APOBEC3 degradation is the primary function of HIV-1 Vif for virus replication in the myeloid cell line THP-1

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Running title: A3 proteins are Vif targets for HIV-1 replication (49/54 characters)

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Abstract

HIV-1 must overcome multiple innate antiviral mechanisms to replicate in CD4\(^+\) T lymphocytes and macrophages. Previous studies have demonstrated that the APOBEC3 (A3) family of proteins (at least A3D, A3F, A3G, and stable A3H haplotypes) contribute to HIV-1 restriction in CD4\(^+\) T lymphocytes. Virus-encoded virion infectivity factor (Vif) counteracts this antiviral activity by degrading A3 enzymes allowing HIV-1 replication in infected cells. In addition to A3 proteins, Vif also targets other cellular proteins in CD4\(^+\) T lymphocytes, including PPP2R5 proteins. However, whether Vif primarily degrades only A3 proteins or has additional essential targets during viral replication is currently unknown.

Herein, we describe the development and characterization of A3F-, A3F/A3G-, and A3A-to-A3G-null THP-1 cells. In comparison to Vif-proficient HIV-1, Vif-deficient viruses have substantially reduced infectivity in parental and A3F-null THP-1 cells, and a more modest decrease in infectivity in A3F/A3G-null cells. Remarkably, disruption of A3A–A3G protein expression completely restores the infectivity of Vif-deficient viruses in THP-1 cells. These results indicate that the primary function of Vif during HIV-1 replication in THP-1 cells is the targeting and degradation of A3 enzymes.
**Importance**

HIV-1 Vif neutralizes the HIV-1 restriction activity of A3 proteins. However, it is currently unclear whether Vif has additional essential cellular targets. To address this question, we disrupted A3A to A3G genes in the THP-1 myeloid cell line using CRISPR and compared the infectivity of wildtype HIV-1 and Vif mutants with the selective A3 neutralization activities. Our results demonstrate that the infectivity of Vif-deficient HIV-1 and the other Vif mutants is fully restored by ablating the expression of cellular A3A to A3G proteins. These results indicate that A3 proteins are the only essential target of Vif that is required for HIV-1 replication in THP-1 cells.
Introduction

The apolipoprotein B mRNA editing enzyme polypeptide-like 3 (APOBEC3, A3) family of proteins comprise seven single-strand DNA cytosine deaminases (A3A–A3D and A3F–A3H) in humans (1-3). A3 enzymes have broad and essential roles in innate antiviral immunity against parasitic DNA-based elements (4-6). Retroviruses are sensitive to A3 enzyme activity due to the obligate step of reverse transcription during viral replication that produces single-stranded cDNA intermediates. These viral cDNA intermediates can act as substrates for A3 enzymes, as demonstrated by C-to-U deamination resulting in G-to-A mutations in the genomic strand. To date, the best-characterized substrate of A3 enzymes is human immunodeficiency virus type 1 (HIV-1). In CD4+ T lymphocytes, four A3 proteins (A3D, A3F, A3G, and stable A3H haplotypes) restrict HIV-1 replication by mutating viral cDNA intermediates and by physically blocking reverse transcription (7-14). A3 enzymes have a preference for specific dinucleotide motifs (5′-CC for A3G and 5′-TC for other A3 enzymes) at target cytosine bases, which appear as 5′-AG or 5′-AA mutations in the genomic strand (7, 8, 15, 16).

Virus-encoded virion infectivity factor (Vif) functions in disrupting the activity of A3 enzymes. Vif forms an E3 ubiquitin ligase complex that degrades A3 enzymes through a proteasome-mediated pathway (2, 3, 17, 18). The central domain of this complex is a Vif heterodimer with the transcription factor, CBF-β, which stabilizes Vif during disruption of A3 protein activity (19, 20). Vif also suppresses the transcription of A3 enzymes by hijacking RUNX/CBF-β complex (21). In addition to these Vif-dependent mechanisms, HIV-1 reverse transcriptase and protease have been shown to disrupt the activity of A3 enzymes via Vif-independent mechanisms (22, 23). Recently, functional proteomic
analyses have demonstrated that Vif has several target proteins, including the PPP2R5 family of proteins, in CD4+ T cell lines and lymphocytes (24, 25). These findings indicate that Vif may have additional essential target proteins during HIV-1 replication in infected cells.

We previously reported that endogenous A3G protein contributes to HIV-1 restriction in a deaminase-dependent manner in THP-1 cells (26). Although disruption of the A3G gene nearly eliminates viral G-to-A mutations, Vif-deficient HIV-1 virions have 50% lower infectivity than wildtype HIV-1 or mutants selectively lacking A3G degradation activity (26). These results indicated that Vif-mediated inhibition of A3G and at least one additional A3 protein is required for efficient HIV-1 replication.

In the present study, we evaluate the effects of other A3 proteins on HIV-1 infectivity by developing and characterizing A3F-, A3F/A3G-, and A3A-to-A3G-null THP-1 cells using HIV-1 Vif mutants with selective A3 neutralization activities. In comparison to wildtype HIV-1, Vif-deficient HIV-1 infectivity is strongly inhibited in A3F-null THP-1 cells and modestly inhibited in A3F/A3G-null THP-1 cells. In contrast, an HIV-1 Vif mutant selectively lacking A3F degradation activity had comparable infectivity to wildtype HIV-1 in A3F-null THP-1 cells and 50% infectivity in parental THP-1 cells, indicating that A3F protein contributes to HIV-1 restriction in THP-1 cells. Furthermore, Vif-deficient HIV-1 infectivity is comparable to wildtype HIV-1 in A3A-to-A3G-null THP-1 cells. These results demonstrate that A3 proteins are the primary target of HIV-1 Vif during virus replication in THP-1 cells.

Results
Endogenous A3H is not involved in HIV-1 restriction in THP-1 cells.

THP-1 cells express significant levels of A3B, A3C, A3F, A3G, and A3H mRNA (26). The results of our previous study indicated that A3G and at least one additional A3 protein are involved in HIV-1 restriction in THP-1 cells (26). Variations in the amino acid sequence of A3 family proteins are known to influence HIV-1 restriction activity (27), and the A3H gene is the most polymorphic of all human A3 genes (10, 22, 28, 29). The A3H allele is grouped into stable and unstable haplotypes according to the combination of amino acid residues at positions 15, 18, 105, 121, and 178 (10, 22, 28, 29). Stable A3H haplotypes are active against HIV-1 whereas unstable A3H haplotypes have absent or minimal activity as they encode proteins with low stability (9, 10, 22, 29, 30). To determine A3H genotypes, we sequenced A3H cDNA from THP-1 cells. Sequencing data identified an unstable haplotype in the THP-1 genome, termed A3H hapI (Fig. 1A). These data suggest that endogenous A3H protein has minimal restriction activity against Vif-deficient HIV-1 in THP-1 cells.

