}) occurred after transmission of the monkey virus to the human population. The nature of the genetic code suggests that the Gly119-encoding virus (G\textsubscript{GA} or G\textsubscript{GG} codon) derived from the Ala119-encoding virus, implying that the viruses with glycine are highly adapted, as also discussed above. A single HIV-2 strain encoding glycine at the 119th CA residue was found in the Los Alamos databases; this strain (7312A) was isolated from a symptomatic 32-years-old man [33], and also was a recombinant between groups A and B (Fig. 1 and 2). This recombinant virus exhibits a genomic organization similar to that of NMC307, NMC716, and NMC842. At present, we do not know whether the emergence of glycine at the 119th position of CA is unique to HIV-2 CRF01_AB. It will be critical to assess the emergence of Gly119 viruses within HIV-2 groups A and B.

It is generally believed that HIV-2 is less pathogenic than HIV-1, and the number of HIV-2 cases is now gradually decreasing in West Africa. However, NMC307, NMC716, and NMC842 were recovered from patients at an advanced stage of AIDS with low CD4+ cell counts and high HIV-2 VLs [19]. It is possible that these HIV-2 CRF01_AB strains are highly pathogenic, unlike other HIV-2 strains. Careful epidemiological and virological studies are necessary to test this hypothesis. In the present study, we found that HIV-2 CRF01_AB CA confers strong resistance to human TRIM5α. In the Caio HIV-2 cohort in West Africa, non-proline residues at position 119 were significantly associated with elevated plasma HIV-2 load [18]. Therefore, resistance to TRIM5α may at least partially explain why these 3 patients in Japan developed AIDS so rapidly, although the possible effects of mutations in regions (e.g., nef, vif, and the long terminal repeats) other than those that encode CA cannot be fully excluded at
present. Our results also suggest that resistance to TRIM5α might be a new marker for the pathogenic potential of HIV-2. The possible emergence of a highly pathogenic HIV-2 strain is an ongoing concern, given that retroviruses can easily evolve to evade host defenses.

Materials and Methods
Phylogenetic Tree Analysis
Multiple sequence alignment was performed using the software CLUSTALW version 2.1. Phylogenetic trees were constructed using the neighbor-joining method. Bootstrap probabilities were calculated by 1000 iterations [34].

Cell Culture
The human 293T [35] and feline CRFK [36] cells were maintained in Dulbecco’s Modified Eagle medium. The human T-cell line CEM-SS [37] was maintained in RPMI medium. All media were supplemented with 10% fetal bovine serum and 1% penicillin-streptomycin.

Plasmid Construction
Recombinant HIV-2 GH123 clones containing the entire CA sequence of the isolates NMC716 or NMC842 (716 or 842, respectively) and 716 or 842 with proline substitutions at the 178th position (716GPP or 842GPP, respectively) were generated by PCR-based mutagenesis. The GH123/G virus was described previously [21]. The 0.6-kb HindIII-XhoI fragment of 842 was replaced with the corresponding fragment of GH123/G, and the resulting plasmid was designated 842Hind. Infectious viruses were prepared by transfection of 293T cells with the resulting proviral DNA clones. Viral titers were determined by measuring P25 (CA) with a RetroTek antigen ELISA kit (ZeptoMetrix, Buffalo, NY).

To construct the wild-type and mutant HIV-2 clones encoding GFP, the 1.6-kb KpnI-XhoI fragment (which encodes the MA, CA and p6) of GH123, 842, or GH123/G, was transferred to pROD-env(-)-GFP [25], a clone in which the env gene is disrupted, and the GFP gene was inserted into the nef region. Infectious viruses were prepared by transfection of 293T cells with proviral DNA clones together with the pMD2G plasmid encoding VSV-G. Viral titers were determined as above.

Construction of recombinant SeV encoding C-terminally HA-tagged CM TRIM5α (CM-TRIM5α-SeV), Rh TRIM5α (Rh-...
Figure 8. Structural models of the HIV-2 capsid C-terminal domain in dimeric form (A) and the HIV-2 GH123 capsid hexamer (B and C). (A) The C-terminal domain dimer model (from the amino acid position 150 to 219) of HIV-2 capsid (CA) is based on the viral sequence of NMC842. HIV-2 CRF01_AB-specific amino acid substitutions are shown in red. (B and C) The space-filling model of CA hexamer from the top (B) and side (C) is shown. Positions of HIV-2 CRF01_AB-specific amino acid substitutions are shown in red. L4/5 and 120P are shown in green and blue, respectively. doi:10.1371/journal.pone.0047757.g008
TRIM5α-SeV), human TRIM5α (Hu-TRIM5α-SeV), and CM TRIM5α lacking the PRYSFYR domain (CM-SPRY(−)-SeV) were described previously [10,15,22].

