

### **RESEARCH ARTICLE**

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# Dynamical Basis for Drug Resistance of HIV-1 Protease

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#### **Abstract**

**Background:** Protease inhibitors designed to bind to protease have become major anti-AIDS drugs. Unfortunately, the emergence of viral mutations severely limits the long-term efficiency of the inhibitors. The resistance mechanism of these diversely located mutations remains unclear.

**Results:** Here I use an elastic network model to probe the connection between the global dynamics of HIV-1 protease and the structural distribution of drug-resistance mutations. The models for study are the crystal structures of unbounded and bound (with the substrate and nine FDA approved inhibitors) forms of HIV-1 protease. Coarse-grained modeling uncovers two groups that couple either with the active site or the flap. These two groups constitute a majority of the drug-resistance residues. In addition, the significance of residues is found to be correlated with their dynamical changes in binding and the results agree well with the complete mutagenesis experiment of HIV-1 protease.

**Conclusions:** The dynamic study of HIV-1 protease elucidates the functional importance of common drug-resistance mutations and suggests a unifying mechanism for drug-resistance residues based on their dynamical properties. The results support the robustness of the elastic network model as a potential predictive tool for drug resistance.

#### **Background**

HIV-1 protease (human immunodeficiency virus type 1 protease) is an enzyme that plays a critical role in the virus replication cycle. It cleaves the *gag* and *pol* viral polyproteins at the active site to process viral maturation [1-3], and without HIV-1 protease the virus was found to be noninfectious [4]. Thus HIV-1 protease is widely considered the major target for AIDS treatment [5,6]. One of the most severe obstacles to protease-inhibiting drugs is the rapid emergence of protease variants. Variants are able to evolve resistance by developing a chain of mutations, and as a result limit the long-term efficiency of these drugs [7,8].

HIV-1 protease is a dimer of C2 symmetry with each monomer consisting of 99 amino acid residues. Each monomer has one  $\alpha$  helix and two antiparallel  $\beta$  sheets in the secondary structure. The enzyme active site is a catalytic triad composed of Asp25-Thr26-Gly27 from each monomer. It is gated by two extended  $\beta$  hairpin

loops (residues 46–56) known as flaps [9]. At the molecular level, resistance to protease inhibition predominantly takes the form of mutations within the protein that preferentially lower the affinity of protease inhibitors with respect to protease substrates, while still maintaining a viable catalytic activity [10]. Mutations associated with drug resistance occur within the active site as well as non-active distal sites [11].

During the past two decades, researchers and clinicians from different disciplines have made enormous efforts to investigate resistance against HIV-1 protease targeted drugs. To elucidate the molecular mechanisms of drug resistance, biochemists and molecular biologists have characterized the structure, energetics and catalytic efficiency of a large number of HIV-1 protease mutants to unravel the resistance mechanism in combination with extensive computational studies [12-15]. Moreover, drug resistance data collected from AIDS patients treated with HIV-1 protease inhibitor drugs [16-19] provide opportunities for researchers to identify resistance-related mutation patterns [20-22]. Recently there have

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been efforts to link protein physical and functional stability with its evolutionary dynamics [23,24].

At the heart of understanding the molecular basis of drug-resistant behaviors of HIV-1 protease is the structural distribution of resistance mutations. Presumably these mutations are not randomly located throughout the protein structure. Although different HIV-1 protease inhibitors elicit different combinations of mutation types to generate distinctive resistance levels, there are 21 most common mutations associated with resistance against all inhibitors [19]. Prediction of resistance mutations of proteins is based on either sequence or structure information [25]. Sequence-based methods predict resistance mutations by analyzing large datasets of sequences with known resistance properties. Thus the availability of those datasets is a prerequisite for such methods [22,26-28]. On the other hand, predicting mutations using protein structure has largely relied on the characterization of binding thermodynamics [29-32], as the mutations with resistance against inhibitors lower the binding affinity of inhibitors far more than that of natural substrates. The accuracy of the prediction is directly related to the accuracy of the potential function used in the calculations and the adequacy of the sampling of the protein conformational space. It is also sensitive to the error/noise in the free energy calculations [32].

Conformational dynamics play an essential role in regulating protein function [33,34]. In the past few years a deepening understanding of the relationship of protein dynamics and function has emerged [35]. Relevant to the study here is the utilization of protein dynamics to identify the sequence regions of functional importance even though their locations may be remote from the active site. Computationally there have been rapid methodological developments in relating protein dynamics to function by probing the long range communications between residues: perturbation method [36,37], clustering analysis of correlation matrix [38], network analysis [39], and energy diffusivity estimation by propagation through vibrational modes [40]. The success of these methods in reproducing experimental results as well as findings from sequence-based methods has established the validity of dynamics-based approaches [38,41].