The A3H hapI results in expression of an unstable protein that has weak anti-HIV-1 activity (28, 29, 31). However, this protein is enzymatically active and has an HIV-1 restriction phenotype similar to the stable A3H haplotype, A3H hapII, when both proteins are expressed at the same levels (31). In addition, A3H protein expression levels are upregulated during HIV-1 infection (10, 22), and A3H hapI is resistant to Vif-mediated degradation (32). Accordingly, we evaluated whether the expression of A3H hapI is associated with HIV-1 restriction in THP-1 cells. To address this question, we utilized HIV-1 Vif mutants that selectively degrade stable A3H (hyper-functional Vif; hyper-Vif) or lack stable A3H degradation (hypo-functional Vif; hypo-Vif) (Fig. 1B). IIIB Vif displays an
intermediate phenotype (Fig. 1B). Of note, hyper-Vif, hypo-Vif, and IIIB Vif have full neutralization activity against A3D, A3F, and A3G proteins (10). VSV-G pseudotyped HIV-1 Vif mutants were produced from HEK293T cells and infected into SupT11 and THP-1 cells to create virus-producing cells (see Pseudo-single cycle infectivity assays in Material & Methods). The produced viruses were then used to measure viral infectivity in TZM-bl cells, evaluate packaging of A3 proteins by western blotting, and analyze the frequency of G-to-A mutations. As shown in Fig. 1C (top panel), hyper-Vif HIV-1, hypo-Vif HIV-1, and IIIB Vif HIV-1 (IIIB) produced in THP-1 cells had similar viral infectivity. While Vif did not degrade A3H protein in THP-1 cells, it was not packaged into viral particles (Fig. 1C, bottom panel). Next, to determine whether G-to-A mutations were introduced into proviral DNA, we recovered proviral DNA from SupT11 cells after infection with each HIV-1 mutant produced from THP-1 cells and sequenced the pol region of these proviruses. Sequencing data demonstrated that hyper-Vif HIV-1, hypo-Vif HIV-1, and IIIB Vif HIV-1 had minimal G-to-A mutations preferred by A3H protein (GA-to-AA signature motif) in proviral DNA (Fig. 1D and E), indicating that endogenous A3H protein expressed in THP-1 cells is not involved in HIV-1 restriction. In contrast, the replication of Vif-null HIV-1 was restricted in THP-1 cells and A3G, the major HIV-1 restrictive A3 protein, was packaged in viral particles, thereby inducing profound G-to-A mutations (10.3 ± 3.5 mutations/kb). Most of mutations were in the GG-to-AG signature motif preferred by A3G (80 ± 10%) in proviral DNA (Fig. 1C-E). The susceptibility of Vif mutants to stable A3H protein was confirmed in SupT11 cells stably expressing stable A3H protein (Fig. 1C-E). Taken together, these results indicate that A3G and other A3 proteins, except A3H, contribute to HIV-1 restriction in THP-1 cells.

A3F protein has a restrictive effect on HIV-1 among A3 family members and is a target of Vif, in addition to A3G, in CD4+ T cell lines and lymphocytes (7, 33-35). To determine whether A3F protein also reduces HIV-1 infectivity in THP-1 cells, we used CRISPR to create A3F and A3F/A3G gene knockout cell lines. Two independent subclones of A3F and A3F/A3G-null THP-1 cells were obtained, as evidenced by the results of genomic DNA sequencing and western blotting (Fig. S1 and S2).

A3 proteins include single- and double-domain deaminases, which are phylogenetically classified into three groups: Z1, Z2, and Z3 domains (3, 4) (Fig. 2A represented in green, yellow, and blue, respectively). A3A, A3B carboxy-terminal domain (CTD), and A3G CTD proteins are classified as Z1 domains (Fig. 2A; represented in green). Of note, exon 4 of the A3A gene, exon 7 of the A3B gene, and exon 7 of A3G gene are highly conserved at the nucleotide level (A3A exon 4 and A3B exon 7 have 95% identity; A3A exon 4 and A3G exon 7 have >99% identity; and A3B exon 7 and A3G exon 7 have 95% identity, respectively). Interestingly, each of these exons has an identical sequence (5′-GAG TGG GAG GCT GCG GGC CA). We therefore designed a guide RNA (gRNA) homologous to this sequence and attempted to delete the entire 125 kbp interval spanning A3A to A3G in THP-1 cells (Fig. 2A; represented in arrows, and S3). We predicted that successful deletion would cause one of the following three scenarios: 1) fusion of exon 4 of the A3A gene with exon 7 of the A3B gene (30 kbp deletion); 2) fusion of exon 7 of the A3B gene with exon 7 of the A3G gene (95 kbp deletion); or 3) fusion of exon 4 of the A3A gene with exon 7 of the A3G gene (125 kbp deletion).
deletion; Fig. 2A). To obtain THP-1 cells lacking expression of A3A to A3G protein, a lentiviral vector expressing gRNA against the target sequence was transduced into THP-1 cells. Finally, two independent subclones (THP-1#11-4 and THP-1#11-7) were obtained, with whole genome sequencing (WGS) analysis demonstrating an extensive deletion between A3A exon 4 and A3G exon 7 at the A3 gene locus (Fig. 2B). In THP-1#11-4, six alleles of the fusion of A3A exon 4 with A3G exon 7 are observed, and each A3A/A3G hybrid exon had six different insertions or deletions (indels) (Fig. S3). THP-1#11-7 harbors three alleles of A3A exon 4 and A3G exon 7 fusions (one may be A3A exon 4) with three different deletions (Fig. S3). Although more than 20 potential off-target sites with two or three nucleotides mismatched with the designed gRNA were predicted, a significant deletion was only found downstream of the predicted A3G pseudogene harboring 2 bp mismatched with the target sequence (Fig. S4; potential target sequence in a yellow box and deletions indicated by green dotted lines). In comparison to parental THP-1 cells, these subclones had similar growth capacities under normal cell culture conditions. RT-qPCR analyses demonstrated that A3B to A3G mRNA is not detectable in either clone (Fig. 2C). However, A3A mRNA expression remained detectable in parental THP-1 cells and the two subclones as the A3A promoter remains intact and potentially functional (Fig. 2A-C). A3A mRNA expression is known to be upregulated 100–1000-fold in THP-1 cell treated with type I interferon (IFN) (36). To confirm the expression of A3A mRNA and protein in THP-1 cells, parental THP-1 cell and the respective subclones were cultured in the presence of type I IFN for 6 hours, and A3 mRNA and protein expression levels were then analyzed by RT-qPCR and western blotting, respectively. In parental THP-1 cells, A3A, A3B, A3F, and A3G mRNA and...
protein expression levels were increased following IFN treatment (Fig. 2C and D). In the THP-1#11-4 subclone, A3A mRNA expression is increased following IFN treatment; however, A3A, A3B, A3C, A3F, and A3G proteins are not detectable, even after IFN treatment (Fig. 2C and D). Further, A3A to A3G proteins are not detectable in the THP-1#11-7 subclone under normal cell culture conditions (Fig. 2D). Interestingly, low levels of a protein with comparable size to A3A are detected in the THP-1#11-7 subclone after IFN treatment (Fig. 2D). Sanger sequence analyses indicated that this protein was an A3A and A3G hybrid with a 3-bp deletion (Fig. S3). Collectively, these data indicate that the THP-1#11-4 and THP-1#11-7 subclones lack expression of A3A to A3G proteins under normal cell culture conditions and that clone THP-1#11-4 is a clean knockout that fails to express functional versions of any of these proteins.

Disruption of A3A to A3G protein expression fully restores the infectivity of Vif-deficient HIV-1 in THP-1 cells.

We next determined whether endogenous A3F protein is degraded by Vif in addition to A3G. HIV-1 Vif mutants with selective A3 neutralization activities were used for pseudo-single cycle infectivity assays as mentioned above. For example, a Vif4A mutant harboring \(^{14}\)AKTK\(^{17}\) substitutions (\(^{14}\)DRMR\(^{17}\) in IIIB) is susceptible to A3D and A3F activity but resistant to A3G activity (37-39) (Fig. 3A). We examined the ability of Vif4A to counteract the activity of A3F as A3D mRNA expression level is relatively low in THP-1 cells (26) (Fig. 2C). As our group and others have previously shown (26, 37, 38, 40), Vif5A containing five alanine substitutions (\(^{40}\)YRHHY\(^{44}\) to \(^{40}\)AAAAA\(^{44}\)) is sensitive to A3G restriction but not the activity of A3D or A3F (Fig. 3A). Vif4A5A is susceptible to A3D,
A3F, and A3G (37) (Fig. 3A). VSV-G pseudotyped HIV-1 and these Vif mutants were used to infect SupT11 derivatives and engineered A3F-null THP-1 cells. First, the susceptibilities of these Vif mutants to A3F and A3G proteins were validated in SupT11 cell lines (Fig. 3B). In SupT11-vector cells, Vif-proficient HIV-1 and all Vif mutants had comparable infectivity in TZM-bl cells (Fig. 3B). As expected, the infectivity of Vif-deficient HIV-1 and the Vif4A and 4A5A mutants was reduced in SupT11-A3F cells as these mutants are unable to degrade A3F protein, thereby leading to packaging of A3F protein in viral particles (Fig. 3B). Further, infection with Vif-deficient HIV-1 or the Vif5A and Vif4A5A mutants resulted in packaging of A3G protein in viral particles from SupT11-A3G cells in addition to reduced infectivity of these Vif mutants (Fig. 3B). These results are consistent with previous reports demonstrating the susceptibilities of Vif mutants to A3 proteins (26, 37-40).

