

Cooperative Effects of Drug-Resistance Mutations in the Flap Region of HIV-1 Protease

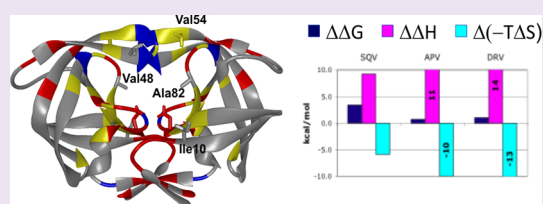
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S Supporting Information

ABSTRACT: Understanding the interdependence of multiple mutations in conferring drug resistance is crucial to the development of novel and robust inhibitors. As HIV-1 protease continues to adapt and evade inhibitors while still maintaining the ability to specifically recognize and efficiently cleave its substrates, the problem of drug resistance has become more complicated. Under the selective pressure of therapy, correlated mutations accumulate throughout the enzyme to compromise inhibitor binding, but characterizing their energetic interdependency is not straightforward. A particular drug resistant variant (L10I/G48V/I54V/V82A) displays extreme entropy–enthalpy compensation relative to wild-type enzyme but a similar variant (L10I/G48V/I54A/V82A) does not. Individual mutations of sites in the flaps (residues 48 and 54) of the enzyme reveal that the thermodynamic effects are not additive. Rather, the thermodynamic profile of the variants is interdependent on the cooperative effects exerted by a particular combination of mutations simultaneously present.



Human immunodeficiency virus 1 (HIV-1) protease is one of the prime therapeutic targets in the treatment for HIV. Protease inhibitors (PIs), especially when used in combination with other drugs in the highly active antiretroviral therapy (HAART), improve the life quality and expectancy in HIV patients.^{1–3} However, accumulation of mutations in protease under selective drug pressure decreases inhibitor binding affinity, compromising the treatment efficacy. Understanding the molecular basis of reduced inhibitor affinity and the associated thermodynamics of binding is essential for developing inhibitors resilient to resistance.

The binding affinity of a ligand to a target is dictated by the free energy of binding, comprising entropic and enthalpic contributions. Either contribution can be the driving force for binding, and potent PIs exist with both entropically and enthalpically driven binding.^{4–6} The first generation inhibitors have entropically driven binding, while the affinity of newer and more potent PIs are mostly due to favorable enthalpy. However, incorporating entropic and enthalpic considerations into rational drug design is not straightforward, as enhancing conformational entropy (increasing bound inhibitor flexibility) is often balanced against the competing enthalpy (maximizing molecular contacts). Additionally, the drug resistance mutations can perturb inhibitor binding thermodynamics in a complex and interdependent manner,^{7,8} including through altering the flexibility of the enzyme.⁹

The drug resistance mutations derived from patient sequences occur throughout HIV-1 protease, not just in the active site. The homodimeric protease is highly polymorphic; only 27 of the 99 residues do not vary either naturally or in

response to selective PI therapy¹⁰ (Figure 1). Many common drug resistant mutations occur in the flaps, which are critical for substrate binding, hence enzymatic activity.^{11,12} When HIV-1 protease binds substrates, the flaps in each monomer encase the substrate in the active site.^{13,14} The protease flap region has been the subject of extensive studies related to sequence tolerance¹⁵ and the mechanism of substrate access to the active site.^{16–20}

Flap⁺_(I54V) is a drug resistant protease variant containing a combination of active site and flap mutations (L10I/G48V/I54V/V82A)²¹ (Figure 1) that often occur together in treated patients.²¹ Two mutations in the flap region, G48V and I54V, have been well correlated with drug resistance to multiple inhibitors,^{22–25} even though Ile54 does not contact either substrates or inhibitors. I54A has a PI susceptibility profile similar to I54V, unlike other substitutions such as I54L and I54M.²⁶ Analysis of patient sequences in the Stanford HIV drug resistance database revealed that I54A also shares many of the mutations frequently observed together with I54V (detailed analysis in Supporting Information). In most of the patients with known preceding sequences, I54A evolves from I54V (Table S1, Supporting Information). Hence, the combination of mutations in the variants studied here, Flap⁺_(I54V) and Flap⁺_(I54A) (L10I/G48V/I54A/V82A), are clinically relevant and correlated.