The dynamics of HIV-1 protease, especially binding dynamics of its ligands are fundamental to the protease inhibitor design and have been a subject of intense computational study [42-49]. Because of limitations of time scale in all-atom simulations, various coarse-grained models have been used to investigate HIV-1 protease binding dynamics and kinetics, shedding light on important dynamics issues [45-49]. The main features of substrate interactions and dynamics at the active site were analyzed within the framework of the coarse-grained model [45,49]. Gaussian models were shown to describe

accurately the correlated motion of HIV-1 protease residues in thermodynamic equilibrium through a series of successful comparisons with an extensive MD simulation [49,50]. There is increasing evidence relating protease's drug-resistance mutations to its dynamics. The impact of some distal mutations on catalytic function of HIV-1 protease was linked to protein flexibility [51,52]. Multi-drug resistance residues of HIV-1 protease were found to overlap the global hinge region identified from coarse-grained normal-mode analysis of the protease [53]. Nevertheless, despite extensive research efforts, a general explanation for drug-resistance mutations of HIV-1 protease is still lacking [54].

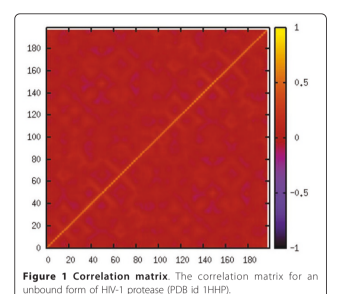
In this study, a coarse-grained elastic network model is used to investigate the dynamics of HIV-1 protease, to probe the connection between its global dynamics and the distribution of drug-resistance mutations, and to examine the potential of the dynamics-based approach as a predictive tool for drug resistance prediction, with an attempt to provide a unifying mechanistic explanation for all residues of resistance based on their dynamical properties. The crystal structures of an unbound form and bound forms with a substrate and nine FDA approved inhibitors of HIV-1 protease are used as model systems. Correlation analysis of the protease at equilibrium focuses on two functional sites of HIV-1 protease: the active site (the Asp25-Thr26-Gly27 triad [45]) and the flap (residues 45-55 [47]). The protease dynamic changes upon ligand binding are examined as well. The implications for resistance mechanisms and protein evolution are discussed.

#### **Results**

Here HIV-1 protease is represented by a coarse-grained network model, and its dynamics is examined in several X-ray crystallographic structures. The linkage between global dynamics and the distribution of drug-resistance mutations is examined first in individual unbound and bound forms, then in the dynamical differences between the unbound and bound forms. The former is a measure of the residual fluctuations in different structures, and the latter is an estimate of dynamical change caused by ligand binding.

## Dynamically coupled regions identified by equilibrium correlations Unbound form

The correlation matrix, consisting of correlations of all residue pairs, captures the essence of protein dynamics. A  $198 \times 198$  correlation matrix (Figure 1) is generated from the elastic network modeling (see Methods section) based on the unbound HIV-1 protease (PDB id 1HHP). The most conspicuous features in the figure are the beta-sheets, represented by the line across the diagonal (residues 19-24, 43-66, 69-78, 118-123, 142-165 and



168-177). Extraction of further information from the matrix requires the application of data analysis tools such as clustering algorithms. The Markov cluster (MCL) algorithm (see Methods section) is the chosen method, and the results from the MCL program for the unbound form consist of five clusters (Table 1). Cluster 1 is the largest cluster with 48 residues. It covers the core domain, which excludes both termini and the flap, and forms the scaffold surrounding the active site of the enzyme. Clusters 2 and 3 are exclusively composed of the residues located at the beginning of the N-terminus and the end of the C-terminus, respectively. The remaining two clusters concentrate on two functional sites (Figure 2). Cluster 4 contains the active site of HIV-1 protease (Asp25-Thr26-Gly27 catalytic triad), one residue at the dimerization region (Leu10), and a large segment of the C-terminal (Ile84-Cys95). Cluster 5 is mainly made up of the residues at the flap region (Pro44-Arg57) with two additional residues from the Cterminal domain (Leu76 and Gly78). These two functional sites are involved in distinct aspects of the protease function: the active site is where the enzymatic catalysis takes place, and the flap controls access to the active site. The clustering results indicate that these two

functional sites belong to separate interaction networks of the protease.

#### **Bound forms**

The ten structures of HIV-1 protease in complex with the natural substrate and nine FDA approved inhibitors are chosen to represent HIV-1 protease in bound forms. Despite their varying degrees of similarity with the unbound form, these ten structures generate very similar clustering results (Table 2). The general clustering pattern of HIV-1 protease unbound form is preserved in all the bound forms: the scaffold, the N- and C- termini, the active site and the flap. Nevertheless, inside the clusters there are reorganizations and splits of the clusters due to ligand binding. Upon binding, the tips of the flaps (residues 48-55) close and cover the active site, and the physical proximity facilitates stronger interactions between the flap region and part of the C-terminal. As a result, the cluster containing the flap grows in the bound form. The cluster containing the flap is enlarged by including residues 79-83, while part of the C-terminal (residues 87-95) is disengaged from the cluster with the active site and forms a new cluster of its own. The diversity of the ligand types of the ten structures does not induce a dramatic impact on the clusters. The number of residues in the cluster of the active site and of the flap ranges from 12 to 18 and from 19 to 24 respectively, but the number of separate sequence segments in these two clusters remains constant.

