Mutations Conferring Resistance to Human Immunodeficiency Virus Type 1 Fusion Inhibitors Are Restricted by gp41 and Rev-Responsive Element Functions

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One of the human immunodeficiency virus (HIV) envelope proteins, gp41, plays a key role in HIV fusion. A gp41-derived peptide, T-20, efficiently inhibits HIV fusion and is currently approved for treatment of HIV-infected individuals. Although resistant variants have been reported, the mechanism of the resistance remains to be defined. To elucidate the mechanism in detail, we generated variants resistant to C34, a peptide derived from the gp41 carboxyl terminus heptad repeat (C-HR) in vitro. The resistant variant had a 5-amino-acid deletion in gp120 and a total of seven amino acid substitutions in gp41. Binding assays revealed that an I37K substitution in the N-terminal heptad repeat (N-HR) impaired the binding of C34, whereas an N126K substitution in the C-HR enhanced the binding to mutated N-HR, indicating that both mutations were directly involved in resistance. On the other hand, substitutions for A30 and D36 seemed to be secondary mutations, located complementary to each other in the Rev-responsive element (RRE), and were mutated simultaneously to maintain the secondary structure of the RRE that was impaired by the mutations at I37. Thus, HIV acquired resistance to C34 by mutations in N-HR, which directly interacted with C34. However, since this region also encoded the RRE, additional mutations were required to maintain viral replication. These results suggest that HIV fusion is one of the attractive targets for HIV chemotherapy.

Peptide inhibitors that block human immunodeficiency virus type 1 (HIV-1) fusion were first reported by Wild et al. (30). Recently, a peptide fusion inhibitor (T-20 or enfuvirtide) has been approved in the United States and Europe for treatment of HIV-infected individuals. The peptide sequence of T-20 is derived from the gp41 C terminus heptad repeat (C-HR) sequence, which corresponds to a linear region of 36 amino acids, and T-20 inhibits fusion by binding to the N-terminal heptad repeat (N-HR) of gp41 and preventing 6-helix bundle formation (4, 30). In HIV-infected patients, the effect of T-20 in combination with an antiretroviral regimen that was optimized with the aid of phenotypic and genotypic resistance testing (TORO 1 and 2) has been reported to suppress drug-resistant HIV replication more efficiently than the optimized regimen alone (17, 18).

The emergence of T-20-resistant HIV-1 was first reported in clinical patients receiving T-20 monotherapy in a phase I clinical trial (28) and subsequently in combined regimens employed in phase II and III trials of T-20 (23, 25). The T-20 susceptibility of recombinant HIV-1 containing the identified substitutions was examined in vitro and considered to be moderately resistant (5.4- to 6.3-fold) (28). However, the detailed mechanism of resistance of these variants still remains to be elucidated. On the other hand, Rimsky et al. revealed that three continuous amino acids in the N-HR (GIV at positions 36 to 38 of gp41) were crucial for the inhibition of HIV-1 entry by T-20 and for efficient association between N-HR and T-20 in vitro (26). Fikker et al. also reported that HIV-1 variants resistant to T-20 contained substitutions in gp41, L33S and N43K, and a deletion of 5 amino acids, FNSTW (ΔFNSTW), in the V4 region of gp120 (9). L33S and N43K contributed to T-20 resistance, whereas the 5-amino-acid deletion alone had little effect on T-20 sensitivity. These results suggest that substitutions in the N-HR directly affect T-20 binding. Although the baseline sensitivity of HIV-1 to T-20 is defined by amino acid substitutions in gp41, coreceptor specificity is influenced by substitutions in the V3 loop in gp120, affects the fusion kinetics, and modulates T-20 sensitivity (4, 5).