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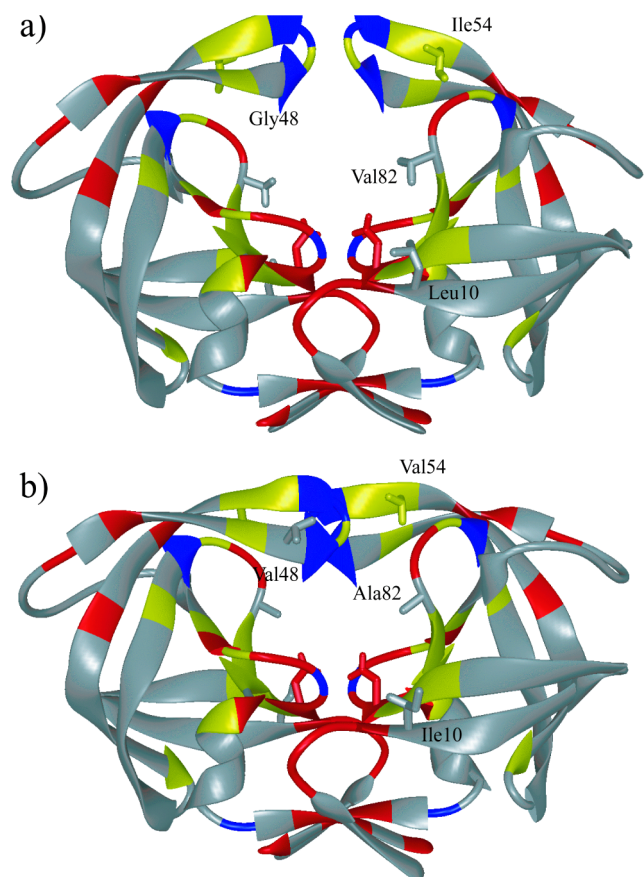


Figure 1. HIV-1 protease colored by sequence conservation.⁴¹ Residues colored red are invariant in both untreated and treated populations of patients. Residues colored yellow are invariant only in the untreated population. Invariant glycine residues are colored blue. (a) The protease structure in unliganded (PDB code: 1HHP) and (b) liganded (PDB code: 1HXB) states. The side chains of the catalytic aspartic acids Asp25 in both monomers, as well as mutation sites Leu10, Gly48, Ile54, and Val82, are displayed.

Curiously, as we previously reported,²⁷ Flap⁺_(I54V) exhibits extreme entropy–enthalpy compensation with respect to wild-type enzyme, regardless of the inhibitor bound. In the present letter, the extraordinary thermodynamic behavior of Flap⁺_(I54V) is compared with that of Flap⁺_(I54A). Additionally, the effect of individual mutations I54V versus I54A on inhibitor binding is compared.

Binding thermodynamics of the two Flap⁺ variants and three single mutants (I54V, I54A, and G48V) were measured by isothermal titration calorimetry for the protease inhibitors saquinavir (SQV), amprevir (APV), and darunavir (DRV) (Table 1; Figure S1, Supporting Information). As with wild-type protease, binding of all variants to SQV was entropically driven. Both G48V and V82A are associated with SQV resistance.²² Here, we found I54A to similarly impair SQV binding, decreasing affinity 47-fold relative to wild-type protease. I54V, however, caused a 1.9-fold decrease in affinity. When these same Ile54 mutations were present in the background of Flap⁺ variants, SQV binding affinity was impaired by 3 orders of magnitude (880- and 170-fold for Flap⁺_(I54A) and Flap⁺_(I54V), respectively). This decrease in SQV affinity in Flap⁺_(I54A) is consistent with an approximately additive effect of single mutations I54A and G48V. However,

I54V in the Flap⁺ background results in a much larger effect than a simple addition of individual mutations.

