Viability of a drug-resistant human immunodeficiency virus type 1 protease variant: structural insights for better antiviral therapy

Moses Prabu-Jeyabalan
University of Massachusetts Medical School

Let us know how access to this document benefits you.
Follow this and additional works at: https://escholarship.umassmed.edu/oapubs

Repository Citation

This material is brought to you by eScholarship@UMassChan. It has been accepted for inclusion in Open Access Publications by UMass Chan Authors by an authorized administrator of eScholarship@UMassChan. For more information, please contact Lisa.Palmer@umassmed.edu.
Viability of a Drug-Resistant Human Immunodeficiency Virus Type 1 Protease Variant: Structural Insights for Better Antiviral Therapy

Moses Prabu-Jeyabalanan, Ellen A. Nalivaika, Nancy M. King, and Celia A. Schiffer*

Department of Biochemistry and Molecular Pharmacology, University of Massachusetts Medical School, Worcester, Massachusetts 01605

Received 28 May 2002/Accepted 11 October 2002

Under the selective pressure of protease inhibitor therapy, patients infected with human immunodeficiency virus (HIV) often develop drug-resistant HIV strains. One of the first drug-resistant mutations to arise in the protease, particularly in patients receiving indinavir or ritonavir treatment, is V82A, which compromises the binding of these and other inhibitors but allows the virus to remain viable. To probe this drug resistance, we solved the crystal structures of three natural substrates and two commercial drugs in complex with an inactive drug-resistant mutant (D25N/V82A) HIV-1 protease. Through structural analysis and comparison of the protein–ligand interactions, we found that Val82 interacts more closely with the drugs than with the natural substrate peptides. The V82A mutation compromises these interactions with the drugs while not greatly affecting the substrate interactions, which is consistent with previously published kinetic data. Coupled with our earlier observations, these findings suggest that future inhibitor design may reduce the probability of the appearance of drug-resistant mutations by targeting residues that are essential for substrate recognition.

The development of the human immunodeficiency virus type 1 (HIV-1) protease inhibitors is regarded as a major success of structure-based drug design (16, 36, 62–64). Indeed, the protease inhibitors are considered the most potent drugs currently available for the treatment of AIDS (63). These drugs are often combined with other drugs to establish highly active antiretroviral therapy, which is credited with an approximately threefold drop in the death rate from AIDS since its introduction (38). Despite this remarkable success, the emergence of HIV-1 mutants that resist current drug regimes (1, 2, 14) remains a critical factor in clinical failure of antiviral therapy (9, 56). The relatively rapid appearance of resistant viral mutants among treated HIV-1 patients is attributable to the high rate of replication of the virus, coupled with a high intrinsic rate of mutation due to the infidelity of the HIV-1 reverse transcriptase (22, 46, 47).

The homodimeric HIV-1 protease is an effective therapeutic target because it allows viral maturation by sequentially cleaving at least 10 asymmetric and nonhomologous sequences in the Gag-Pol polyproteins (8, 19, 42). The six Food and Drug Administration-approved HIV-1 protease inhibitors, amprenavir (APV), indinavir (IDV), nelfinavir (NFV), saquinavir (SQV), ritonavir (RTV), and lopinavir (LPV), that are on the market are all competitive inhibitors (17), binding at the active site. Therefore, they compete directly with the enzyme’s ability to recognize substrates (33, 34, 43, 44, 49, 60). These drugs are peptidomimetics that resulted from structure-based drug design efforts (7, 23–25, 55, 59). All of them have large, generally hydrophobic moieties that interact with the mainly hydrophobic S2–S2′ pockets in the active site (62). Despite chemical differences, these inhibitors occupy a similar space in the active site, and hence similar mutations in HIV-1 protease can cause multidrug resistance without substantially altering substrate binding (1, 45, 51).