In summary, there exist two independent clusters containing two important functional sites of HIV-1 protease. These two networks are relatively robust to perturbations caused by the different types of ligand.

### Dynamical behavior differences between the unbound and bound forms

Without directly engaging the active site, another way to influence a protein's function is to perturb the motions essential to its function. For enzymes, these essential motions are the conformational changes accompanied by the association and dissociation of ligands [55]. In HIV-1 protease the large scale open/close conformational change of the flap along the reaction pathway is the major structural reorganization induced by ligand

Table 1 Clustering result of the unbound form

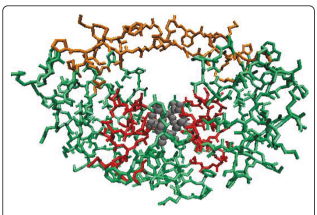
Scaffold 11 12 13 14 15 16 17 18 19 20 21 22 32 33 34 35 36 37 38 39 40 41 42 43 58 59 60 61 62 63 64 65 66 67 68 69 70

71 72 73 74 75 77 79 80 81 82 83

N-terminal 1 2 3 4 5 6 7 8 9 C-terminal 96 97 98 99

Coupled with the active site 10 23 24 25 26 27 28 29 30 31 84 85 86 87 88 89 90 91 92 93 94 95

Coupled with the flap region 44 45 46 47 48 49 50 51 52 53 54 55 56 57 76 78



**Figure 2 Two clusters containing functional sites**. Residues in Clusters 4 (red) and 5 (yellow) are displayed in an unbound form of HIV-1 protease (PDB id 1HHP). The active site is highlighted by a ball representation.

binding [43-47]. Extensive investigations of 73 X-ray mutant and complex structures of HIV-1 protease revealed that a common and predominant dynamic behavior was found among the protease in complex with different ligands [56]. The focus of study here is the dynamical changes from the unbound to bound forms of HIV-1 protease at the residue level. The dynamical property of each residue is characterized by the sum of its couplings with other residues (see Methods). The binding-induced changes in the dynamical behaviors of individual residues were very similar among the ten bound structures (Figure 3). Overall, the perturbations to the residues are not isotropic, and the regions exhibiting the largest deviations are signaled by peaks. The ten bound proteases all share very similar locations and magnitudes of these peaks (residues 32-42, 44-57 and 77-82). The most pronounced difference between the unbound and bound protease is the loss of flexibility in the flap tip upon binding, as indicated by the highest peak (around residues 44-57) in Figure 3.

#### Discussion

Using a coarse-grained network model, functionally important residues of HIV-1 protease were identified based on correlation analysis of either equilibrium fluctuations or dynamical changes. Experimentally the residues of functional importance can be directly probed by mutagenesis experiments. The complete mutagenesis experiments carried out on HIV-1 protease revealed three mutationally sensitive sequence domains [57]: the active site region (Ala22-Leu33), the flap region (Ile47-Gly52) and the part of C-terminal (Thr74-Arg87). The active site and flap region have obvious functional significance. The influence the C-terminal residues exert on the catalytic cycle is most likely through long-range

couplings with the functional sites. Correlation analysis suggests that some residues of the C-terminal (Ile84-Cys95) are functionally important due to their strong interactions with the active site, while the others (Leu76 and Gly78) may influence the dynamics of the protein through their couplings with the flap. The dynamical changes analysis also indicates that residues 77-82 are involved in the binding process. Structurally, residues 79-84 form a wall of the active site, and their motions were shown by previous simulations to correlate well with the open/closed dynamics of the flap [52].

Currently there are nine FDA approved protease inhibitors, and the most up-to-date clinical data indicate 21 most common drug-resistant mutation positions ("most common" defined as mutation position shared by at least two inhibitors among the nine currently approved protease inhibitors):10, 20, 24, 32, 33, 36, 46, 47, 48, 50, 53, 54, 62, 71, 73, 76, 77, 82, 84, 88 and 90 [19]. These mutations are located at the dimer interface (residues 88 and 90), the core domain (residues 10, 20, 24, 32, 33, 36, 62, 71, 73, 76, 77, 82, and 84), and the flap domain (residues 46, 47, 48, 50, 53, and 54), respectively. The clustering analysis of HIV-1 protease in various forms identified 10, 24, 32, 46, 47, 48, 50, 53, 54, 76, 77, 82, 84, 88 and 90 (15 out of the total 21 resistance sites) as those coupled either with the active site or the flap. The dynamical change analysis identifies two additional residues 33 and 36 as residues of importance. The undetected residues (20, 62, 71 and 73) are located in the hydrophobic core of the protease, and their mutations likely affect the protease activity through influencing the protease structure and stability [30]. Another noteworthy fact is that there are far more residues discovered by clustering analysis than by clinical studies (15 out of the total 34 residues in the two clusters containing functional sites are the residues of resistance). Drug resistance residues not only influence protease binding, but also generate differential binding affinity between the substrate and inhibitors. The cluster analysis can only detect residues that may influence the binding. Therefore the pool of residues identified by the clustering analysis is larger than those found with drug resistance affinity in clinical studies. The similar clustering results from different protease inhibitor complexes further suggest that global dynamics are preserved among different complexes of HIV-1 protease inhibitors. The convergence of results regardless of the conformation of the protein was also found by the Gaussian network modeling of HIV-1 protease [50]. Computational studies using atomistic MD simulations reached the same conclusion that correlation matrix based analysis does not differentiate the essential modes of motion of the protein native forms from those of the mutants [58]. Nevertheless, the dynamic approach proposed here