Viral Infection
CEM-SS cells (1×10⁶) were infected with SeVs encoding the respective TRIM5α proteins at a multiplicity of infection of 10 plaque-forming units per cell and incubated at 37°C for 9 h. Aliquots of 1×10⁷ cells were then superinfected with GH123, GH123/G, 716, 716GPP, 842, 842GPP, or 842Hind virus. Each superinfection used a titer of virus corresponding to 20 ng of p25 (CA). Experiment was performed three separate times with duplicate samples. For viral infection of cells producing physiological levels of TRIM5α, TRIM5α knock-down Jurkat cells (TRIM5α-KD Jurkat) and control cells (Luci-siRNA Jurkat) were infected with GH123, GH123/G, 716, or 842 virus. Each infection used a titer of virus corresponding to 100 ng of p25. The culture supernatants were collected periodically, and the level of p25 (CA) was measured as described above. Experiment was performed two separate times with duplicate samples.

Western Blot
CEM-SS cells (1×10⁶) infected with recombinant SeVs expressing HA-tagged TRIM5α proteins were lysed in lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Nonidet P40, 0.5% sodium deoxycholate). TRIM5α proteins in the lysates were subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis. Proteins in the gel were then electronically transferred onto a membrane (Immobilon; Millipore, Billerica, MA). Blots were blocked and probed with anti-HA high-affinity rat monoclonal antibody (Roche, Indianapolis, IN) overnight at 4°C. Each blot was then incubated with peroxidase-conjugated anti-rat IgG (Nacalai Tesque, Kyoto, Japan). Blots were visualized with a Chemilumi-One chemiluminescent kit (Nacalai Tesque, Kyoto, Japan).

Single-round Infection Assay
SeV-infected CRFK cells (4×10⁵) were infected with a titer of pROD-env(-)-GFP derivative virus corresponding to 500 ng of p25 (CA). Two days after infection, the cells were fixed by formaldehyde, and GFP-producing cells were counted by flow cytometry. Experiment was performed three separate times with triplicate samples.

Molecular Modeling and MD Simulation
We used molecular dynamic (MD) simulations [38] to analyze the structural dynamics of the HIV-2 CA NTDs. First, initial CA structures for MD simulation were constructed by homology modeling [39] using the Molecular Operating Environment, MOE (Chemical Computing Group Inc., Montreal, Canada) as described previously [15,40]. We used the high-resolution crystal structure of the HIV-2 CA CTD at a resolution of 1.70 Å (PDB code: 1A8O) [17]. The amino acid sequence identity of HIV-1 (1A8O) and HIV-2 CA (NMCG42 in this study) is about 76%. The sequence similarity is sufficient to construct a structural model with an r.m.s. deviation of approximately 1.5 Å for the main chain between the predicted and actual structures [39]. The 3-D structures were optimized thermodynamically by energy minimization using MOE and an AMBER99 force field [44] and further refined the physically unacceptable local structures on the basis of evaluation of unusual dihedral angles, phi and psi, by the Ramachandran plot using MOE. The binding energies of the CA dimer models, Ebind, were calculated as described elsewhere [45,46], using the formula $E_{\text{bind}} = E_{\text{dimer}} - 2E_{\text{monomer}}$, where $E_{\text{dimer}}$ is the energy of the CA dimer; $E_{\text{monomer}}$ is the energy of the CA monomer.

Conclusions
The CA of HIV-2 CRF01_AB isolates have a unique feature distinct from that of other HIV-2 strains; CRF01_AB-specific amino acid substitutions in the CA CTD were necessary for strong resistance to human TRIM5α.

Supporting Information

Figure S1 Western blot analysis of TRIM5α proteins.
HA-tagged TRIM5α proteins in lysate of CEM-SS cells infected with recombinant SeV were visualized by western blotting with an antibody against HA. S(-), Hu, CM, and Rh denote CM SPRY(−), human, cynomolgus monkey, and rhesus TRIM5α, respectively. Molecular weight makers are shown on the left. (EPS)

Acknowledgments
We thank Dr. Y. Tian, Dr. S. Nakamura and Dr. T. Yasunaga for helpful discussions, and Ms. S. Bando and Ms. N. Teramoto for assistance.

Author Contributions
Conceived and designed the experiments: TM EEN MY ST KK. Analyzed the data: TM EEN MY SI HS TS. Contributed reagents/materials/analysis tools: SI KK JL WS HS TS. Performed the experiments: TM EEN MY ST KK. Wrote the paper: TM EEN MY SI JL HS TS.

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