Since the primary function of HIV-1 protease is to cleave its substrates rather than bind inhibitors, our laboratory analyzed the substrate recognition of this enzyme and attempted to arrive at a structural rationale for what constitutes a substrate. We determined the crystal structures of peptides bound to an inactive wild-type HIV-1 protease (44). Analysis of these complexes found that all the conserved protease–substrate hydrogen bonds involve only backbone atoms of the substrate and therefore are unlikely to be the primary determinant of substrate specificity; also, the conformation of the unprimed side of the peptide is conserved and asymmetric, resembling a toroid in the S1–S3 pockets, while the prime side of the peptide remains extended. Based on these two observations, we proposed that the major determinant of specificity for HIV-1 protease is that the conformation of the unprimed side of the substrate can form a toroid.

To further understand the substrate binding, drug resistance, and protein adaptability of HIV-1 protease (26, 43, 44, 53), we examined one of the first drug-resistant mutations to occur in HIV-1 protease among patients receiving antiviral therapy, V82A (54). This mutation occurs particularly in those patients receiving IDV or RTV (9, 10, 12, 39). Structurally, the Val82 residue is located at the P1-loops (Gly78–Asn83) near the active site (see Fig. 1a). The effect of this mutation on the binding to RTV, IDV, and NFV inhibitors is to significantly reduce their affinity (18, 27), whereas in contrast, the effect of this mutation on the ability of the enzyme to cleave substrates is smaller (18, 27). Understanding how the drug-resistant protease recognizes both substrates and inhibitors becomes crucial for future rational drug design. To address this issue, we have determined the crystal structures of the inactive drug-resistant variant D25N/V82A of HIV-1 protease in complex with either...
of three Gag peptide substrates—matrix-capsid (MA-CA), capsid-p2 (CA-p2), and p1–p6—and two clinically used drugs—SQV and RTV. We conducted structural analysis using these complexes in relevance to ligand-protease interaction, which includes shape complementarity, hydrogen bonds, van der Waals (VDW) packing, and estimation of VDW interaction energy. Since the wild-type structures for these ligands are already available in the Protein Data Bank (7, 23–25, 55), it allowed us to make a direct structural comparison. The findings of this study suggest that Val82 is not crucial for substrate recognition but is critical for inhibitor binding.

**MATERIALS AND METHODS**

**Nomenclature.** Ligand acronyms with corresponding subscripts, such as V82A for the mutant or WT for wild-type protease, are used to distinguish HIV-1 protease variants throughout this article. For example, p1–p6V82A denotes p1–p6 cleavage site peptide complexed to the V82A mutant protease.

**Mutagenesis, protein purification, and crystallization.** The protease gene containing the D25N modification was a gift from C. S. Craik, University of California San Francisco; the D25N mutation is required for complexing protease with its substrates without cleaving them. The alanine mutation at position 82 was introduced by using the QuickChange site-directed mutagenesis kit (Stratagene, La Jolla, Calif.). The protocol used for protein overexpression and purification has been described previously (21, 43).

The purified protein was concentrated to 2.5 mg ml⁻¹ and equilibrated on ice for 30 min with a fivefold molar excess of the ligand. Crystals were obtained in hanging drops under more than one condition. The crystals used here were grown with a reservoir solution of 126 mM phosphate buffer (pH 6.2), 63 mM sodium citrate, and ammonium sulfate (28 to 31% and 22% for the substrates and inhibitors, respectively) (43, 57). The crystals were brick shaped, tiny, and colorless, with a maximum dimension between 0.1 and 0.2 mm; the protein concentration ranged between 0.4 and 1.9 mg ml⁻¹.

**Data collection.** Intensity data were collected by using a Rigaku R-axis image plate mounted on a Rigaku rotating-anode X-ray generator, with the latter operating at 50 kV and 100 mA. The Yale mirror system was used to focus the X-ray beam. Data for the CA-p2V82A, p1–p6V82A, and SQVV82A complexes were collected at 25°C, while data for the MA-CAV82A and RTVV82A complexes were collected at −80°C. The crystal system and cell dimensions were determined using the program DENZO (37, 40a). Indexing of reflections revealed that crystals for all of the complexes grew in a similar P2₁ 2₁ 2₁ cell, except for the RTVV82A crystals, which grew in a P2₁ cell with β = 99.8°. However, its unit cell vectors, a, b, and c, were similar to those of the remaining complexes. The final data collection statistics for the five structures in this study are shown in Table 1.