Table 2 Clustering results of the ten bound forms

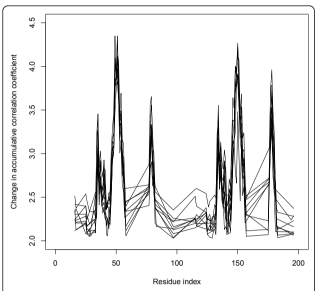
PDB id	Mutation	RMSD (Å)	Coupled with the active site	Coupled with the flap region
2FNS 7 (substrate)	Q7K, D25N, L63P, l64V	0.52	<u>10</u> 23 <u>24</u> 25 26 27 28 29 30 31 <u>84</u> 85 86 87	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 78 79 80 81 <u>82</u> 83
1HPV 6 (Amprenavir)	None	0.54	<u>10</u> 23 <u>24</u> 25 26 27 28 29 30 31 <u>32</u> 85 86	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 78 79 80 81 <u>82</u> 83 <u>84</u>
1HXB 6 (Saquinavir)	None	0.31	<u>1023 24 25 26 27 28 29</u> 30 31 <u>328</u> 5 86	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 78 79 80 81 <u>82</u> 83 <u>84</u>
1MUI 5 (Lopinavir)	N37S	0.53	<u>1023 24 25 26 27 28 29</u> 30 31 <u>3284</u> 85 86	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 78 79 80 81 <u>82</u> 83
2O4K 8 (Atazanavir)	Q7K	0.35	23 <u>24</u> 25 26 27 28 29 30 31 <u>32</u> 83 <u>84</u> 85 86 87 <u>88</u> 89 <u>90</u>	43 44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 57 58 <u>7677</u> 78 <i>79</i> 80 81 <u>82</u>
3JVY 6 (Darunavir)	Q7K, L33I*, L63I, C67A, G86A, C95A	0.38	23 <u>24</u> 25 26 27 28 29 30 31 <u>32</u> 85 86	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 79 80 81 <u>82</u> 83 <u>84</u>
2O4P 8 (Tipranavir)	Q7K	0.39	23 <u>24</u> 25 26 27 28 29 30 31 <u>32</u> 83 <u>84</u> 85 86 87 <u>88</u> 89 <u>90</u>	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 79 80 81 <u>82</u> 83 <u>84</u>
1HXW 6 (Ritonavir)	None	0.40	<u>10</u> 23 <u>24</u> 25 26 27 28 29 30 31 85 86 87	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 79 80 81 <u>82</u> 83 <u>84</u>
2PYN 6 (Nelfinavir)	D30N, A71V*	0.26	<u>1023 24 25 26 27 28 29</u> 30 31 <u>3284</u> 85 86	44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 78 79 80 81 <u>82</u> 83
2B7Z 7 (Indinavir)	K20R*, V32l*, L33F*, M36l*, M46l*, L63P, A71V*,V82A*, I84V*, L90M*	0.32	<u>10</u> 22 23 <u>24</u> 25 26 27 28 29 <u>84</u> 85 86	34 43 44 45 <u>464748</u> 49 <u>50</u> 51 52 <u>5354</u> 55 56 57 <u>7677</u> 78 79 80 81 <u>82</u> 83

The type of the ligand is listed in brackets. The global average RMSD of backbone atoms between the unbound (PDB id 1HHP) and bound forms is given. The mutations of each protease are listed. Only the clusters containing the active site and the flap are shown here.

<sup>(1) \*</sup>drug-resistance mutations

<sup>(2)</sup> Residues belonging to the 21 common resistant mutations are underlined.

<sup>(3)</sup> Residues shared by all the bound forms are in italics.