To elucidate the mechanism of resistance to the peptide fusion inhibitors, we generated and characterized HIV-1 variants resistant to C34, a gp41 C-HR-derived peptide (2, 22) (Fig. 1A). During the selection of C34-resistant variants, we observed a 5-amino-acid deletion in the gp120 V4 region and a total of seven amino acid substitutions in gp41. Among the deletion and the substitutions, I37K and N126K play a key role in the resistance to C-HR-derived peptides, including T-20. Other deletions or substitutions were considered to enhance C34 resistance and/or improve the impaired replication kinetics. A30V and D36G maintained the Rev-responsive element (RRE) structure destabilized by I37T and I37K, respectively. Thus, these results reveal that the deletions or substitutions conferring resistance are restricted by both gp41 and RRE functions, suggesting that HIV-1 fusion is one of the most ideal targets for chemotherapy.

MATERIALS AND METHODS

Cells and viruses. MT-2 and Cos-7 cells were grown in RPMI 1640- and Dulbecco’s modified Eagle medium-based culture medium, respectively. HeLa-
CD4-LTR-β-gal cells were kindly provided by M. Emerman through the AIDS Research and Reference Reagent Program, Division of AIDS, National Institute of Allergy and Infectious Disease (Bethesda, Md.), and used for the drug susceptibility assay (multinuclear activation of galactosidase indicator [MAGI] assay) as described previously (12, 14, 21). An HIV-1 infectious clone, pNL4-3, which was kindly provided by H. Sakai, Institute for Virus Research, Kyoto University (Kyoto, Japan), was used for constructions and the production of HIV-1 variants. A wild-type HIV-1, HIV-1WT, was generated by transfection of pNL4-3 into Cos-7 cells.

**Antiviral agents.** The peptides used were N36, derived from the N-HR of gp41, and C34 and T-20, derived from the C-HR of gp41. The peptides were synthesized as described previously (24) and are depicted in Fig. 1A. 22-Dideoxycytidine (ddC) was purchased from Sigma (St. Louis, Mo.).

**Determination of drug susceptibility of HIV-1.** The peptide sensitivity of infectious clones was determined by the MAGI assay with some modifications (14, 21). Briefly, the target cells (HeLa-CD4-LTR-β-gal; 10^4 cells/well) were plated in 96-well flat microtiter culture plates. On the following day, the cells were inoculated with the HIV-1 clones (60 MAGI U/well, giving 60 blue cells after 48 h of incubation) and cultured in the presence of various concentrations of drugs in fresh medium. Forty-eight hours after viral exposure, all the blue cells stained with X-Gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) were counted in each well. The activity of test compounds was determined as the concentration that blocked HIV-1 replication by 50% (50% effective concentration [EC50]).

**Construction of recombinant HIV-1 clones.** Recombinant infectious HIV-1 clones carrying various mutations in gp120 and/or gp41 were generated by using pNL4-3. Briefly, the desired mutations were introduced into the NheI-BamHI region (1,220 bp) of pSLgp41WT, which encoded nucleotides 7250 to 8469 of pNL4-3, by an oligonucleotide-based mutagenesis method (29). NheI-BamHI fragments were inserted into pNL4-3, generating various molecular clones with the desired mutations. Each molecular clone was transfected into Cos-7 cells (10^5 cells/six-well culture plate). After 48 h, MT-2 cells (10^6 cells/well) were added and cocultured with the Cos-7 cells for an additional 24 h. When an extensive cytopathic effect (CPE) was observed, the supernatants were harvested and stored at -80°C until use.

**Generation of HIV-1 variants resistant to C34.** MT-2 cells were exposed to HIV-1WT and cultured in the presence of C34 at an initial concentration of 0.0001 μM. Cultures were incubated at 37°C until an extensive CPE was observed. The culture supernatants were used for further passages in MT-2 cells in the presence of twofold increasing concentrations of C34 when massive CPEs were observed. At the indicated passages, proviral DNAs from the lysates of infected cells were sequenced, and the EC50s of the HIV-1 variants were determined with the MAGI assay.