**Structure solution and crystallographic refinement.** Structure solution and crystallographic refinement for all five structures were carried out using the Crystallography & NMR System version 0.9 (5). A previously published inhibitor complex (PDB:1MTR) (35) was used as the starting model for solving substrate
TABLE 1. Data collection and refinement statistics for the five proteaseV82A complexes

<table>
<thead>
<tr>
<th>Variable</th>
<th>MA-CA</th>
<th>CA-p2</th>
<th>p1-p6</th>
<th>SQV</th>
<th>RTV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ligand sequence</td>
<td>VSQNY-PIVQN</td>
<td>KARVL-AEAMS</td>
<td>RPGNF-LQSRP</td>
<td>See Fig. 3</td>
<td>See Fig. 3</td>
</tr>
<tr>
<td>Data collection</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Space group</td>
<td>P2₁2₁2₁</td>
<td>P2₁2₁2₁</td>
<td>P2₁2₁2₁</td>
<td>P2₁2₁2₁</td>
<td>P2₁</td>
</tr>
<tr>
<td>a (Å)</td>
<td>50.97</td>
<td>51.56</td>
<td>51.27</td>
<td>51.44</td>
<td>51.62</td>
</tr>
<tr>
<td>b (Å)</td>
<td>57.86</td>
<td>59.38</td>
<td>59.13</td>
<td>60.04</td>
<td>59.04</td>
</tr>
<tr>
<td>c (Å)</td>
<td>62.07</td>
<td>61.87</td>
<td>61.93</td>
<td>62.02</td>
<td>61.35</td>
</tr>
<tr>
<td>β (°)</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>99.8</td>
</tr>
<tr>
<td>Z</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Temp (°C)</td>
<td>−80</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td>−80</td>
</tr>
<tr>
<td>Resolution (Å)</td>
<td>1.9 (1.97–1.9)</td>
<td>2.15 (2.23–2.15)</td>
<td>2.0 (2.07–2.0)</td>
<td>2.5 (2.59–2.5)</td>
<td>2.2 (2.28–2.5)</td>
</tr>
<tr>
<td>Total no. of reflections</td>
<td>48,168</td>
<td>25,200</td>
<td>98,506</td>
<td>28,072</td>
<td>19,193</td>
</tr>
<tr>
<td>No. of unique reflections</td>
<td>14,278 (1086)</td>
<td>10,094 (834)</td>
<td>12,659 (1182)</td>
<td>6,743 (496)</td>
<td>13,097 (1183)</td>
</tr>
<tr>
<td>Rmerge (%)</td>
<td>5.7 (19)</td>
<td>7.4 (31)</td>
<td>9.0 (33)</td>
<td>8.0 (36)</td>
<td>6.4 (27)</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>94.9 (74)</td>
<td>95.2 (76)</td>
<td>95.4 (92)</td>
<td>93.4 (71)</td>
<td>71.2 (66)</td>
</tr>
<tr>
<td>I/σ(I)</td>
<td>5.8</td>
<td>7.5</td>
<td>7.7</td>
<td>7.0</td>
<td>7.6</td>
</tr>
<tr>
<td>Refinement</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R value (%)</td>
<td>20.1 (26)</td>
<td>20.8 (32)</td>
<td>17.9 (22)</td>
<td>19.1 (30)</td>
<td>22.3 (32)</td>
</tr>
<tr>
<td>Rfree (%)</td>
<td>22.7 (32)</td>
<td>24.3 (34)</td>
<td>21.0 (29)</td>
<td>25.7 (36)</td>
<td>28.4 (37)</td>
</tr>
<tr>
<td>Sigma cutoff</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>RMSD in:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bond lengths (Å)</td>
<td>0.005</td>
<td>0.006</td>
<td>0.005</td>
<td>0.008</td>
<td>0.008</td>
</tr>
<tr>
<td>Bond angles (°)</td>
<td>1.3</td>
<td>1.4</td>
<td>1.8</td>
<td>1.3</td>
<td>1.4</td>
</tr>
</tbody>
</table>

* Values in parentheses are the statistics corresponding to the highest-resolution shell.

b RMSD root mean square deviation.