This cooperative effect of mutations in Flap⁺_(I54V) has also been observed for APV binding. In the WT protease background, single mutations I54A and I54V had the opposite effect on APV susceptibility, with I54V rendering the enzyme hyper-susceptible by 0.3-fold. In striking contrast, however, Flap⁺_(I54V) lost 3-fold affinity to APV binding, even though G48V alone did not have any considerable effect on binding affinity (0.9-fold relative to WT). Flap⁺_(I54V) was also unique in being the only variant that had an unfavorable binding enthalpy to both APV and DRV, while all other variants bound with favorable enthalpy and entropy (Table 1).

DRV binding to I54V and I54A mutants had comparable free energies, -14.8 and -14.7 kcal mol⁻¹, respectively, yet with varied enthalpic and entropic contributions. Both single I54 mutants caused about a 2-fold affinity lost, while G48V decreased DRV susceptibility 16-fold. Unlike the case for APV, Flap⁺ background did not result in a cooperative effect to impair binding affinity but profoundly changed the entropic and enthalpic contributions to the affinity. As with APV, only the Flap⁺_(I54V) variant had an unfavorable binding enthalpy to DRV. Hence, mutations in Flap⁺_(I54V) cooperatively altered the binding thermodynamics for both APV and DRV binding, yielding solely entropy-driven inhibitor binding by this variant. Interestingly, Flap⁺_(I54V) binding to APV and DRV had relative enthalpic and entropic contributions similar to WT protease binding to the first-generation inhibitor SQV.

The changes in thermodynamic parameters with respect to WT protease (Figure 2) further illustrate that the combined effect of mutations in Flap⁺_(I54V) is greater than a simple summation of the changes induced by the individual mutations G48V and I54V. G48V and I54V altered the enthalpy and entropy of binding to APV by less than 1 kcal mol⁻¹. However, the enthalpy and entropy of binding to the same inhibitor for Flap⁺_(I54V) changed by 11 and 10 kcal mol⁻¹, respectively. This cooperative effect was also observed for binding DRV: Flap⁺_(I54V) had a larger change in enthalpy and entropy with respect to WT (14.1 and -13.1 , kcal mol⁻¹, respectively) than the sum of the changes observed for G48V or I54V (Figure 2). Besides G48V or I54V, Flap⁺_(I54V) has an additional active site mutation V82A, as does Flap⁺_(I54A). However, these dramatic changes in the enthalpy and entropy of binding for Flap⁺_(I54V) are not due to the V82A mutation because these changes are much less pronounced for the binding of the same inhibitors to Flap⁺_(I54A), which contains the same additional mutation.

Thus, the extreme entropy–enthalpy compensation previously reported for Flap⁺_(I54V)²⁷ is not due to any individual mutation or a simple additive effect of these mutations, but rather a result of cooperative effects when both flap mutations, G48V and I54V, simultaneously occur. Both Gly48 and Ile54 are in the flap region of HIV-1 protease, but these two residues are not in physical contact with each other (Figure 3). This kind of cooperativity among residues that are not in contact with each other has been previously reported.^{7,28} However, the causes and molecular basis of such effects are yet to be elucidated.

The two flap mutations might exert their effects by altering the dynamic movements of this critical region. The flexibility of the flaps is central to protease activity, controlling the access of substrates and inhibitors to the active site. The flaps in unliganded protease is the most dynamic region, sampling a wide range of conformations.^{18,20,29,30} Gly48 is at the tip of the

Table 1. Thermodynamic Parameters of Binding for Each Protease Variant to SQV, APV, and DRV at 20°C