**FIG. 1.** Schematic view of HIV-1 gp41 (A) and induction of C34-resistant HIV-1 (B). The locations of the fusion peptide (FP), N-terminal heptad repeat region (N-HR), C-terminal heptad repeat region (C-HR), transmembrane domain (TM), various gp41-derived peptides, and the Rev-responsive element (RRE) coding region are shown (A). The residue numbers of each peptide correspond to their positions in gp41. The bold underlined letters in the boxes indicate the novel mutations that have been reported in T-20-resistant HIV-1 variants (T-20r) in vitro (9, 26) and that have been observed in C34-resistant HIV-1 variants (C34r). (B) HIV-1WT was passaged in the presence of increasing concentrations of C34 in MT-2 cells. The dose-escalating selection was carried out for a total of 93 passages, with compound concentrations ranging from 0.0001 to 20 μM. At the indicated passages, proviral DNAs from the lysates of infected cells were sequenced, and the EC50s of the HIV-1 variants were determined with the MAGI assay.
were seen in the earlier periods. Such dose-escalating culture was performed until resistant variants were obtained. This selection was carried out for a total of 93 passages. At the indicated passages (Fig. 1B), the sequence of the env region was determined by direct sequencing of the proviral DNA extracted from the infected MT-2 cells.

Viral replication kinetics assay. MT-2 cells (10⁶ cells/5 ml) were infected with each virus preparation (500 MAGI U) for 4 h. The infected cells were then washed and cultured in a final volume of 5 ml. The culture supernatants (100 µl) were harvested on days 1, 2, 4, 6, and 8 after infection, and the p24 antigen amounts were determined.

For competitive HIV-1 replication assays (CHRA), two titrated infectious clones to be examined were mixed and added to MT-2 cells (10⁶ cells/3 ml) as described previously (15) with some modifications. To ensure that the two infectious clones being compared were of approximately equal infectivity, a fixed amount (500 MAGI U) of one infectious clone was mixed with three different amounts (250, 500, and 1,000 MAGI U) of the other infectious clone. On day 1, one third of the infected MT-2 cells were harvested and washed twice with phosphate-buffered saline, and the cellular DNA was extracted. The purified DNA was subjected to nested PCR and then direct DNA sequencing. The HIV-1 cocculture which best approximated a 50:50 mixture on day 1 was further propagated. Every 6 to 7 days, the cell-free supernatant of the virus coculture (1 ml) was transmitted to new uninfected MT-2 cells. The cells harvested at the end of each passage were subjected to direct sequencing, and the viral population change was determined.

Binding assay. Each peptide (40 µM) was mixed with 10 mM phosphate-buffered saline-140 mM NaCl, pH 7.4, in an Avid model 202 D8 spectrometer equipped with a thermostatted temperature controller. The thermal stability was assessed by monitoring the change in the circular dichroism signal at 222 nm. The midpoint of the thermal unfolding transition (melting temperature [Tm]) of each complex was determined as described previously (24).

Gel shift assay. RNA of the RRE region and recombinant Rev were prepared as described previously (10) with some modifications. Briefly, the RRE region of the variants (nucleotides 7748 to 8009 of pNL4-3) was introduced into pBluescript (Stratagene, La Jolla, Calif.). In vitro RNA transcription was performed with T7 RNA polymerase and [32P]UTP. Recombinant Rev was generated by use of pGEX-6P-1/B2L1 expression system (Amersham Biosciences, Piscataway, N.J.). The RNA and Rev were mixed at 25°C for 20 min in binding buffer (50 mM Tris-HCl, pH 7.5, 150 mM KCl, 1 mM dithiothreitol, 8% glycerol, 50 µg of tRNA/ml, and 100 µg of bovine serum albumin/ml) and subjected to native acrylamide gel electrophoresis.