**Figure 3 Fluctuation differences between the unbound and bound forms.** Fluctuation difference between the unbound (PDB id 1HHP) and ten bound forms of HIV-1 protease plotted as a function of residues. Only residues with larger than average difference are plotted.

locates a majority of drug-resistance mutations, and provides insight on the drug-resistance mechanism of HIV-1 protease. Drug resistance mutations of HIV-1 protease can be classified as active site or non-active site mutations, depending on their location within the protein. Active site mutations are located in the vicinity of the active site and directly affect the protease-inhibitor interactions. Thus their action on inhibitor binding affinity can be readily understood in structural terms. On the other hand, non-active site mutations influence binding from various distal locations and their mechanism of action is not immediately apparent. Although some residue-specific explanations and suggestions have been proposed, the overall mechanism by which these diversely located non-active site residues influence inhibitor-binding remains unclear [19]. Dynamic studies presented here suggest a simple yet general explanation for the distribution of the drug resistance residues in HIV-1 protease. Except for these residues of resistance influencing protease binding affinity by acting on the structural stability of HIV-1 protease, the drug-resistance residues belong to clusters that are either the "coupled with the active site" or "coupled with the flap". It is noteworthy that residues coupled with the functional sites (Clusters 4 and 5) do not all locate in the physical vicinity of each other. Long distance communications play an important role in mediating the interactions between the functional sites and distal residues. The residues clustered with the functional sites can exert an impact on protein function, even though they may not locate near the active site. Leu10 and Leu90, among the residues clustered with the active site (Cluster 3), are well-known residues whose mutations lead to drug resistant variants [17]. The finding that Leu76 and Gly78 are coupled with the flap region is also corroborated by a detailed protein flexibility study of HIV-1 protease which concluded that mutations at residues Thr74-Val77, although far from the active site, reduce protease activity because of their correlated motions with the flap region [55]. Traditionally, computer methods of predicting resistance mutations based on protein structure have been largely focused on energetic analysis [25], in which atomistic molecular mechanics and/or molecular dynamics are used to investigate ligand-protein binding affinity. The mechanism of non-active site mutations has largely remained a challenge for energetic analysis because of the minimal structural and energetic perturbations caused by those mutations [51]. These difficulties, however, open the door for dynamics-based study, especially via coarse-grained methods such as elastic network modeling, which provide an efficient way of sampling the global dynamics of proteins. The interaction network identified by the correlation analysis contains both active site and non-active site residues. These residues influence the inhibitor binding by coupling with the active site and the flap, regardless of their physical proximity. Thus the dynamic approach in this study is able to detect both active and non-active drugresistance sites based on coarse-grained protein models.

It is believed that only a few amino acid sites are responsible for adaptive evolution in almost all proteins [59], although the nature of these positively selected residues is yet to be elucidated. The findings from this study indicate that the interaction networks of globally distributed residues involving the functional sites play a dominant role in the evolutionary pathways of HIV-1 protease, and they become the major sites that develop resistance under selective antiviral pressure. Pathogenic proteins such as HIV-1 protease escape the challenges to their survival imposed by drug inhibition through mutations at these amino acid residues. Structure-based modeling of proteins confirms the decisive role of physical interactions in the evolution of virus proteins and raises the possibility of constructing a protein fitness landscape by means of physical modeling of proteins.

#### **Conclusions**

This study examines the functional significance of common drug-resistance mutations of HIV-1 protease by characterizing its global dynamics using coarse-grained modeling. The calculations show that most residues of drug-resistance are coupled either with the active site or with the flap. These couplings are rather robust to the perturbation of ligand binding. These findings result in

a unifying mechanism for all drug-resistance residues based on their dynamical properties. They also indicate that global dynamics of HIV-1 protease are intrinsically connected to the structural distribution of drug-resistance mutations, thus dynamic study provides a simple yet general and useful tool to examine the tendency of drug resistance of residues in addition to traditional energetic analysis.

#### **Methods**

#### Elastic network modelling

The elastic network model was applied according to the standard protocol [60]. The details of the correlation matrix can be found elsewhere [41]. In this study the cutoff distance  $R_c$  is set to be 10Å, but the choice of  $R_c$  was shown not to noticeably affect the results based on the correlation matrix generated from the model [61]. The structures of the unbound and ligand-bound forms of HIV-1 protease with various ligands were used. For the ligand-bound protein, only the  $C_\alpha$  atoms of the protein are represented by the network model and the ligand is not incorporated in the model. The online server http://ignmtest.ccbb.pitt.edu/cgi-bin/anm/anm1.cgi [60] was used to generate the correlation matrix, and the element in the correlation matrix is defined as

$$C_{ij} = \frac{\langle \Delta R_i \Delta R_j \rangle}{\left[\langle \Delta R_i^2 \rangle \langle \Delta R_i^2 \rangle\right]^{\frac{1}{2}}}$$
(1)

where  $\Delta R_i$  and  $\Delta R_j$  are the fluctuations of nodes i and j, respectively.