variant	K_a (M^{-1})	K_d (nM)	K_d ratio	ΔH kcal mol $^{-1}$	$-T\Delta S$ kcal mol $^{-1}$	ΔG kcal mol $^{-1}$
SQV						
wild-type ^a	$(2.0 \pm 0.1) \times 10^9$	0.50 ± 0.03	1	3.6 ± 0.1	-16.1 ± 0.1	-12.5 ± 0.1
Flap+(I54V) ^b	$(5.7 \pm 0.1) \times 10^6$	170 ± 4	350	12.9 ± 0.1	-21.9 ± 0.1	-9.1 ± 0.1
I54V	$(1.1 \pm 0.6) \times 10^9$	0.93 ± 0.6	1.9	7.0 ± 0.2	-19.1 ± 0.4	-12.1 ± 0.3
Flap+(I54A)	$(1.1 \pm 0.1) \times 10^6$	880 ± 40	1800	10.7 ± 0.3	-18.8 ± 0.3	-8.1 ± 0.1
I54A	$(4.3 \pm 0.3) \times 10^7$	23 ± 2	47	8.0 ± 0.1	-18.2 ± 0.1	-10.2 ± 0.1
G48V	$(5.8 \pm 0.3) \times 10^7$	17 ± 1	34	6.9 ± 0.1	-17.3 ± 0.1	-10.4 ± 0.1
APV						
wild-type ^c	$(2.6 \pm 1.3) \times 10^9$	0.39 ± 0.20	1	-7.3 ± 0.9	-5.3 ± 0.9	-12.6 ± 0.3
Flap+(I54V) ^b	$(7.6 \pm 0.1) \times 10^8$	1.31 ± 0.01	3.3	3.3 ± 0.5	-15.2 ± 0.5	-11.9 ± 0.1
I54V	$(7.4 \pm 0.4) \times 10^9$	0.14 ± 0.01	0.3	-7.2 ± 0.5	-6.0 ± 0.5	-13.2 ± 0.1
Flap+(I54A)	$(1.0 \pm 0.1) \times 10^9$	0.99 ± 0.09	2.5	-7.0 ± 0.8	-5.1 ± 0.8	-12.1 ± 0.1
I54A	$(1.5 \pm 0.1) \times 10^9$	0.66 ± 0.04	1.7	-7.3 ± 0.2	-5.0 ± 0.2	-12.3 ± 0.1
G48V	$(2.9 \pm 0.3) \times 10^9$	0.34 ± 0.03	0.9	-7.4 ± 0.5	-5.3 ± 0.5	-12.7 ± 0.1
DRV						
wild-type ^c	$(2.2 \pm 1.1) \times 10^{11}$	0.0045 ± 0.002	1	-12.1 ± 0.9	-3.1 ± 0.9	-15.2 ± 0.3
Flap+(I54V) ^b	$(3.9 \pm 0.7) \times 10^{10}$	0.026 ± 0.005	5.7	2.0 ± 0.6	-16.2 ± 0.6	-14.2 ± 0.1
I54V	$(1.2 \pm 0.3) \times 10^{11}$	0.0083 ± 0.002	1.9	-8.9 ± 0.3	-5.9 ± 0.4	-14.8 ± 0.1
Flap+(I54A)	$(1.4 \pm 0.2) \times 10^{10}$	0.074 ± 0.01	16	-7.0 ± 0.4	-6.6 ± 0.4	-13.6 ± 0.1
I54A	$(9.5 \pm 0.1) \times 10^{10}$	0.011 ± 0.001	2.3	-10.6 ± 0.1	-4.1 ± 0.1	-14.7 ± 0.1
G48V	$(1.4 \pm 0.1) \times 10^{10}$	0.073 ± 0.004	16	-10.3 ± 1.6	-3.3 ± 1.6	-13.6 ± 0.1