RESULTS

Amino acid substitutions identified in the env region of C34-resistant HIV-1. At passage 14 (P-14) in the culture where HIV-1 was propagating in the presence of C34 (0.0032 µM), one amino acid substitution, glutamine to histidine at position 39 (Q39H), in the N-HR, was newly identified in the C-HR, whereas Q39H had returned to the original wild-type amino acid. At P-41 (0.026 µM), two substitutions, A30V and I37I/T (mixture of I and T), were observed in the N-HR in addition to N126K, while at P-50 (0.077 µM), definitive I37T was detected (A30V/I37T/N126K) (Fig. 1B). At P-75 (2.5 µM), A30V had returned to the original wild-type amino acid, D36G was detected, I37T was substituted for I37K, and L204I which was located in the cytoplasamic domain of gp41, was identified (D36G/I37K/N126K/L204I) (Fig. 1B). At P-92 (20 µM), a deletion of five amino acids, FNSTW, in the V4 loop of gp120 (∆FNSTW) was observed together with the four substitutions (∆FNSTW/D36G/I37K/N126K/L204I) (Fig. 1B). In addition to the env region, we also examined the Tat and Rev-encoding regions but did not observe any substitutions. These results suggest that, in order to develop a higher resistance to C34, HIV-1 acquires not only multiple substitutions in gp41 but also the 5-amino-acid deletion in gp120.

Susceptibility of the different env recombinant viruses to fusion inhibitors. To clarify which substitutions among the identified changes were responsible for C34 resistance, we first generated infectious HIV-1 clones containing the deletion (∆FNSTW) in gp120 or the single amino acid substitutions (A30V, D36G, I37T, I37K, Q39H, N126K, or L204I) in gp41 that were observed during the selection procedure (Fig. 1B). We also evaluated the activities of the gp41-derived peptides N36, T-20, and C34 and a reverse transcriptase inhibitor used as a control, ddC, against these strains with the MAGI assay (Table 1).

HIV-1∆FNSTW, HIV-1A30V, HIV-1Q39H, and HIV-1L204I showed weak resistance to C34 compared with HIV-1WT (less than fivefold). Interestingly, D36G, observed in the majority of HIV-1 strains (16), conferred an increased T-20 susceptibility to HIV-1 (10-fold), in agreement with previous reports (20, 26), whereas D36G did not contribute C34 resistance by itself (0.8-fold). Although I37T has also been reported as one of the T-20 resistance mutations in vitro, its detailed mechanism of resistance remains unknown (20, 26). In our experiments, I37T conferred T-20 and C34 resistance to HIV-1 (13- and 11-fold, respectively), and I38K also conferred both T-20 and C34 resistance (212- and 13-fold, respectively). HIV-1N126K showed moderate resistance to C34 (6.8-fold). Neither the deletion in gp120 nor any of the substitutions in gp41 conferred resistance to N36 or ddC (Table 1).

Although the I37 substitutions appeared to be primarily responsible for C34 resistance, the C34 resistance levels of the I37 substitution variants were not comparable to that of the selected virus at P-93 (EC50, 0.78 µM). Therefore, we generated infectious HIV-1 clones containing the identified substitutions combined with I37T or I37K and determined their susceptibilities to the peptides (Table 1). The combination of I37K and N126K enhanced C34 resistance (13- to 28-fold), whereas HIV-1I37T, HIV-1I37T/N126K, and HIV-1A30V/I37T/N126K showed comparable resistance levels to C34. Moreover, I37K/N126K combined with D36G (D36G/I37K/N126K) enhanced C34 resistance (72-fold), although the L204I substitution combined with D36G/I37K/N126K decreased the levels of resistance to both T-20 and C34 (10- and 54-fold, respectively). A clone containing the deletion in gp120 and four substitutions in gp41, HIV-1∆FNSTW/D36G/I37K/N126K/L204I, showed the highest resistance to C34 (83-fold) and cross-resistance to T-20 (64-fold). These results indicate that the I37K substitution is mainly responsible for C34 resistance, whereas the other substitutions enhance the resistance or improve the impaired viral replication kinetics.