#### Clustering analysis

The correlation matrix is submitted for clustering analysis by the Markov cluster (MCL) algorithm [62]. The MCL algorithm is one of the most successful clustering procedures in identifying protein-protein interactions from genomic data [63] and has been shown to be robust and outperform other clustering algorithms [64]. Relevant to the study here is the application of MCL to clustering protein residues based on the interaction correlation matrix [38]. MCL finds cluster structure in graphs by performing a random walk through the graphs. The process computes the probabilities of random walks through the graph, and uses expansion and inflation to change the probabilities associated with the random walks departing from one particular node. It results in the separation of the graph into different segments. Cluster granularity is controlled by the inflation parameter which is the only variable in the MCL program used in this study. In order to reduce the noise, the correlation matrix has to be adjusted before being submitted to the MCL program. First the absolute values of the correlation coefficients are taken. Then a cutoff value is applied to produce a condensed version. Correlations less than the cutoff value are set to zero and the cutoff value is subtracted from the remaining correlations. The cutoff and the inflation parameter (0.08 and 1.5, 0.075 and 1.4 for the unbound and bound protease, respectively) were chosen to produce a total number of five clusters for the unbound protease, and six to seven clusters for the bound protease. The number of clusters is chosen to be five or six because the resulting clusters make the most physical sense. All the bound forms are subject to the same parameters.

## Fluctuation difference between unbound and bound forms

The structural differences between the unbound (ligand-free) and bound (ligand-bound) proteins are usually significant. The structural changes are not to be identified with the dynamical behavior changes. The change in dynamical behavior caused by binding for residue i,  $\Delta C_i$ , is calculated as the sum of the absolute values of the difference between the correlation coefficients of residue i in the unbound and bound forms,

$$\Delta C_i = \sum_j |C_{ij} - C'_{ij}| \tag{2}$$

In Equation 2, the sum is over all the residues in the protein.  $C_{ij}$  and  $C'_{ij}$  denote the correlation coefficient of residues i and j in the unbound and bound forms, respectively.

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#### Authors' contributions

YM is responsible for all the work related to the manuscript.

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#### References

- 1. Katz RA: The retroviral enzymes. Annu Rev Biochem 1994, 63:133-173.
- Wlodawer A: Rational approach to AIDS drug design through structural biology. Ann Red Med 2002, 53:595-614.
- Kurup A, Mekapati SB, Garg R, Hansch C: HIV-1 protease inhibitors: a comparative QSAR analysis. Curr Med Chem 2003, 10:1679-1688.
- Kohl NE, Emini EA, Schleif WA, Davis LJ, Heimbach JC, Dixon RA, Scolnick EM, Sigal IS: Active human immunodeficiency virus protease is required for viral infectivity. Proc Natl Acad Sci USA 1988, 85:4686-4690.
- Wlodawer A, Vondrasek J: Inhibitors of HIV-1 protease: A major success of structureassisted drug design. Annu Rev Biophys Biomol Struct 1998, 27:249-284.