^aFoulkes et al., 2006.³⁹ ^bKing et al., 2012.²⁷ ^cKing et al., 2002.⁴⁰

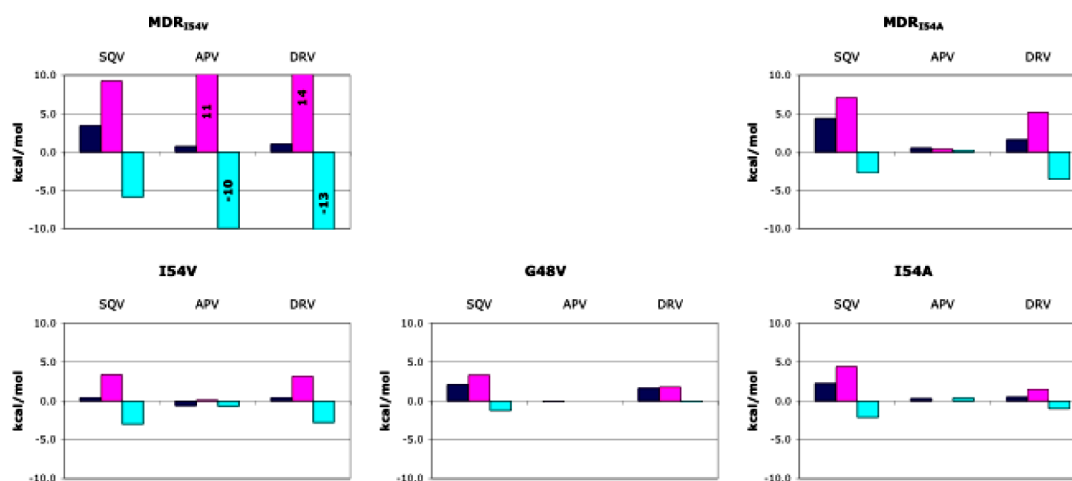


Figure 2. Changes in the thermodynamic parameters of binding relative to wild-type (WT) protease. The dark blue, magenta, and cyan bars are changes in the free energy ($\Delta\Delta G$), enthalpy ($\Delta\Delta H$), and entropy ($\Delta(-T\Delta S)$) of binding, respectively.

flap, and a mutation to valine would significantly restrict the backbone flexibility at this site, thereby altering the conformational range of the tip of the flap. Ile54 is in contact with Ile50 of the other monomer in crystal structures of the bound protease (Figure 3). Even in unliganded protease, the flaps sample a minor population where the flap tips contact each other.³¹ I54V mutation may be destabilizing this minor conformation, further restricting the conformational range of the flaps. Hence, the observed cooperativity of G48V and I54V mutations is most likely caused by their effects on flap dynamics. The change in flap dynamics due to I54V and G48V mutations has recently been confirmed for unliganded Flap+(I54V) by MD and NMR studies.⁹

When Flap+(I54V) is compared with Flap+(I54A), the extreme entropy–enthalpy compensation and considerable amount of the cooperativity is lost upon shortening of the side chain not in contact with the inhibitors. There are significant differences

in the enthalpy and entropy of binding, despite only small differences between the free energy of binding to any inhibitor for Flap+(I54V) and Flap+(I54A), only 0.2 and 0.6 kcal mol $^{-1}$ for APV and DRV, respectively. However, the enthalpy of binding to APV was 10 kcal mol $^{-1}$ less favorable for Flap+(I54V) than for Flap+(I54A), and the entropy of binding was 10 kcal mol $^{-1}$ more favorable (Figure 2). Similarly, in the case of DRV, the enthalpy of binding to Flap+(I54V) was 9 kcal mol $^{-1}$ less favorable, and the entropy of binding was 9 kcal mol $^{-1}$ more favorable than binding to Flap+(I54A). These changes affected not only the magnitude of the enthalpy and entropy of binding but also the ratio of their contribution to the free energy of binding. As mentioned previously, the binding of Flap+(I54V) to both APV and DRV was enthalpically unfavorable, and these reactions were entirely driven by the entropy of binding (Table 1). In contrast, the enthalpic and entropic components of binding contributed approximately equally to the free energy of binding