Pseudo-single cycle infectivity assays were then performed in parental THP-1, A3G-null, and A3F-null cells using these Vif mutants. Vif-proficient HIV-1 degraded A3F and A3G proteins in THP-1 cells, and lower amounts of these A3 proteins were packaged into viral particles (Fig. 3C; THP-1 parent). In contrast, Vif-deficient HIV-1 was unable to degrade A3F and A3G proteins, thereby leading to reduced viral infectivity compared to Vif-proficient HIV-1 (Fig. 3C; THP-1 parent). The infectivity of A3F-susceptible Vif mutants, Vif4A and Vif4A5A, was lower than that of Vif-proficient HIV-1, indicating that endogenous A3F protein contributes to Vif-deficient HIV-1 restriction in THP-1 cells (Fig. 3C; THP-1 parent). This finding was supported by results in A3G-null THP-1 cells where Vif4A mutants are restricted, as observed in parental THP-1 cells (Fig. 3C; THP-1 ΔA3G). The involvement of endogenous A3G protein in HIV-1 restriction was confirmed in A3G-
null THP-1 cells, as reported (26) (Fig. 3C; THP-1 ∆A3G). To determine whether endogenous A3F protein contributes to HIV-1 restriction in THP-1 cells, pseudo-single cycle infectivity assays were performed according to the methods described above in two independent A3F-null THP-1 clones (Fig. S1). Vif-deficient HIV-1 and the Vif5A and Vif4A5A mutants had reduced infectivity in A3F-null subclones due to the inhibitory effect of A3G (Fig. 3C; THP-1 ∆A3F#1 and #2). However, the infectivity of the Vif4A mutant was restored to near wildtype levels following disruption of A3F expression in THP-1 cells. These data demonstrate that endogenous A3F protein contributes to Vif-deficient HIV-1 restriction in THP-1 cells, and that Vif degrades A3F and thereby prevents packaging and restriction upon target cell infection.

A3F and A3G proteins are involved in Vif-deficient HIV-1 restriction in THP-1 cells and are degraded by Vif (26) (Fig. 3C). However, it is unclear whether only these A3 proteins are associated with Vif-deficient HIV-1 restriction in THP-1 cells. To address this issue, we performed pseudo-single cycle infectivity assays in A3F/A3G-null THP-1 cells using separation-of-function Vif mutants. Although Vif-deficient HIV-1 had greater infectivity defects in parental, A3G-null, and A3F-null THP-1 cells compared to wildtype HIV-1 (parent: <10% infectivity, ∆A3G: 30 to 40% infectivity, and ∆A3F: 20% infectivity, respectively), the infectivity of Vif-deficient HIV-1 was 30% lower in A3F/A3G-null THP-1 cells (Fig. 3C; THP-1 parent, ∆A3G, ∆A3F#1 and #2, and ∆A3F/A3G#1 and #2). On the other hand, the Vif4A, Vif5A, and Vif4A5A mutants had similar infectivity to wildtype HIV-1 in A3F/A3G-null THP-1 cells (Fig. 3C; THP-1 ∆A3F/A3G#1 and #2). These data indicate that other A3 proteins, in addition to A3F and A3G, contribute to Vif-deficient HIV-
1 restriction in THP-1 cells or that Vif disrupts an additional essential target during viral replication in THP-1 cells.

The universally recognized primary target of Vif is the A3 family of proteins (2, 3, 17, 18). However, Vif-mediated A3 degradation may mask an additional A3-independent Vif function required for viral replication. To address this issue, we constructed two independent A3A-to-A3G-null THP-1 clones (Fig. 2) and characterized HIV-1 infection using pseudo-single cycle infectivity assays with Vif mutants. As mentioned above, the disruption of A3F and A3G protein expression results in Vif-deficient HIV-1 having 70% of wildtype HIV-1 infectivity in THP-1 cells (Fig. 3C; THP-1A3F/A3G#1 and #2). Remarkably, Vif-deficient HIV-1 and the other Vif mutants have comparable infectivity to Vif-proficient HIV-1 lacking expression of A3A-to-A3G in THP-1 cells (Fig. 3C; THP-1#11-4 and #11-7). These results indicate that A3 degradation is the only function of Vif required for viral replication in THP-1 cells.

A3 proteins restrict HIV-1 replication via deaminase-dependent and deaminase-independent mechanisms in THP-1 cells.

Our previous results indicated that A3G protein is the primary source of A3 mutagenesis in THP-1 cells (26). To further investigate the G-to-A mutation spectra in each A3-null THP-1 subclone, the pol region was cloned and sequenced from the proviruses used in the aforementioned infectivity assays. As expected, GG-to-AG mutations are observed in the proviral DNA of Vif mutants lacking A3G neutralization activity (Vif-deficient HIV-1 and Vif5A and Vif4A5A mutants) produced from SupT11-A3G cells (Fig. 4A-B; SupT11-A3G). Consistent with a previous report (26), THP-1 expresses A3G protein capable of mutating
A3G-susceptible Vif mutants, including Vif-deficient HIV-1 and Vif5A and Vif4A5A mutants, as seen in parental THP-1 cells. These GG-to-AG mutations are not observed in A3G-null THP-1 cells (Fig. 4A-B; THP-1 parent and ∆A3G). Similarly, GG-to-AG mutations preferred by A3G were seen in the proviruses of the A3G-susceptible Vif mutants produced from two independent A3F-null THP-1 cells, with disruption of A3G nearly completely eliminating these mutations in THP-1 cells (Fig. 4A and B; THP-1ΔA3F#1 and #2, ΔA3F/A3G#1 and #2, #11-4, and #11-7). These data indicate that A3G protein is the primary source of G-to-A mutations in HIV-1 proviruses produced by THP-1 cells.

Although the Vif mutants lacking A3F neutralization activity (Vif-deficient HIV-1 and Vif4A and Vif4A5A mutants) produced from SupT11-A3F cells have a relatively low number of G-to-A mutations, the observed G-to-A mutations are predominantly within the GA-to-AA sequence motif preferred by A3F (Fig. 4A-B; SupT11-A3F). However, A3F-preferred GA-to-AA mutations are not observed in proviruses of A3F-susceptible Vif mutants produced from parental or A3G-null THP-1 cells, in support of prior observations (26) (Fig. 4A-B; THP-1 parent and ∆A3G). In addition, fewer GA-to-AA mutations are observed in THP-1 cells, even after disruption of A3F protein expression (Fig. 4A-B; THP-1ΔA3F#1 and #2, ΔA3F/A3G#1 and #2, #11-4, and #11-7). Accordingly, these results combine to indicate that A3F protein in THP-1 cells is involved in Vif-deficient HIV-1 restriction via a deaminase-independent mechanism.