Structural analysis. (i) Difference distance plots. The difference distance plots are constructed to assess intramolecular relative shifts in response to ligand binding (26). Initially, Cα-Cα distances (dα) were computed between all of the CA atoms within a protease dimer of a complex, A, and repeated for a second complex, B, with which the first structure is compared. The difference in these distances (Dα = |dα|A − |dα|B) was plotted as a two-dimensional contour plot as a function of residue numbers (i and j) by using GNUPLOT (61).

(ii) Buried surface area. The classic Lee and Richards program (31) was used to evaluate the surface area buried by the ligands on complexation. Accessible surface area (ASA) was computed for each of the ligand-bound protease structures. The ASA calculation was repeated for the unbound ligands in the absence of the protease coordinates as well. The difference in these two ASA values is defined as the surface area buried on complexation.

(iii) Shape complementarity. Besides buried surface area, another crucial feature that has been suggested to determine the specificity and stability of protein-ligand association is shape complementarity (SC) (30). SC is related to the dot product of the normals of the apposing surfaces and the exponent of the distances between them (for the complete mathematical formulation, see reference 30). The protocol includes the following: construction of molecular surface for both ligands and protein, calculation of normal vectors (nligand and nprotein) over a user-designated grid space, and computation of the intermolecular distance matrix between each intermolecular grid point (rilm = rligand − rprotein). The SC by the ligand on the protein is then given by the relationship SCligand−protein = (nligand·nprotein)exp[−rilm/γ], where γ is a parameter that controls the shape complementarity. This SC is then computed for each grid point, summed, and averaged. The calculation is repeated to obtain the SC protein−ligand. Finally, the average of the two SC values is the actual SC. In this analysis, we quote SC values as SCV82A/SCWT ratios.

(iv) VDW interaction energy. Estimates of VDW interaction energy were computed to provide a theoretical quantitative assessment for the ligand-protein nonbonded interactions. VDW energy between ith and jth atoms is calculated using the Lennard-Jones potential function. The Crystallography & NMR System was used to perform this task; it utilizes data from reference 13 to obtain interatomic distances and incorporates the PROLSQ REPEL function (20) for treating nonbonded interactions.

RESULTS

Overall structural features. Three decameric substrate peptides from the Gag cleavage sites of matrix-capsid (MA-CA: VSQNY-PIVQN), capsid-p2 (CA-p2: KARVL-AEAMS) and p1-p6 (RPGNF-LQSRP) and two protease inhibitors, SQV and RTV, were crystallized in complex with the inactive (D25N) variant of HIV-1 protease with the drug resistant V82A mutation. The crystallographic statistics for these structures are listed in Table 1. The inhibitors and the substrate peptides were unambiguously located in the two difference Fourier maps, 2Fobs − F and Fcal − Fobs, and each of the ligands was uniquely oriented within the protease dimer. This uniqueness
in the orientation is crucial for accurately determining ligand-protease interactions.

As in wild-type HIV-1 protease-substrate complexes (43, 44) and protease-substrate analogue structures (33, 34, 60), the substrate peptides in the V82A complexes also assume a relatively extended conformation (Fig. 1a and b). Also preserved in the V82A complexes is the toroid shape on the unprimed side of the substrate (Fig. 1c). This shape is achieved by the packing of P3 and/or P4 against the P1 side chain. In spite of having a glycine at P3, as seen in the corresponding wild-type complex, the p1–p6V82A peptide also forms a toroid through changes in the peptide backbone conformation (Fig. 1b). In contrast, the shapes of the five commercially available drugs, as observed in the wild-type complexes (7, 23–25, 55) do not form this toroid (Fig. 1d). Thus, the inhibitors do not mimic the conserved structural motif exhibited by the substrate sequences.