- Blair WS: HIV-1 entry- an expanding portal for drug discovery. Drug Discovery Today 2000, 5:183-194.
- Condra JH, Schleif WA, Blahy OM, Gabryelski LJ, Graham DJ, Quintero JC, Rhodes A, Robbins HL, Roth E, Shivaprakash M, Titus D, Yang T, Tepplert H, Squires KE, Deutsch PJ, EMini EA: In-vivo emergence of hiv-1 variants resistant to multiple protease inhibitors. Nature 1995, 374:569-571.
- Ho DD, Toyoshima T, Mo H, Kempf DJ, Norbeck D, Chen CM, Wideburg NE, Burt SK, Erickson JW, Singh MK: Characterization of human immunodeficiency virus type 1 variants with increased resistant to a C2-symmetric protease inhibitor. J Virol 1994. 68:2016-2020.
- Mager PP: The active site of HIV-1 protease. Med Res Rev 2001, 21:348-353.
- 10. Freire E: Overcoming HIV-1 resistance to protease inhibitors. *Drug Discov Today* 2006, **3**:281-286.
- Muzammil S, Ross P, Freire E: A major role for a set of non-active site mutations in the development of HIV-1 protease drug resistance. *Biochemistry* 2003, 42:631-638.
- Chen XF, Weber IT, Harrison RW: Molecular dynamics simulations of 14 HIV protease mutants in complexes with indinavir. J Mol Model 2004, 10:373-381.
- Hou TJ, McLaughlin WA, Wang W: Evaluating the potency of HIV-1 protease drugs to combat resistance. Proteins 2008, 71:1163-1174.
- Muzammil S, Armstrong AA, Kang LW, Jakalian A, Bonneau PR, Schmelmer V, Amzel LM, Freire E: Unique thermodynamic response of tipranavir to human immunodeficiency virus type 1 protease drug resistance mutations. J Virol 2007, 81:5144-5154.
- Prabu-Jeyabalan M, Nalivaika E, Schiffer CA: Substrate shape determines specificity of recognition for HIV-1 protease: analysis of crystal structures of six substrate complexes. Structure 2002, 10:369-381.
- Hertogs K, Bloor S, Kemp SD, Van den Eynde C, Alcorn TM, Pauwels R, Houtte MV, Staszewski S, Miller V, Larder BA: Phenotypic and genotypic analysis of clinical HIV-1 isolates reveals extensive protease inhibitor cross-resistance: a survey of over 6000 samples. AIDS 2000, 14:1203-1210.
- Shafer RW: Genotypic testing for human immunodeficiency virus type 1 drug resistance. Clin Microbiol Rev 2002, 15:247-277.
- Chen L, Perlina A, Lee CJ: Positive selection detection in 40,000 human immunodeficiency virus (HIV) type 1 sequences automatically identifies drug resistance and positive fitness mutations in HIV protease and reverse transcriptase. J Virol 2004, 78:3722-3732.
- Johnson VA, Brun-Vezinet F, Clotet B, Gunthard HF, Kuritzkes DR, Pillay D, Schapiro JM, Richman DD: Update of the drug resistance mutations in HIV-1: December 2009. Top HIV Med 2009, 17:138-145.
- Vercauteren J, Vandamme AM: Algorithms for the interpretation of HIV-1 genotypic drug resistance information. Antivir Res 2006, 71:335-342.
- Beerenwinkel N, Schmidt B, Walter H, Kaiser R, Lengauer T, Hoffmann D, Korn K, Selbig J: Diversity and complexity of HIV-1 drug resistance: a bioinformatics approach to predicting phenotype from genotype. Proc Natl Acad Sci USA 2002, 99:8271-8276.
- Rhee SY, Taylor J, Wadhera G, Ben-Hur A, Brutlag DL, Shafer RW: Genotypic predictors of human immunodeficiency virus type 1 drug resistance. Proc Natl Acad Sci USA 2006, 103:17355-17360.
- Hamacher K: Relating sequence evolution of HIV1-protease to its underlying molecular mechanics. Gene 2008, 422:30-36.
- Zhang J, Hou T, Wang W, Liu JS: Detecting and understanding combinatorial mutation patterns responsible for HIV drug resistance. Proc Natl Acad Sci USA 2010, 107:1321-1326.
- Cao ZW, Han LY, Zheng CJ, Ji ZL, Chen X, Lin HH, Chen YZ: Computer prediction of drug resistance mutations in proteins. *Drug Discov Today* 2005, 10:521-529.
- Saigo H, Uno T, Tsuda K: Mining complex genotypic features for predicting HIV-1 drug resistance. Bioinformatics 2007, 23:2455-2462
- Wang K, Jenwitheesuk E, Samudrala R, Mittler JE: Simple linear model provides highly accurate genotypic predictions of HIV-1 drug resistance. Antivir Ther 2004, 9:343-352.
- Zazzi M, Romano L, Venturi G, Shafer RW, Reid C, Dal Bello F, Parolin C, Palù G, Valensin PE: Comparative evaluation of three computerized algorithms for prediction of antiretroviral susceptibility from HIV type 1 genotype. J Antimicrob Chemother 2004, 53:356-360.
- Wang W, Kollman PA: Computational study of protein specificity: the molecular basis of HIV-1 protease drug resistance. Proc Natl Acad Sci USA 2001, 98:14937-14942.