Next, we generated a T-20-resistant molecular clone which had been previously reported (26), HIV-1D36G/V38M, and evaluated the susceptibility to N36, T-20, and C34. HIV-1D36G/V38M showed moderate resistance to both T-20 and C34 (5.1- and 15-fold, respectively). Combined with the finding that I37K is the major mutation for resistance to C34, this region, positions 37 and 38 of gp41, appears to be involved in resistance to both T-20 and C34, while changes at position 36 appear to be largely restricted in their effects to T-20.
Peptide binding affinity. To clarify the effect of the substitutions on the interaction of N-HR and C-HR, the binding affinity of the peptides in vitro was examined with the synthesized peptides (Table 2). The affinity between N36D36G/I37K and C34 was unstable even at 37°C, indicating that the peptide inhibitor C34 hardly bound to N36D36G/I37K. However, it is still unclear whether it is a direct effect of the N-HR mutations decreasing the affinity of C34 binding or an indirect effect of the N-HR mutations destabilizing the N-HR trimer formation. In contrast, C34N126K, with the substitution responsible for the resistance, showed enhanced binding affinity not only to N36 but also to N36D36G/I37K. Thus, there are two implications of mutations in gp41 for conferring C34 resistance: the decreased affinity of C34 for N36D36G/I37K and the increased affinity of C34N126K for both N36 WT and N36D36G/I37K. In other words, the D36G and I37K substitutions in the N-HR interfere with the binding of the peptide inhibitors, such as T-20 and C34, and N126K in the C-HR enhances the intra-gp41 binding of N-HR and C-HR compared with the peptide inhibitors.

Replication kinetics of C34-resistant variants. To determine the effects of the identified deletion and mutations on HIV-1 replication, we first examined the replication kinetics of HIV-1 variants by p24 production in the culture supernatants. The p24 production by the variants ranged from 14 to 34% of that of HIV-1WT (HIV-1I37T/N126K, 14%; HIV-1 N126K, 30%; and HIV-1A30V/I37T/N126K, 34%) as determined at day 8 (Fig. 2). Next, the replication levels of variants with representative substitutions were compared by CHRA. The resistances (n-fold) of the variants are also shown in Fig. 2. Since HIV-1Q39H was considered to be one of the polymorphisms and appeared only transiently, we first compared the replication of HIV-1WT and HIV-1N126K and found an impaired replication profile for HIV-1N126K. The variant with a combination including the I37T substitution, HIV-1 I37T/N126K, which was not observed during selection, showed the slowest replication profile. To develop an HIV-1A30V/I37T/N126K variant, the A30V substitution was introduced first, and then the I37T substitution was introduced (Fig. 1B). This was consistent with the results of the CHRA that the replication profile of HIV-1A30V/I37T/N126K was

![FIG. 2. Replication kinetics of the resistant variants. The replication kinetics determined by p24 antigen production and the CHRA are summarized. The data are depicted as the resistance (x axis) and replication (y axis) compared with those of HIV-1WT. Variants observed (continuous arrows) and not observed (dashed arrows) in the selection are shown in the order of their emergence.](http://jvi.asm.org/)
greater than that of HIV-1 \text{I37T/N126K}, suggesting that the A30V substitution improved the impaired replication profile of HIV-1 \text{I37T/N126K}. Since both HIV-1 \text{I37K} and HIV-1 \text{I37K/N126K} could replicate for only a few passages, these variants could not be studied. For \text{I37K/N126K} substitutions, D36G instead of A30V substitution seemed to be suitable in order to maintain resistance and replication, although the replication of HIV-1 \text{D36G/I37K/N126K} still remained impaired. The L204I substitution improved the replication kinetics of HIV-1 \text{D36G/I37K/N126K}. The deletion in the V4 region contributed to both the increased resistance and replication. These results indicate that among the deletion and the substitutions, \text{I37T/K} and \text{N126K}, and especially \text{I37K}, are related to resistance, while A30V, D36G, and L204I are associated with improvement of replication and the deletion in the V4 region had modest effects on increased resistance and replication.