A3F protein has been shown to inhibit the accumulation of reverse transcription (RT) products (14). To investigate a potential effect on RT, SupT11 cells were infected with viruses from the pseudo-single cycle infectivity assays described above, and late RT
(LRT) products were examined by quantitative PCR (qPCR). As expected, all Vif mutants were decreased in LRT products in comparison to wildtype virus when these mutants were produced in parental THP-1 cells and used to infect SupT11 cells (Fig. 4C; THP-1 parent). LRT products of Vif5A and Vif4A mutants were restored to levels comparable to Vif-proficient HIV-1 following the disruption of A3G or A3F protein expression in THP-1 cells (Fig. 4C; THP-1 ∆A3G, and ∆A3F#1 and #2), indicating that both A3G and A3F proteins inhibit HIV-1 via a deaminase-independent mechanism. However, double knockout of A3G and A3F in THP-1 cells did not increase the LRT products of Vif-deficient HIV-1 compared to those of Vif-proficient virus (Fig. 4C; THP-1 ∆A3F/A3G#1 and #2), indicating other A3 proteins, in addition to A3F and A3G, may contribute to the restriction of HIV-1 in THP-1 cells via a deaminase-independent mechanism or that a separate protein targeted by Vif blocks the accumulation of RT products. To test this hypothesis, we measured LRT products by infecting SupT11 cells with HIV-1 Vif mutants produced in A3A-to-A3G-null clones. Consistent with the results of the pseudo-single cycle infectivity assays (Fig. 3C), Vif-deficient HIV-1 and other Vif mutants had comparable levels of LRT products to Vif-proficient HIV-1 lacking expression of A3A to A3G protein in THP-1 cells (Fig. 4C; THP-1#11-4 and #11-7). These data indicate that Vif-mediated A3 degradation is required for viral replication in THP-1 to counteract deaminase-dependent and -independent HIV-1 restriction by A3 proteins.

Transmitted/founder (TF) HIV-1 Vif also only targets A3 family proteins to enable virus replication in THP-1 cells.
We finally examined whether the A3-dependent function of Vif was present in TF viruses. To address this issue, Vif-proficient and deficient versions of the CH58 TF virus were produced from parental THP-1 and A3A-to-A3G-null cells, with viral infectivity measured in TZM-bl cells (Fig. 5). Similar to the results observed with IIIB viruses, Vif-deficient CH58 virus was restricted in parental THP-1 cells; however, this restriction is completely abolished by disruption of the A3A to A3G genes (Fig. 5). These data indicate that TF viruses also utilize a primarily A3-dependent function of Vif during replication in THP-1 cells.

Discussion

Vif-mediated A3 degradation is critical for HIV-1 replication in CD4⁺ T lymphocytes and myeloid cells (2, 3, 17, 18). In CD4⁺ T lymphocytes, at least A3D, A3F, A3G, and A3H (only stable haplotypes) are involved in Vif-deficient HIV-1 restriction, and Vif is required to degrade A3 enzymes and allow efficient viral replication (2, 3, 17, 18). However, the degradation of A3 enzymes by Vif during HIV-1 replication in myeloid lineage cells has yet to be fully elucidated. We previously reported that A3G protein contributes to Vif-deficient HIV-1 restriction in a deaminase-dependent manner in THP-1 cells (26). Herein, we demonstrate that A3F protein also inhibits Vif-deficient HIV-1 in a largely deaminase-independent manner and that Vif avoids this HIV-1 restriction mechanism by degrading A3F protein (Fig. 3-4). Importantly, the results of pseudo-single cycle infectivity assays demonstrate that the disruption of A3A to A3G protein confers comparable infectivity to wildtype HIV-1 in a Vif-deficient lab-adapted virus (IIIB) and TF virus (CH58) (Fig. 3-5).
These results indicate that Vif-mediated A3 degradation is the primary function of Vif during HIV-1 replication in THP-1 cells. Our results demonstrate that A3F and A3G but not A3H proteins restrict Vif-deficient HIV-1 via deaminase-dependent and -independent mechanisms in THP-1 cells (Fig. 1, 3 and 4). In addition to A3F and A3G proteins, our findings indicate that at least one additional A3 protein is involved in Vif-deficient HIV-1 restriction via a deaminase-independent mechanism (Fig. 3-4). Accordingly, the remaining four A3 proteins (A3A, A3B, A3C, and A3D) may contribute to Vif-deficient HIV-1 restriction in a deaminase-independent manner in THP-1 cells (Fig. 4). However, A3A and A3B are highly unlikely to contribute in this manner as A3A mRNA and protein expression levels are very low or undetectable in THP-1 cells without IFN treatment (Fig. 2C-D). Further, both A3A and A3B are resistant to degradation by HIV-1 Vif (7, 34, 41-43). It is therefore plausible that A3C and A3D proteins contribute to Vif-deficient HIV-1 restriction in THP-1 cells. An A3C-isoleucine 188 variant is reportedly more active against HIV-1 than a serine 188 variant (44, 45). To ask which A3C variant is expressed by THP-1 cells, we determined the A3C genotypes of THP-1 cells using cDNA sequencing. These results demonstrated that the amino acid residue of A3C at position 188 is serine. This result indicates that A3C has a modest effect on Vif-deficient HIV-1 restriction via a deaminase-independent mechanism in THP-1 cells, consistent with prior studies (45). Similarly, the results of previous studies indicate that A3D has a weak effect on Vif-deficient HIV-1 restriction in HEK293, SupT11, and CEM2n cells (7, 8, 37, 46, 47). Nevertheless, the fact that Vif-deficient HIV-1 has 20% lower infectivity indicates that a synergistic mechanism may enhance the effect of
A3 proteins on HIV-1 infectivity (48, 49). Further studies are required to fully elucidate the mechanisms underlying the effect of A3 proteins on HIV-1 infectivity.

Similar to CD4+ T lymphocytes, HIV-1 can also target myeloid cells such as monocytes and macrophages, and these infections are associated with viral dissemination, persistence, and latency (50, 51). Accordingly, it is important to understand the role of restriction factors, including A3 proteins, in myeloid cells. In monocytes, A3A mRNA levels are 10–1000 times higher than other A3 mRNA expression levels, and A3A mRNA expression is reduced by 10–100-fold after differentiation into monocyte-derived macrophages (MDMs) (52-54). In contrast, A3G mRNA expression levels are reduced approximately 10-fold lower after differentiation of monocytes into MDMs (52, 53). A3F mRNA expression levels are less variable during the differentiation of monocytes into MDMs (52). Interestingly, suppression of A3A and A3G protein levels by siRNA reportedly leads to a 4–5-fold increase in p24 production by HIV-1-infected monocytes (53). As MDMs are generally more sensitive to HIV-1 infection than monocytes, it is highly likely that A3A and A3G contribute to the susceptibility of MDMs to HIV-1 infection. However, as previous studies have reported that A3A is less active against HIV-1 in HEK293T and SupT11 cell lines (7, 34, 55), further studies are required to determine the contribution of A3A to HIV-1 restriction in monocytes.

In addition to A3A and A3G, A3F and A3H may be involved in HIV-1 restriction in monocytes. Although A3F mRNA expression levels are essentially unchanged during differentiation from monocytes into MDMs (53), A3F mRNA expression levels are comparable to A3G mRNA expression levels (53, 54), indicating that A3F protein likely contributes to HIV-1 restriction in monocytes. It is possible that only stable A3H
haplotypes and A3C-I188 are associated with HIV-1 restriction in monocytes. According to previous observations in HEK293, SupT11, and CEM2n cells (7, 8, 37, 46, 47), A3D may modestly contribute to HIV-1 restriction in monocytes. As A3B mRNA expression levels are relatively low, it is unlikely that this A3B inhibits HIV-1 in monocytes. However, the contribution of A3 proteins other than A3A and A3G to HIV-1 suppression in monocytes remains unclear, and the antiviral activities of these A3 proteins warrant further investigation.

In MDMs, A3A appears to be associated with anti-HIV-1 activity as increasing HIV-1 infectivity has been reported following siRNA knockdown of A3A (53, 54). In addition, HIV-1 replication assays in MDMs using HIV-1 Vif4A and Vif5A mutants demonstrated that the replication kinetics of both mutants were slower than that of the Vif-proficient HIV-1, indicating that A3D, A3F, and A3G contribute to HIV-1 restriction in MDMs (39). However, the effects of A3D and A3F on HIV-1 replication are donor-dependent, likely due to their respective expression levels (39). As the antiviral activity of A3B, A3C, and A3H proteins has not been reported in MDMs, further studies are required to address these issues.