- 30. Chen YZ, Gu XL, Cao ZW: Can an optimization/scoring procedure in ligand protein docking be employed to probe drug resistant mutations in proteins? *J Mol Graph Model* 2001, **19**:560-570.
- 31. Shenderovich MD, Kagan RM, Heseltine PNR, Ramnarayan K: **Structure-based phenotyping predicts HIV-1 protease inhibitor resistance.** *Protein Sci* 2003, **12**:1706-1718.
- 32. Hou T, Zhang W, Wang J, Wang W: Predicting drug resistance of the HIV-1 protease using molecular interaction energy components. *Proteins* 2009, **74**:837-846.
- Eisenmesser EZ, Millet O, Labeikovsky W, Korzhnev DM, Wolf-Watz M, Bosco DA, Skalicky JJ, Kay LE, Kern D: Intrinsic dynamics of an enzyme underlies catalysis. Nature 2005, 438:117-121.
- 34. Kern D, Zuiderweg ERP: The role of dynamics in allosteric regulation. *Nature* 2003, **13**:748-757.
- 35. Bahar I, Lezon TR, Yang LW, Eyal E: Global dynamics of proteins: bridging between structure and function. *Annu Rev Biophys* 2010, **39**:23-42.
- Zheng W, Tekpinar M: Large-scale evaluation of dynamically important residues in proteins predicted by the perturbation analysis of a coarsegrained elastic model. BMC Struct Biol 2009, 9:45.
- Sharp K, Skinner JJ: Pump-probe molecular dynamics as a tool for studying protein motion and long range coupling. Proteins 2006, 65:347-361.
- Kong Y, Karplus M: Signaling pathways of PDZ2 domain: A molecular dynamics interaction correlation analysis. Proteins 2009, 74:145-154.
- Cusack MP, Thibert B, Bredesen DE, del Rio G: Efficient identification of critical residues based only on protein structure by network analysis. PLoS ONE 2007, 2:e421.
- 40. Leitner DM: Frequency-resolved communication maps for proteins and other nanoscale materials. *J Chem Phys* 2009, **130**:195101-195109.
- 41. Mao Y: Dynamics studies of luciferase using elastic network model: how the sequence distribution of luciferase determines its color. *Protein Eng Des Sel* 2011, **24**:341-349.
- Song Y, Zhang Y, Shen T, Bajaj CL, McCammon JA, Baker NA: Finite element solution of the steady-state Smoluchowski equation for rate constant calculations. *Biophys J* 2004, 86:2017-2029.
- Hornak V, Okur A, Rizzo RC, Simmerling C: HIV-1 protease flaps spontaneously close to the correct structure in simulations following manual placement of an inhibitor into the open state. J Am Chem Soc 2006. 128:2812-2813.
- 44. Toth G, Borics A: Closing of the flaps of HIV-1 protease induced by substrate binding: A model of flap closing mechanism in retroviral aspartic proteases. *Biochemistry* 2006, 45:6606-6614.
- 45. Li D, Liu MS, Ji B, Hwang K: Coarse-grained molecular dynamics of ligands binding into protein: the case of HIV-1 protease inhibitors. J Chem Phys 2009, 130:215102.
- Tozzini V, McCammon JA: A coarse grained model for the dynamics of flap opening in HIV-1 protease. Chem Phys Lett 2005, 413:123-128.
- Tozzini V, Trylska J, Chang C, McCammon JA: Flap opening dynamics in HIV-1 protease explored with a coarse-grained model. J Struct Biol 2007, 157:606-615.
- Pandey RB, Farmer BL: Residue energy and mobility in sequence to global structure and dynamics of a HIV-1 protease (1DIFA) by a coarsegrained Monte Carlo simulation. J Chem Phys 2009, 130:044906.
- Micheletti C, Carloni P, Maritan A: Accurate and efficient description of protein vibrational dynamics: comparing molecular dynamics and Gaussian models. Proteins 2004, 55:635-645.
- Kurt N, Scott WR, Schiffer CA, Haliloglu T: Cooperative fluctuations of unliganded and substrate-bound HIV-1 protease: A structure-based analysis on a variety of conformations from crystallography and molecular dynamics simulations. *Proteins* 2003, 51:409-422.
- Piana S, Carloni P, Rothlisberger U: Drug resistance in HIV-1 protease: Flexibility assisted mechanism of compensatory mutations. Protein Sci 2002. 11:2393-2402.
- Perryman AL, Lin JH, McCammon JA: HIV-1 protease molecular dynamics of a wild-type and of the V82F/I84V mutant: possible contributions to drug resistance and a potential new target site for drugs. Protein Sci 2004. 13:1108-1123.
- 53. Liu Y, Eyal E, Bahar I: Analysis of correlated mutations in hiv-1 protease using spectral clustering. *Bioinformatics* 2008, **24**:1243-1250.
- Ali A, Bandaranayake RM, Cai Y, King NM, Kolli M, Mittal S, Murzycki JF, Nalam MNL, Nalivaika EA, Özen A, Prabu-Jeyabalan MM, Thayer K,

- Schiffer CA: Molecular basis for drug resistance in HIV-1 protease. Viruses 2010, 2:2509-2535.
- Szarecka A, Xu Y, Tang P: Dynamics of firefly luciferase inhibition by general anesthetics: Gaussian and anisotropic network analyses. *Biophys* J 2007, 93:1895-1905.
- Zoete V, Michielin O, Karplus M: Relation between sequence and structure of HIV-1 protease inhibitor complexes: a model system for the analysis of protein flexibility. J Mol Biol 2002, 315:21-52.
- Loeb DD, Swanstrom R, Everitt L, Manchester M, Stamper SE, Hutchison CA
  Complete mutagenesis of the HIV-1 protease. Nature 1989, 340:397-340.
- 58. Genoni A, Morra G, Merz KM Jr, Colombo G: Computational study of the resistance shown by the subtype B/HIV-1 protease to currently known inhibitors. *Biochemistry* 2010, **49**:4283-4295.
- Guindon S, Rodrigo AG, Dyer KA, Huelsenbeck JP: Modeling the sitespecific variation of selection patterns along lineages. Proc Natl Acad Sci USA 2004, 101:12957-12962.
- Chennubhotla C, Rader AJ, Yang LW, Bahar I: Elastic network models for understanding biomolecular machinery: from enzymes to supramolecular assemblies. *Phys Biol* 2005, 2:S173-S180.
- Atilgan AR, Durrel SR, Jernigan RL, Demirel MC, Keskin O, Bahar I: Anisotropy of fluctuation dynamics of proteins with an elastic network model. *Biophys J* 2001, 80:505-515.
- Enright AJ, Van Dongen S, Ouzounis CA: An efficient algorithm for largescale detection of protein families. Nucleic Acids Res 2002, 30:1575-1584.
- Vlasblom J, Wodak SJ: Markov clustering versus affinity propagation for the partitioning of protein interaction graphs. BMC Bioinformatics 2009, 10-99
- 64. Brohee S, van Helden J: Evaluation of clustering algorithms for proteinprotein interaction networks. *BMC Bioinformatics* 2006, 7:488.

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