To probe the effect of the V82A mutation, we compared the structures of the ligandV82A complexes with their wild-type counterparts. To elucidate the distant structural alterations (26), difference distance plots were generated between the V82A and wild-type structures (24, 44, 55) of each ligand (Fig. 2). The plots involving substrate structures indicate only minor deviations (Fig. 2a and b), while the inhibitorV82A complexes, on the other hand, exhibit substantial shifts from the corresponding inhibitorWT structures, especially near the protease flaps (Lys45–Ile54) (Fig. 2c and d). Thus, the three substrateV82A complexes are very structurally similar to their substrateWT complexes, while inhibitorV82A complexes are not.

Protease-ligand hydrogen bonds. In protein-ligand interactions, hydrogen bonds play a crucial role in stabilizing the complex and, in many instances, also determining specificity. Analysis of the wild-type protease-ligand hydrogen bonds involving five inhibitors and six substrates suggests that the substrates form almost twice the number of hydrogen bonds with the protease (about 16) compared with the inhibitors (about 9) (Fig. 3a). Furthermore, the number of hydrogen bonds in each substrateWT complex is similar to that in the substrateV82A complexes whereas four hydrogen bonds were lost between the inhibitorWT and inhibitorV82A complexes. This difference (Fig. 3a) indicates that the mutation has specifically affected protease-inhibitor hydrogen bonding.

As previously observed in substrateWT complexes (44), the hydrogen bonds conserved among these three substrateV82A complexes involve only the backbone atoms of the peptides (Fig. 3b). Nine hydrogen bonds are conserved among the three substrateV82A complexes, eight of which are conserved among all six substrateWT complexes as well. Thus, the conserved hydrogen bonding pattern present among the substrateWT complexes is retained among the substrateV82A complexes as well, indicating that the protease-substrate hydrogen bonding is not perturbed by the V82A mutation.

In contrast, the inhibitorWT complexes, SQVWT and RTVWT, make 13 and 11 protease-inhibitor hydrogen bonds, respectively, 9 of which are conserved between them (Fig. 3c and d). In the SQVV82A and RTVV82A complexes, however, there are nine and seven protease-inhibitor hydrogen bonds, respectively, five of which are conserved between them. The catalytic aspartic acids, Asp25/25', make four hydrogen bonds with one atom each of the inhibitor in the inhibitorWT complexes, and these four hydrogen bonds are also seen in the inhibitorV82A complexes with nearly identical interatomic distances. The conservation of these four hydrogen bonds supports the likelihood that the D25N mutation has not caused any change in the complex. A large-scale shift that is seen in the protease flaps (1.2 Å in the RTVV82A relative to the RTVWT complex) and that is in the proximity of Asn25/25' substantially alters hydrogen bonding patterns in the inhibitorV82A complexes. Thus, the differences in protease-inhibitor hydrogen bonding patterns between the inhibitorWT and the inhibitorV82A complexes unambiguously reveal that inhibitor binding has been substantially affected in the V82A drug-resistant protease.

VDW interactions. The surface area buried by the ligands on binding the protease molecule, which is an indicative parameter of the extensiveness of the ligand-protein interface, was computed by the method of Lee and Richards (31) (Table 2). Typically, substrates buried 850 to 1000 Å² of their accessible surface area while the inhibitors account for only 500 to 650 Å². This amounts to a 30 to 40% reduction of buried surface area by the inhibitors relative to the substrates. Although the change in buried surface area due to mutation is not very significant (Table 2), the difference in area between the substrates and inhibitors indicates that the substrate-protease association is more extensive than the inhibitor-protease interface.

Complementation of shape by the VDW surface of protease residue 82 with the VDW surfaces of the ligands is likely to provide further evidence for the influence of mutation (Fig. 4a and b). The VDW surface of residue 82 (valine or alanine) does not make any contact with VDW surface on the primed side of the substrates, P1′–P3′ (Fig. 4a). On the unprimed side of the peptide, however, Ala82' contacts all the substrates tangentially while Val82' contacts only PheP1 in the p1–p6WT complex, closely resembling the pattern observed in the p1–p6V82A complex. While the SQVWT and RTVWT complexes exhibit a high degree of packing between the VDW surfaces of the inhibitor and Val82/82' (Fig. 4b), the VDW surfaces of both Alav82 and Alav82' fail to contact those of the inhibitor in the SQVV82A and RTVV82A complexes (Fig. 4b).