Vif is required for HIV-1 replication in CD4+ T lymphocytes and macrophages (2, 3, 17, 18). In the absence of Vif, HIV-1 is attacked by A3 proteins in CD4+ T lymphocytes, macrophages, monocytes, dendritic cells, and CD4+ T cell lines, and massive G-to-A mutations accumulate in HIV-1 proviral DNA (7, 8, 10, 15, 23, 26, 39, 56, 57). HIV-1 Vif recruits A3 proteins into an E3 ubiquitin ligase complex, thereby avoiding the antiviral activity of these proteins by promoting their degradation through a proteasome-mediated pathway (2, 3, 17, 18). The primary function of Vif has long been posited to be the
suppression of the antiviral activity of A3 proteins. On the other hand, Vif causes G2/M cell cycle arrest (58-60). As the amino acid residues of Vif responsible for G2/M cell cycle arrest do not completely match with the amino acid residues required for Vif-mediated A3 degradation, these functions of Vif may be independent of each other (61-63). In 2016, a functional proteomic analysis identified the PPP2R5 family of proteins, which function as regulators of protein phosphatase 2A (PP2A), as novel targets of Vif (25). Subsequently, Salamango et al. revealed that Vif induces G2/M arrest by degrading PPP2R5 proteins (60). Vif-induced G2/M arrest has been observed in many cell types, including HEK293T, SupT11, CEM-SS, and THP-1 cells and CD4+ T lymphocytes (25, 61, 63). However, Vif-mediated G2/M arrest is not required for HIV-1 replication, supporting our findings that A3 family proteins are the sole essential substrate of Vif during viral replication in THP-1 cells under normal cell culture conditions (Fig. 3-5). It has recently been reported that fragile X mental retardation 1 (FMR1) and diphthamide biosynthesis 7 (DPH7) are degraded by Vif in CD4+ T lymphocytes (24). Further studies are required to determine whether a substrate of Vif other than A3 proteins is required for HIV-1 replication in vivo.

In summary, the findings of the present study demonstrate that the primary target of Vif is the A3 family of proteins during HIV-1 replication in THP-1 cells. Whether this observation is applicable to primary CD4+ T lymphocytes and myeloid cells, such as monocytes and macrophages, is important for the development of antiviral therapies targeting the A3-Vif axis. Such studies may contribute to a functional cure for HIV-1 by manipulating A3 mutagenesis.
Material & Methods

Cell lines and culture conditions

HEK293T (CRL-3216) was obtained from American Type Culture Collection. TZM-bl (#8129) (64) was obtained from the NIH AIDS Reagent Program (NARP). The creation and characterization of the permissive T cell line SupT11 and the SupT11 single clones stably expressing untagged A3 (SupT11-vector, -A3F, -A3G and -A3H hapII high) have been reported (10, 33). CEM-GXR (CEM-GFP expressing CCR5) was provided by Dr. Todd Allen (Harvard University, USA) (65). THP-1 was provided by Dr. Andrea Cimarelli (INSERM, France) (53). The generation and characterization of THP-1 ΔA3G#1 have been reported (26). Adherent cells were cultured in DMEM (Wako, Cat# 044-29765) supplemented with 10% fetal bovine serum (FBS) (NICHIREI, Cat#175012) and 1% penicillin/streptomycin (P/S) (Wako, Cat# 168-23191). Suspension cells were maintained in RPMI (Thermo Fisher Scientific, Cat# C11875500BT) with 10% FBS and 1% P/S.

Genotyping of A3C and A3H genes

Total RNA was isolated from THP-1 by RNA Premium Kit (NIPPON Genetics, Cat# FG-81250). Then, cDNA was synthesized by Transcriptor Reverse Transcriptase (Roche, Cat# 03531287001) and used to amplify A3C or A3H gene with the following primers [A3C outer primers: (5’-GCG CTT CAG AAA AGA GTG GG) and (5’-GGA GAC AGA CCA TGA GGC). A3C inner primers: (5’-ACA TGA ATC CAC AGA TCA GAA A) and (5’-CCC TCT ACT GGA GAC TCT CC). A3H outer primers: (5’-CCA GAA GCA CAG ATG AGG CAA). A3H inner primers: (5’-TGT TAA CAG CCG AAA CAT TCC) and (5’-TCT TGA GTT GCT TCT TGA TAA T)]. The
amplified fragments were cloned into the pJET cloning vector (Thermo Fisher Scientific, Cat# K1231). At least 10 independent clones were subjected to Sanger sequencing (AZENTA) and sequence data were analyzed by Sequencher v5.4.6 (Gene Codes Corporation).

Construction of pLentiCRISPR-Blast

The pLentiCRISPR1000 system was previously described (66). pLentiCRISPR1000-Blast was generated by restriction digest with BsmBI and MluI to excise the P2A-puromycin cassette. An oligo containing a P2A-blasticidin cassette was purchased from IDT (5’-AGC GGA GCT ACT AAC TTC AGC CTG CTG AAG CAG GCT GGC GAC GTG GAG GAG AAC CCT GGA CCT ACC GGT ATG GCC AAG CCA CTG TCC CAA GAA GAG TCA ACT CTG ATC GAG AGG GCC ACT GCA ACC ATT AAT AGC ATT CCC ATC TCT GAA GAC TAT AGC GTA GCT AGT GCC GCA CTC AGC TCT GAT GGA CGC ATA TTC ACC GGC GTT AAT GTC TAC CAC TTC ACC ACC GGC GGA CCC TGC GCC GAA CTG GTC GTG CTG GGG ACC GCA GCC GCG GCC GCT GCC GGG AAT TTG ACG TGC ATT GTT GCA ATA GGC AAC GAG AAT AGG GGC ATC CTG TCA CCT TGC GGC CGG TGT CGG CAA GTG CTG CTG GAC CTG CAC CCC GGC ATC AAG GCC ATA GTC AAG GAT AGT GAT GCC CAG CCG ACC GCC GTT GGG ATT CGA GAA CTT CTG CCT TCT GGG TAC TCT GGG TAC GTC TGG GAA GGC TAG) and amplified with the primers (5’-CAA GAC TAG TGG AAG CGG AGC TAC TAA CTT CCC AGA CGT ACC C) using high-fidelity Phusion polymerase (NEB, Cat#
The PCR fragment was digested with BmtI and MluI, and ligated into the cut pLentiCRISPR1000, producing pLentiCRISPR1000-Blast.

Creation of THP-1 cells disrupting A3 genes

An A3F specific guide for exon 3 was designed (Fig. S1A and S2A) and evaluated manually for specificity to the A3F target sequence via an alignment with the most related members of the A3 family as described previously (26). Oligos with ends compatible with the Esp3I sites in pLentiCRISPR1000-Blast were purchased from IDT [ΔA3F gRNA: (5’-CAC CGG TAG TAG TAG AGG CGG GCG G) and (5’-CCA TCA TCA TCT CCG CCC GCC CAA G)]. The targeting construct was generated by annealing oligos and cloned by Golden Gate ligation into pLentiCRISPR1000-Blast. A guide with a common sequence among A3A exon 4, A3B exon 7 and A3G exon 7 was designed (Fig. 2A) and oligos with ends compatible with the Esp3I sites in pLentiCRISPR1000 (66) were purchased from IDT [PanZ1 gRNA: (5’-CAC CGT GGC CCG CAG CCT CCC ACT C) and (5’-GAA CGA GTG GGA GGC TGC GGG CCA C)]. The targeting construct was generated by annealing oligos and cloned by Golden Gate ligation into pLentiCRISPR1000 (66). All constructs were confirmed by Sanger sequencing (AZENTA) and sequence data were analyzed by Sequencher v5.4.6 (Gene Codes Corporation).

For transduction, VSV-G pseudotyped virus was generated by transfecting 2.5 μg of the pLentiCRISPR1000 or pLentiCRISPR1000-Blast targeting construct along with 1.67 μg of pΔ-NRF (HIV-1 gag, pol, rev, tat genes) (67) and 0.83 μg of pMD.G (VSV-G) expression vectors using TransIT-LT1 (Takara, Cat# MIR2306) into 293T cells. At 48 hours post-transfection, viral supernatants were harvested, filtered with 0.45 μm filters...
(Merck, Cat# SLHVR33RB), and concentrated by centrifugation (26,200 × g, 4°C, 2 hours).