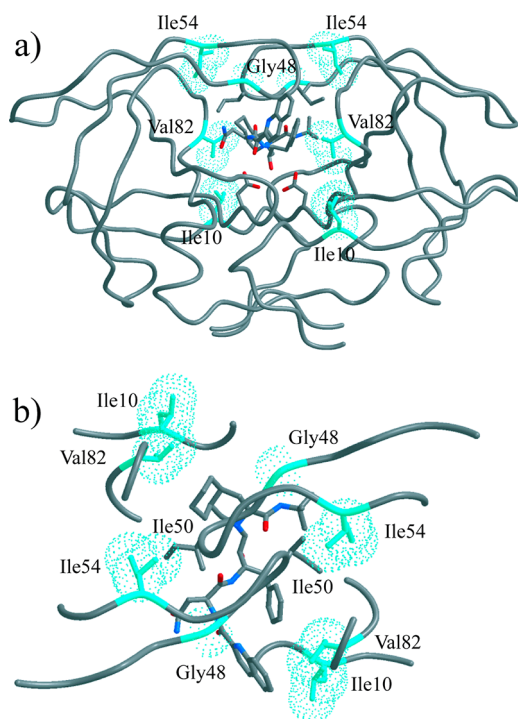


Figure 3. Positions of mutated residues displayed on inhibitor-bound HIV-1 protease structure (WT protease bound to SQV, PDB code 1HXB). Ile10, Gly48, Ile54, and Val82 side chains are in cyan van der Waal spheres. The SQV molecule is in stick representation and colored according to atom type, with gray for carbon, blue for nitrogen, and red for oxygen. (a) Front view, showing that neither residue 10 nor residue 54 is in contact with the inhibitor. (b) Top view, showing that residue 54 is in close contact with Ile50 of the other monomer.

between Flap^{+(I54A)} and APV or DRV. These drastic differences between Flap^{+(I54V)} and Flap^{+(I54A)} clearly indicate that relatively small changes at Ile54 can alter binding of inhibitors.

Our results indicate that residues in the flap region have inhibitor-dependent complex cooperative effects on the thermodynamics of inhibitor binding. The entropy-driven binding and the large cooperative entropy enhancement upon simultaneous G48V/I54V mutations could, in theory, be due to either enhancement of solvation entropy or conformational dynamics. Even though the overall structure and shape of the binding site are not expected to be considerably altered due to the mutations introduced, changes in solvation cannot be completely ruled out. However, protein flexibility, which dictates conformational entropy, has been reported to be the significant component to entropy in certain protein systems and modulate ligand binding.^{32,33} In addition to the free form,⁹ dynamics of inhibitor-bound wild-type and mutant protease variants will help elucidate the effect of flap flexibility on conformational entropy and hence the observed extensive cooperativity in entropy–enthalpy compensation.

METHODS

Protease-Gene Construction. The synthetic gene for the wild-type (WT) HIV-1 protease sequence was made using codons optimized for protein expression in *Escherichia coli*; the gene also included a substitution of Q7K to prevent autoproteolysis.³⁴ Mutagenesis was performed using a QuikChange Site-Directed Mutagenesis Kit (Stratagene) and confirmed by DNA sequencing.

All protease genes, including wild-type (WT), contained the polymorphism L63P.

Protease Expression and Purification. Overexpression and purification of HIV-1 protease was previously described.^{4,13} Briefly, the HIV-1 protease gene was cloned into the plasmid pXC-35 (ATCC),³⁵ which was then transformed into *Escherichia coli* TAP56. Transformed cells were grown in a 12 L fermenter and, following protein expression, lysed to release inclusion bodies containing the protease.³⁶ The inclusion bodies were isolated by centrifugation, and the pellet was dissolved in 50% (v/v) acetic acid to extract the protease. High molecular weight proteins were separated from the desired protease by size-exclusion chromatography on a 2.1 L Sephadex G-75 superfine column (Sigma) equilibrated with 50% (v/v) acetic acid. Refolding was accomplished by rapidly diluting the protease solution into a 10-fold excess of refolding buffer. Excess acetic acid was removed through dialysis.

Isothermal Titration Calorimetry. The binding affinity and enthalpy were measured using direct titrations for SQV and competitive displacement isothermal titration calorimetry^{5,6,37,38} for APV and DRV on a VP isothermal titration calorimeter (MicroCal) at 20 °C, as previously described.^{4,39} The buffer used consisted of 10 mM sodium acetate (pH 5.0), 2.0% dimethyl sulfoxide, and 2 mM tris(2-carboxyethyl) phosphine. Depending on the variant and inhibitor being used, protease concentrations were between 12 and 45 μM, as determined by absorbance at 280 nm. The protease solution was first saturated with 0.3 to 0.4 mM pepstatin. The pepstatin was then displaced from the protease by titrating 0.066 to 0.2 mM inhibitor. Heats of dilution were subtracted from the corresponding heats of reaction to obtain the heat caused solely by binding of the ligand to the enzyme. Data was analyzed using Origin7 software (OriginLab). Final results represent the average of at least three independent measurements.

ASSOCIATED CONTENT

Supporting Information

Thermodynamic parameters for the binding of each protease variant to APV, DRV, and SQV. Patient sequence analysis for I54A mutation. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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The authors declare no competing financial interest.

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