In particular, VDW packing in the RTVV82A complex is achieved by both side chain atoms of Val82', CG1 and CG2, contacting the isopropyl group of the 2-isopropyl-4-thiazolyl at the P3 position of the drug. In addition, the isopropyl group extends the farthest at this site compared with any drug or substrate probed in this investigation (Fig. 1b). This isopropyl group appears to have been designed to target the pocket by the Val82 residue, and when an alanine mutation occurs, the P3 group is no longer complementary (Fig. 4b).

SQV, on the other hand, due to its similarities to the naturally occurring substrates, at least in the P2–P2' region (Fig. 1b), is less vulnerable to V82A mutation. In fact, the computed shape complementarity (30) decreases by only 18% in SQV compared with 28% in RTV for the inhibitorV82A relative to corresponding wild-type complexes. However, with the exception of the MA-CA complexes, the substrate-protease shape complementarity between the wild type and V82A are very similar (Table 2).

The effect of mutation was further investigated by computing ligand-protease VDW interaction energy (Table 2). A comparison of the estimated VDW interaction energy between the
wild-type and V82A substrate structures shows a difference of less than 1.5 kcal/mol. The SQV\textsubscript{V82A} complex, surprisingly, decreased VDW interaction energy by approximately 12 kcal/mol. However, this is probably because SQV\textsubscript{WT} complex (55) was refined with an earlier force field in X-PLOR 2.1 and even small differences in VDW radii can lead to substantial differences in the estimates of energies if different force fields are used. In the RTV\textsubscript{V82A} structure, where the refinement force field is the same as that used to refine the RTV\textsubscript{WT} complex (24), the increase in computed VDW interaction energy, by nearly 10 kcal/mol, indicates that RTV\textsubscript{V82A} is less favorable than the RTV\textsubscript{WT} complex.

FIG. 2. Double difference plots. Relative shifts in the V82A complexes in reference to the corresponding wild-type complexes. (a) CA-p2; (b) p1-p6; (c) SQV; (d) RTV. Each contour line represents a deviation by 0.25 Å. The different colors black, green, red, and blue distinguish the contour ranges −1.0 Å and below, −0.5 to −1.0 Å, 1.0 Å and above, and 0.5 to 1.0 Å, respectively.
The number of substrate atoms within a VDW radius of residue 82 was compared in nine substrate complexes (three substrateV82A and six substrateWT complexes [43, 44]) and seven inhibitor structures (two inhibitor V82A and five inhibitorWT structures [7, 23–25, 55]) (Fig. 4c). In all drug complexes except APV, six or more atoms of the drug molecule surround the side chain atoms of Val82 (Fig. 4c), explaining the importance of Val82 for inhibitor binding. In contrast, five or fewer atoms of the substrate peptides surround Val82 in most of the substrateWT complexes. The only substrate complex for which this is an exception is the p1–p6WT complex, where 10 substrate atoms lie within the VDW contact distance of the Val82 side chain.

The number of substrate atoms within a VDW radius of residue 82 was compared in nine substrate complexes (three substrateV82A and six substrateWT complexes [43, 44]) and seven inhibitor structures (two inhibitor V82A and five inhibitorWT structures [7, 23–25, 55]) (Fig. 4c). In all drug complexes except APV, six or more atoms of the drug molecule surround the side chain atoms of Val82 (Fig. 4c), explaining the importance of Val82 for inhibitor binding. In contrast, five or fewer atoms of the substrate peptides surround Val82 in most of the substrateWT complexes. The only substrate complex for which this is an exception is the p1–p6WT complex, where 10 substrate atoms lie within the VDW contact distance of the Val82 side chain.