Then, viral pellets were resuspended in 10% FBS/RPMI and incubated with cells for 48 hours. Forty-eight hours later, cells were placed under drug selection in 10% FBS/RPMI containing 1 µg/ml puromycin (InvivoGen, Cat# ant-pr) or 6 ng/ml blasticidin (InvivoGen, Cat# ant-bl). Single-cell clones were isolated by the limiting dilution of the drug-resistant cell pool and expanded. The expression levels of A3F protein in THP-1 ΔA3F#1 and #2, and THP-1ΔA3F/A3G#1 and #2 cells were confirmed by immunoblots (see Western blots).

To confirm indels in the A3F target sequence of the selected clones, genomic DNA was isolated by DNeasy Blood & Tissue Kits (Qiagen, Cat# 69504) and amplified with Choice-Taq DNA polymerase (Denville Scientific, Cat# CB4050-2) using primers (5’-GCT GAA GTC GCC CTT GAA TAA ACA CGC and 5’-TGT CAG TGC TGG CCC CG). The amplified PCR products were cloned into the pJET clonin g vector (Thermo Fisher Scientific, Cat# K1231) and subjected to Sanger sequencing (AZENTA). To confirm indels in the A3A, A3B and A3G target sequences of the selected clones (THP-1#11-4 and #11-7), genomic DNA was isolated by DNeasy Blood & Tissue Kits (Qiagen, Cat# 69504) and subjected to whole genome sequencing (WGS) (macrogen). The sequencing data were aligned by Isaac aligner (iSAAC-04.18.11.09). Off-target sites were analyzed by Cas-OFFinder (http://www.rgenome.net/cas-offinder/). For further analysis of indels between A3A and A3G, genomic DNAs from THP-1#11-4 and #11-7 were amplified using primers (5’-GGG GCT TTC TGA AAG AAT GAG AAC TGG GC and 5’-TGT CAG TGC TGG CCC CG). The amplified PCR products were cloned into the pJET cloning vector (Thermo Fisher Scientific, Cat# K1231) and subjected to Sanger sequencing (AZENTA). All sequence data were analyzed by Sequencher v5.4.6 (Gene Codes Corporation). To
assess the expression levels of A3 mRNAs and proteins, THP-1 parent, #11-4, and #11-7 were incubated in 10% FBS/RPMI including 500 units/ml IFN (R & D Systems, Cat# 11200-2) for 6 hours. Then, cells were harvested and subjected to RT-qPCR (see RT-qPCR) (Fig. 2C) and Western blot (see Western blot) (Fig. 2D).

Pseudo-single cycle infectivity assays

Vif-proficient and Vif-deficient (X26 and X27) HIV-1 IIIB C200 proviral expression constructs have been reported (68). HIV-1 IIIB C200 mutants with hyper- (H48 and 60EKGE63) and hypo- (V39) functional Vifs have been reported (10). An HIV-1 IIIB C200 Vif 5A mutant (40AAAAA44) has been described (26). HIV-1 IIIB C200 Vif 4A (14AKTK18) and 4A5A (14AKTK18 and 40AAAAA44) mutants were created by digesting pNLCSFV3-4A, and -4A5A proviral DNA construct [(37); kindly provided by Dr. Kei Sato, University of Tokyo, Japan] at Swal and Sall sites and cloned into pIIIB C200 proviral construct. The proviral expression vector encoding full length TF virus, CH58 (#11856) was obtained from the NARP. The creation of Vif-deficient CH58 mutant has been described previously (69).

HIV-1 single-cycle assays using VSV-G pseudotyped viruses were performed as described previously (23, 26). 293T cells were cotransfected with 2.4 μg of proviral DNA construct and 0.6 μg of VSV-G expression vector using TransIT-LT1 reagent (Takara, Cat# MIR2306) into 293T cells (3 × 10⁵). Forty-eight hours later, supernatants were harvested, filtered (0.45 μm filters, Merck, Cat# SLHVR33RB), and used to titrate on 2.5 × 10⁴ CEM-GXR reporter cells for MOI determinations. GFP+ cells were measured using a FACS Canto II (BD Biosciences) and the data were analyzed using FlowJo software.
v10.7.1 (BD Biosciences). 1 or 5 × 10⁶ target cells were infected with an MOI of 0.05 (for SupT11 derivatives) or 0.25 (for THP-1 derivatives) and washed with PBS twice at 24 hours post-infection and then incubated for an additional 24 hours. After 24 hours, supernatants were collected and filtered. The resulting viral particles were quantified by p24 ELISA (ZeptoMetrix, Cat# 0801008) and used to infect 1 × 10⁴ TZM-bl cells (1 or 2 ng of p24). At 48 hours postinfection, the infected cells were lysed with a Bright-Glo luciferase assay system (Promega, Cat# E2650) and the intracellular luciferase activity was measured by a Synergy H1 microplate reader (BioTek) or Centro XS3 LB960 microplate luminometer (Berthold Technologies).

**Quantification of LRT products**

Viruses were produced by infecting VSV-G pseudotyped virus into THP-1 cells as described above (see HIV-1 infectivity assays) and the resulting viral particles were quantified by p24 ELISA (ZeptoMetrix, Cat# 0801008). The viral supernatants including 20 ng of p24 antigen were used for infection into SupT11 cells. At 12 hours postinfection, cells were harvested and washed with PBS twice. Then, total DNA was isolated by DNeasy Blood & Tissue Kits (Qiagen, Cat# 69504) and treated with RNase A (Qiagen, Cat# 19101) according to the manufacturer’s instruction. Following DpnI digestion, 50 ng of DNA was used to amplify LRT products and CCR5 gene with the following primers; LRT forward: (5’-CGT CTG TTG TGT GAC TCT GG) and LRT reverse: (5’-TTT TGG CGT ACT CAC CAG TCG). CCR5 forward: (5’-CCA GAA GAG CTG AGA CAT CCG) and CCR5 reverse (5’-GCC AAG CAG CTG AGA GGT TAC T). qPCR was performed using Power SYBR Green PCR Master Mix (Thermo Fisher Scientific, Cat# 4367659) and
fluorescent signals from resulting PCR products were acquired using a Thermal Cycler Dice Real Time System III (Takara). Finally, each LRT product was represented as values normalized by the quantity of the CCR5 gene (Fig. 4C).

**RT-qPCR**

Cells were harvested and washed with PBS twice. Then, total RNA was isolated by RNA Premium Kit (NIPPON Genetics, Cat# FG-81250) and cDNA was synthesized by Transcriptor Reverse Transcriptase (Roche, Cat# 03531287001) with random hexamer.

RT-qPCR was performed using Power SYBR Green PCR Master Mix (Thermo Fisher Scientific, Cat# 4367659). Primers for each A3 mRNA have been reported previously (70, 71). A3A forward: (5'-GAG AAG GGA CAA GCA CAT GG) and A3A reverse: (5'-TGG ATC CAT CAA GTG TCT GG). A3B forward: (5'-GAC CCT TTG GTC CTT CGA C) and A3B reverse: (5'-GCA CAG CCC CAG GAG AAG). A3C forward: (5'-AGC GCT TCA GAA AGT GG) and A3C reverse: (5'-AAG TTT CGT TCC GAT CGT TG). A3D forward: (5'-ACC CAA ACG TCA GTC GAA TC) and A3D reverse: (5'-CAC ATT TCT GCG TGG TTC TC). A3F forward: (5'-CCG TTT GGA CGC AAA GAT) and A3F reverse: (5'-CCA GGT GAT CTG GAA ACA CTT). A3G forward: (5'-CCG AGG ACC CGA AGG TTA C) and A3G reverse: (5'-TCC AAC AGT GCT GAA ATT CG). A3H forward: (5'-AGC TGT GGC CAG AAG CAC) and A3H reverse: (5'-CGG AAT GTT TCG GCT GTT). TATA-binding protein (TBP) forward: (5'-CCC ATG ACT CCC ATG ACC) and TBP reverse: (5'-TTT ACA ACC AAG ATT CAC TGT GG). Fluorescent signals from resulting PCR products were acquired using a Thermal Cycler Dice Real Time System III (Takara).
Finally, each A3 mRNA expression level was represented as values normalized by TBP mRNA expression levels (Fig. 2C).