**DISCUSSION**

After adsorption to cells followed by conformational changes of gp120, HIV fusion takes place by interaction and binding of the N-HR and C-HR of gp41. C-HR-derived peptides, e.g., T-20 or C34, inhibit HIV replication as a decoy for C-HR (2, 30). Previous reports showed that mutations in N-HR (19) were associated with resistance to T-20 (9, 25, 26, 28). In this study, we showed that competition between C34 and C-HR also plays a key role in acquiring resistance. However, in order to develop high resistance to C34, changes in the binding affinity of N-HR, as observed in T-20 resistance, are insufficient on their own, and an additional mutation in C-HR is necessary. According to the crystal structure of gp41 (2), none of the substitutions observed in this study are located on the binding surface of the N-HR or C-HR. Even one of the primary mutations that directly contribute to the resistance, \text{I37K}, is not located on the binding surface of C-HR. We also identified a primary mutation, N126K, that is located outside of the binding surface of C-HR. Although the significance of these locations still remains to be defined at present, further structural analysis may provide insights into the mechanism of binding.
enhancement. Our observations also provide evidence that small compounds interacting directly with the binding surface seem to inhibit HIV replication efficiently, since HIV would hardly be mutated on the binding surface.

During the selection of the C34-resistant variants, the substitutions were introduced in the following order: a substitution (N126K) associated with susceptibility to C34 was introduced first, followed by a substitution (A30V) associated with replication. L204I also improved the replication kinetics of HIV-1. It is likely that substitutions that are associated with resistance usually impair the replication kinetics, resulting in selection of HIV-1 variants containing substitutions that improve the replication disadvantages. This hypothesis has been proved with analyses of replication kinetics of T20-resistant variants described previously (20). Moreover, such substitution patterns have previously been observed in multi-dideoxyribonucleoside-resistant variants (15, 21). However, the mechanism of replication improvement in multi-dideoxyribonucleoside-resistant variants remains unknown. In this study, I37 is one of the key amino acids for C34 resistance, and it is located in an important region for the Rev-RRE interaction (13). The significance of the secondary mutations, A30V and D36G, for improvement of the RRE structural stability impaired by I37T or I37K is thought to be that they maintain both gp41 and RRE functions. In contrast to the C34 resistance mutations, nucleotides encoding some T20 resistance mutations, L33S and N43K (9), are located in a single-stranded bulge region of the stem IIc loop top (UUA to UCA) (Fig. 3A) and in the bulge region of stem III, indicating that the structural changes to the RRE would be minimal, while other T20 resistance mutations (20), such as G36D, G36S, I37K, and A30V and Q39H. These substitutions are also observed in some treatment-naive clinical isolates (16). It is well known that HIV reverse transcriptase makes several nucleotides miss incorporation during the reverse transcription, suggesting that each HIV isolate, even in the wild-type population, contains several substitutions in the integrated DNA genome. D36 is identified only in pNL4-3-derived clones, although the G36/ I37/V38 motif is well conserved, not only in HIV-1 but also in HIV-2 and simian immunodeficiency virus strains (16). Furthermore, the 5-amino-acid deletion in gp120 was reported not only in a fusion inhibitor, T20 (9), but also in CD4-gp120-binding inhibitors DSS000 (8) and AR177 (Zintev) (7), CXCR4 antagonists, bicyclams JM2763 and SID791 (6), and SDF1α-resistant variants (27). In these reports, pNL4-3-derived viruses were also used for the selection of the resistant variants. Only pNL4-3 has the 5-amino-acid tandem sequence FNSTWFNSTW in the gp120 V4 region. Therefore, this deletion is thought to be specific for HIV-1 and also in HIV-2 and simian immunodeficiency virus strains (16). Furthermore, the 5-amino-acid deletion conferred weak C34 resistance. These results indicate that we should be careful before concluding that such substitutions are involved in the resistance or replication kinetics.

In conclusion, HIV acquires resistance against C34 by mutations in both N-HR and C-HR. However, mutations in N-HR are restricted by Rev-RRE and/or gp120-gp41 interactions, suggesting that HIV-1 fusion is one of the most attractive targets for blocking HIV infection.

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