### Table 2. Comparison of the structural analysis performed on the five V82A complexes and their wild-type equivalents

<table>
<thead>
<tr>
<th>Structure</th>
<th>Surface area buried by ligand on complexation (Å²)</th>
<th>Ratio in shape complementarity V82A/WT</th>
<th>No. of ligand-protease VDW contacts</th>
<th>Ligand-protease estimated VDW interaction energy (kcal/mole)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>WT</td>
<td>V82A</td>
<td>WT</td>
<td>V82A</td>
</tr>
<tr>
<td>MA-CA</td>
<td>928</td>
<td>867</td>
<td>0.83</td>
<td>174 (5)</td>
</tr>
<tr>
<td>CA-p2</td>
<td>977</td>
<td>923</td>
<td>1.00</td>
<td>167 (1)</td>
</tr>
<tr>
<td>p1-p6</td>
<td>1,029</td>
<td>1,038</td>
<td>1.07</td>
<td>197 (10)</td>
</tr>
<tr>
<td>SQV</td>
<td>612</td>
<td>630</td>
<td>0.82</td>
<td>165 (7)</td>
</tr>
<tr>
<td>RTV</td>
<td>526</td>
<td>500</td>
<td>0.72</td>
<td>165 (11)</td>
</tr>
</tbody>
</table>

*The structural analysis includes surface area buried by the ligand on complexation, shape complementarity, number of ligand-protease VDW contacts, and ligand protease estimated VDW interaction energy. The numbers shown in parentheses highlight the number of atoms located within VDW radius distance of residue 82 in the corresponding structures.

² The distance criterion used for the VDW contacts is 2.4 to 4.2 Å.

³ In the comparison of the peptide complexes, only equivalent residues were used, since the termini were disordered in some of the crystal structures.

⁴ The refinement of SQVWT was carried out using X-PLOR version 2.1 which had slightly different non-bonded energy parameters.
FIG. 4. Protease-ligand VDW interactions. (a and b) VDW surfaces of residue 82, viewed down the dimer twofold axis, and the neighboring atoms of the corresponding substrates (a) and inhibitors (b). The ligands and the protease are shown in gray and black, respectively. A Ca trace of the overlying flaps is also shown. (c) Histogram representation of numbers of ligand atoms within VDW contact of residue 82 in the crystal structures determined here. The number of atoms in the substrateWT and inhibitorWT complexes are shown in white, and those in the substrateV82A and inhibitorV82A complexes are shown in black. Abbreviations are defined in the legend to Fig. 3.
side chain. The p1–p6V82A is also the only complex where contact of seven atoms within VDW radii of Ala82 is lost, thus reducing the VDW interaction considerably (Fig. 4c) compared with the p1–p6WT complex. However, this reduction seems to be compensated in the p1–p6V82A complex by the conservation of the total number of substrate-protease VDW contacts (Table 2). This compensation is achieved by subtle side chain alterations undergone by Asn at the P2 position of the p1–p6 substrate in combination with slight changes in Val32 and Ile47 of the protease. Thus, as seen in the CA-p2WT structure (43), the protease as well as its substrates are highly adaptable to each other.

In comparison with the inhibitor WT complexes, the inhibitorV82A complexes have far fewer atoms making VDW contacts with Ala82 (Fig. 4c). Only one atom comes within VDW contact of Ala82 in the SQVV82A complex, while seven atoms surround Val82 in the SQVWT complex. The decrease is even larger in the RTVV82A complex, where only 2 atoms come within the VDW distance of Ala82, while 11 atoms surround Val82 in the RTVWT complex. This clustering of atoms around Val82 in the inhibitor complexes suggests that the inhibitors were designed to contact the valine residue.

As an extension to VDW contacts by Val82 (or Ala82) with ligands, assessment of VDW contacts between the ligands and the entire protein is also crucial. The change in the number of substrate-protease VDW contacts was either absent (as in Ca-p2 and p1–p6) or nominal (as in MA-CA) (Table 2). However, significant loss in the number of inhibitor-protease VDW contacts was observed between the SQV complexes (165 in SQVWT and 142 in SQVV82A) (Table 2) and the RTV complexes (165 in RTVWT and 130 in RTVV82A). Thus, the overall effect of V82A mutation in the substrate complexes is minimal compared to its affect on the inhibitor complexes.