**Hypermutation analyses**

Hypermutation analyses were performed as previously described (23, 26, 45). Genomic DNAs containing HIV-1 proviruses were recovered by infecting viruses produced in derivatives of THP-1 or SupT11 cells into SupT11 using DNeasy Blood & Tissue Kits (Qiagen, Cat# 69504). Following DpnI digestion, the viral pol region was amplified by nested PCR with outer primers (876 bp) [(5'-TCC ART ATT TRC CAT AAA RAA AAA) and (5'-TTY AGA TTT TTA AAT GGY TYT TGA)] and inner primers (564 bp) [(5'-AAT ATT CCA RTR TAR CAT RAC AAA AAT) and (5'-AAT GGY TYT TGA TAA ATT TGA TAT GT)]. The resulting 564 bp amplicon was subjected to pJET cloning. At least 10 independent clones were Sanger sequenced (AZENTA) for each condition and analyzed by the HIV sequence database (https://www.hiv.lanl.gov/content/sequence/HYPERMUT/hypermut.html). Clones with identical mutations were eliminated.

**Western blot**

Western blot for cell and viral lysates were performed as described previously (23, 26, 72). Cells were harvested, washed with PBS twice, and lysed in lysis buffer [25 mM HEPES (pH7.2), 20% glycerol, 125 mM NaCl, 1% Nonidet P40 (NP40) substitute (Nacalai Tesque, Cat# 18558-54)]. After quantification of total protein by protein assay dye (Bio-Rad, Cat# 5000006), lysates were diluted with 2 × SDS sample buffer [100 mM Tris-HCl...
(pH 6.8), 4% SDS, 12% β-mercaptoethanol, 20% glycerol, 0.05% bromophenol blue] and
boiled for 10 minutes. Virions were dissolved in 2 × SDS sample buffer and boiled for 10
minutes after pelleting down using 20% sucrose (26,200 × g, 4˚C, 2 hours). Then, the
quantity of p24 antigen was measured by p24 ELISA (ZeptoMetrix, Cat# 0801008).

Proteins in the cell and viral lysates (5 μg of total protein and 10 ng of p24 antigen)
were separated by SDS-PAGE and transferred to PVDF membranes (Millipore, Cat#
IPVH00010). Membranes were blocked with 5% milk in PBS containing 0.1% Tween 20
(0.1% PBST) and incubated in 4% milk/0.1% PBST containing primary antibodies: mouse
anti-HSP90 (BD Transduction Laboratories, Cat# 610418, 1:5,000); rabbit anti-A3B
(5210-87-13, 1:1,000) (73); rabbit anti-A3C (Proteintech, Cat# 10591-1-AP, 1:1,000);
rabbit anti-A3F (675, 1:1,000) (74); rabbit anti-A3G (NARP, #10201, 1:2,500); rabbit anti-
A3H (Novus Biologicals, NBP1-91682, 1:5,000): mouse anti-Vif (NARP, #6459, 1:2,000);
mouse anti-p24 (NARP, #1513, 1:2,000). Subsequently, the membranes were incubated
with horseradish peroxidase (HRP)-conjugated secondary antibodies: donkey anti-rabbit
IgG-HRP (Jackson ImmunoResearch, 711-035-152; 1:5,000); donkey anti-mouse IgG-
HRP (Jackson ImmunoResearch, 715-035-150). SuperSignal West Femto Maximum
Sensitivity Substrate (Thermo Fisher Scientific, Cat# 34095) or Super signal atto (Thermo
Fisher Scientific, Cat# A38555) was used for HRP detection. Bands were visualized by
the Amersham Imager 600 (Amersham).

Statistical analyses
Statistical significance was performed using a two-sided paired t test (Fig. 1C, 2C,
3B, 3C, 4C, and 5). GraphPad Prism software v8.4.3 was used for these statistical tests.
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The authors declare that they have no competing interests.

References


Figure legends

Figure 1. Endogenous A3H does not inhibit HIV-1 in THP-1 cells.

(A) A3H haplotypes in THP-1 cells. The indicated positions are key amino acid residues that determine the expression of unstable (hapI) or stable (hapII) A3H protein.

(B) Schematic of the susceptibility of Vif mutants to stable A3H haplotypes. Key amino acid residues that determine the susceptibility of HIV-1 IIIB Vif to restriction by stable A3H haplotypes. -, full resistance; +, partial resistance; ++++, sensitivity.

(C) Representative infectivity of hyper- and hypo-functional Vif HIV-1 mutants. Top panels show the infectivity of hyper-Vif, hypo-Vif, and IIIB Vif, and Vif-deficient HIV-1 mutants produced in THP-1 cells compared to the same viruses produced in SupT11 cells with stable expression of the control vector or A3H haplotype II. The amounts of produced viruses used to infect TZM-bl cells was normalized to p24 levels. Each bar shows the average of four independent experiments with the standard deviation (SD). Data are represented as relative infectivity compared to hyper-Vif HIV-1. Statistical significance was determined using the two-sided paired t test. *P < 0.05 compared with the infectivity of hyper-Vif HIV-1. The bottom panels are representative Western blots of three independent experiments. The levels of viral and cellular proteins in virus-like particles (VLPs) and whole cell lysates are shown. p24 and HSP90 were used as loading controls.

(D) G-to-A mutations. Average number of G-to-A mutations in the 564 bp pol gene after infection with hyper-Vif, hypo-Vif, IIIB Vif, or Vif deficient HIV-1 produced from THP-1 or SupT11 cells expressing either the vector control or A3H hapII. Each bar depicts the average of three independent experiments with SD.
(E) G-to-A mutation profile. Dinucleotide sequence contexts of G-to-A mutations in the 564 bp pol gene after infection with the indicated viruses produced from indicated cell lines. Each vertical line indicates the location of the dinucleotide sequence contexts described in the legend within the 564 bp amplicon (horizontal line).

Figure 2. Disruption of the A3A to A3G genes in THP-1 cells.

(A) Schematic of the A3 gene at the A3 locus. The A3 family of genes comprises seven members with one or two Z domains (single- or double-domain deaminases) which belong to three phylogenetically distinct groups shown in green, yellow, and blue. Three sites with an identical sequence (5′-GAG TGG GAG GCT GCG GGC CA) in exon 4 of the A3A gene, exon 7 of the A3B gene, and exon 7 of the A3G gene are targeted by gRNA, as indicated by arrows. The three predicted scenarios are shown. Bar represents 15,000 bp.

(B) Mapping of WGS sequencing data to the A3 locus. Genomic DNA from parental THP-1, THP-1#11-4, and #11-7 cells were subjected to WGS analysis, with an extensive deletion including the A3A–A3G genes observed in THP-1#11-4 and #11-7 clones.

(C) RT-qPCR data. Parental THP-1, THP-1#11-4, and #11-7 cells were treated with 500 units/ml type I IFN. Total RNA was isolated after 6 hours. A3 mRNA expression levels were quantified by RT-qPCR and are normalized to TBP mRNA levels. Each bar represents the average of three independent experiments with SD. Statistical significance was determined using the two-sided paired t test. *, P < 0.05 compared to untreated cells.
(D) Representative Western blots of three independent experiments. Levels of indicated A3 proteins in whole cell lysates from cells treated with or without type I IFN are shown. HSP90 was used as a loading control.

Figure 3. Pseudo-single cycle infectivity assays for each HIV-1 mutant in A3-null THP-1 cells.

(A) Schematic of the susceptibility of Vif mutants to A3F and A3G. Key amino acid residues that determine the susceptibility of HIV-1 IIIB Vif to restriction by A3F and A3G. -, resistance; +, sensitivity.

(B) Representative infectivity of Vif-proficient, Vif-deficient, Vif4A, Vif5A, and Vif4A5A HIV-1 mutants in SupT11 cells stably expressing vector control, A3F, or A3G. Top panels show the infectivity of indicated HIV-1 mutants produced in SupT11 cells stably expressing vector control, A3F, or A3G. The amounts of produced viruses used to infect TZM-bl cells was normalized to p24 levels. Each bar represents the average of four independent experiments with SD. Data are presented as relative infectivity compared to Vif-proficient HIV-1 (WT). Statistical significance was assessed using the two-sided paired t test. *P < 0.05 compared to Vif-proficient HIV-1. Bottom panels are representative Western blots of three independent experiments. Levels of indicated viral and cellular proteins in VLPs and whole cell lysates are shown. p24 and HSP90 were used as loading controls.

(C) Representative infectivity of Vif-proficient, Vif-deficient, Vif4A, Vif5A, and Vif4A5A HIV-1 mutants in A3-null THP-1 cells. Top panels show the infectivity of indicated HIV-1 mutants produced in parental or A3-null THP-1 cells. The amounts of produced viruses
used to infect TZM-bl cells was normalized to p24 levels. Each bar represents the average of four independent experiments with SD. Data are presented as infectivity relative to Vif-proficient HIV-1 (WT). Statistical significance was assessed using the two-sided paired t test. *P < 0.05 compared to Vif-proficient HIV-1. Bottom panels are representative Western blots of three independent experiments. Levels of indicated viral and cellular proteins in VLPs and whole cell lysates are shown. p24 and HSP90 were used as loading controls.

**Figure 4. A3 proteins inhibit Vif-deficient HIV-1 by both deaminase-dependent and independent mechanisms in THP-1 cells.**

**(A)** G-to-A mutations. Average number of G-to-A mutations in the 564 bp pol gene after infection with hyper-Vif, hypo-Vif, IIIB Vif, or Vif-deficient HIV-1 produced from THP-1 or SupT11 expressing either vector control or A3H hapII. Each bar depicts the average of three independent experiments with SD.

**(B)** G-to-A mutation profile. Dinucleotide sequence contexts of G-to-A mutations in the 564 bp pol gene after infection with the indicated viruses produced from indicated cell lines. Each vertical line indicates the location of the dinucleotide sequence contexts described in the legend within the 564 bp amplicon (horizontal line).

**(C)** Representative LRT quantification data for Vif-proficient, Vif-deficient, Vif4A, Vif5A, and Vif4A5A HIV-1 mutants in each A3-null THP-1 subclone. Data show LRT products of the indicated HIV-1 mutants produced in parental or indicated A3-null THP-1 cells. The amount of produced viruses used to infect SupT11 cells was normalized to p24 levels. LRT products were measured by qPCR. Each bar represents the average of four
independent experiments with SD. LRT products were normalized to the quantity of the CCR5 gene relative to Vif-proficient HIV-1 (WT). Statistical significance was assessed using the two-sided paired t test. *P < 0.05 compared to Vif-proficient HIV-1 LRT products.

Figure 5. Pseudo-single cycle infectivity assays of TF virus molecular clone in A3A-to-A3G-null THP-1 cells.

Infectivity of Vif-proficient and Vif-deficient CH58 viruses. Top panels show the infectivity of Vif-proficient and Vif-deficient HIV-1 produced in parental THP-1, THP-1#11-4, or THP-1#11-7 cells. The amounts of produced viruses used to infect TZM-bl cells was normalized to p24 levels. Each bar represents the average of four independent experiments with SD. Data are represented as relative to Vif-proficient HIV-1 (WT). Statistical significance was assessed using the two-sided paired t test. *P < 0.05 compared to Vif-proficient HIV-1. The bottom panels are representative Western blots of three independent experiments. The levels of indicated viral and cellular proteins in VLPs and whole cell lysates are shown. p24 and HSP90 were used as loading controls.

Figure S1. Development of A3F-null THP-1 cells.

(A) A3F exon 3 sequences encompassing the gRNA target site in parental THP-1 and two independent A3F-null THP-1 cells. Indels in two alleles for each A3F-null THP-1 clone are shown.

(B) Representative Western blots of three independent experiments. Levels of A3F and A3G protein in whole cell lysates are shown. HSP90 was used as a loading control.
Figure S2. Development of A3F/A3G-null THP-1 cells.

(A) A3F exon 3 sequences encompassing the gRNA target site in parental THP-1 and two independent A3F/A3G-null THP-1 cells. Indels in two alleles for each A3F/A3G-null THP-1 clone are shown.

(B) Representative Western blots of three independent experiments. Levels of A3F and A3G protein in whole cell lysates are shown. HSP90 was used as a loading control.

Fig. S3 Sequence analysis of flanking region targeted by gRNA in THP-1#11-4 and #11-7.

(A) A3A exon 4 and A3G exon 7 hybrid sequences encompassing the gRNA target site in THP-1#11-4 cells. Only one nucleotide difference (>99% identity) was observed between A3A exon 4 and A3G exon 7 and is shown in purple (A3A, cytosine) or green (A3G, adenine). Indels in six alleles of the THP-1#11-4 clone are shown.

(B) A3A exon 4 and A3G exon 7 hybrid sequences encompassing the gRNA target site in THP-1#11-7 cells. Only one nucleotide difference (>99% identity) was observed between A3A exon 4 and A3G exon 7 and is shown in purple (A3A, cytosine) or green (A3G, adenine). Indels in three alleles of the THP-1#11-7 clone are shown.

Fig. S4 Deletions around predicted A3G pseudogene.

Mapping of WGS sequencing data to off-target and downstream regions on chromosome 12. Genomic DNA from parental THP-1, THP-1#11-4, and THP-1#11-7 cells were subjected to WGS analysis. The yellow box indicates the off-target sequence in the
predicted pseudogene. Several deletions were observed in the regions indicated by green dot boxes in THP-1#11-4 and THP-1#11-7 clones.
Figure 1

A

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B

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<td>ΔVif</td>
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C

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E

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Figure 2

A

B

C

D

Mapping of whole-genome sequencing data to the A3 locus

- Scenario 1 (29.9 kbp deletion): A3B, A3F, A3G, A3H
- Scenario 2 (95 kbp deletion): A3A, A3B, A3G
- Scenario 3 (125 kbp deletion): A3A, A3B, A3C

Time after IFN treatment (h)

A3A
A3B
A3C
A3D
A3F
A3G
A3H

A3A
A3B
A3C
A3G

Parent #11-4 #11-7

5210
87-13

IFN
(-) (+) (+) (+)

A3A
A3B
A3C
A3F
A3G
A3H
HSP90

kDa
40
35
25
40
40
15
100

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(https://doi.org/10.1101/2023.03.28.534666)
Figure 3

A

Sensitivity to Vif

Parent

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

B

SupT11

Vector

A3F

A3G

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

C

THP-1

Parent

ΔA3G

ΔA3F#1

ΔA3F#2

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)

*p24

p24

Vif

WT

DRNR — YRIHY / // + +

4A

AKTK — YRIHY / // + —

Vif

5A

DRNR — AAAA / // — + +

4A5A

AKTK — AAAA / // — + +

ΔA3F/A3G#1

ΔA3F/A3G#2

#11-4

#11-7

(ΔA3A to A3G)

(ΔA3G to A3G)
Figure 4

(A) SupT1

(B) THP-1

(C) Relative LRT

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Figure 5

Relative infectivity

CH58

THP-1#11-4

THP-1#11-7

Vif WT

Δ

No virus

VLP

25

40

25

40

kDa

kDa

Vif WT

Δ

No virus

A3G

40

25

100

kDa

Vif

25

100

kDa

HSP90