**DISCUSSION**

Our results reveal high structural similarity among the three V82A substrate complexes, CA-p2V82A, p1–p6V82A, and MA-CAV82A, and their previously reported wild-type counterparts (43, 44), particularly around the active-site region, suggesting that the V82A mutation has not influenced substrate binding. Conformational changes were observed, however, at the active site between the inhibitorV82A structures and their previously published wild-type counterparts (24, 55), suggesting that the protease mutation has indeed influenced inhibitor binding. In particular, the movement by the protease flaps in the RTVV82A complex by more than 1 Å in comparison to the RTVWT complex (24) is another indication of modified inhibitor binding. Difference distance plots show almost no shifts in the backbone as a result of V82A on substrate recognition (Fig. 2a and b) but show significant shifts in the inhibitor complexes (Fig. 2c and d). Thus, the V82A mutation appears to minimally perturb substrate binding while strongly affecting inhibitor binding.

Protease-ligand hydrogen bonds are crucial for retaining the stability of the complexes. The lack of difference in protease-substrate hydrogen bonds between the substrateV82A and substrateWT complexes (Fig. 3a) suggests that the specificity and stability of protease-substrate complexes is unaltered. The inhibitorV82A complexes, on the other hand, exhibit large-scale shifts with respect to their wild-type complexes, resulting in a loss of several hydrogen bonds. The substrates, which are longer than inhibitors, form more extensive hydrogen bond network with the protease, involving a variety of its residues and encompassing the entire active site (Fig. 3b). This network is probably a key factor in allowing the protease to maintain its function despite having many mutations. Because inhibitors in the inhibitorWT complexes have so few hydrogen bond acceptors and donors, the loss of hydrogen bonds due to the V82A mutation further decreases the specificity and affinity of the complex.

The asymmetric consensus volume occupied by the substrates compared to the relatively symmetric inhibitors illustrates the crucial differences between them (Fig. 1c and d). Coupled with our earlier findings with the wild-type protease-substrate complexes (44), this observation supports the hypothesis that the asymmetric toroidal shape on the unprimed side of the substrate is important for substrate specificity. The inhibitors, however, do not possess this shape (Fig. 1c). The toroidal shape of the substrates is the only conserved structural motif exhibited by the otherwise nonhomologous cleavage site sequences and could be utilized in designing the next generation of inhibitors. In fact, the positions corresponding to the P3–P1’ sites of the substrates may be used to model inhibitors instead of using the positions of P2–P2’. A recent study has revealed that inhibitors with a macrocyclic group connecting the P3 and P1 residues are likely to be more efficient (35).

Since hydrophobic residues in HIV-1 protease have a higher propensity for conferring drug-resistant mutations (54), a study of changes in VDW interactions around the mutation site is essential for understanding drug resistance. Lack of tight pairing between the VDW surfaces of Val82/82’ and the substrate, seen in the substrateWT complexes, suggests that the valine residue may not be extensively utilized during substrate binding (Fig. 4a). The similarity between the substrateWT and substrateV82A complexes, particularly in relation to the VDW packing of Ala82/82’ with the substrates, is further supported by the conservation of the estimated VDW interaction energy (Table 2). On the other hand, the apposition of the VDW surfaces of the Val82/82’ residue and inhibitors (Fig. 4b) in the inhibitorWT complexes suggests that the alanine mutation at residue 82 will probably affect inhibitor binding. A previous
study using the V82A mutant in complex with a C2-symmetry-based diol analogue, A-77003, pointed out that the protease is highly adaptable to accommodating the valine-to-alanine alteration (1). The present study, however, reveals substantial loss of substrate recognition as contact residues for new inhibitors. Perhaps this type of directed inhibitor-design will make it more difficult for further drug resistance to evolve.

ACKNOWLEDGMENTS

Saquinavir and ritonavir were obtained through the NIH AIDS Research and Reference Reagent Program, Division of AIDS, NIADDK, NIH. We thank Charles Craik from UCSF for the original D25N plasmid of HIV-1 protease. We also thank Balaji Bhryavbhata for technical support during data collection, Vincent Chou and Nicole M. King for graphics assistance, and Claire Baldwin for editorial advice. This research was supported by the American Cancer Society (RPG-99-213-01-MBC) and the National Institutes of Health (GM64849).

REFERENCES
