Research article

Open Access Structural and functional implications of p53 missense cancer mutations Yuhong Tan and Ray Luo*

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Abstract

Most human cancers contain mutations in the transcription factor p53 and majority of these are missense and located in the DNA binding core domain. In this study, the stabilities of all core domain missense mutations are predicted and are used to infer their likely inactivation mechanisms. Overall, 47.0% non-PRO/GLY mutants are stable ($\Delta\Delta G < 1.0$ kT) and 36.3% mutants are unstable ($\Delta\Delta G$ > 3.0 kT), 12.2% mutants are with 1.0 kT < $\Delta\Delta G$ < 3.0 kT. Only 4.5% mutants are with no conclusive predictions. Certain types of either stable or unstable mutations are found not to depend on their local structures. Y, I, C, V, F and W (W, R and F) are the most common residues before (after) mutation in unstable mutants. Q, N, K, D, A, S and T (I, T, L and V) are the most common residues before (after) mutation in stable mutants. The stability correlations with sequence, structure, and molecular contacts are also analyzed. No direct correlation between secondary structure and stability is apparent, but a strong correlation between solvent exposure and stability is noticeable. Our correlation analysis shows that loss of protein-protein contacts may be an alternative cause for p53 inactivation. Correlation with clinical data shows that loss of stability and loss of DNA contacts are the two main inactivation mechanisms. Finally, correlation with functional data shows that most mutations which retain functions are stable, and most mutations that gain functions are unstable, indicating destabilized and deformed p53 proteins are more likely to find new binding partners.

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I. Introduction

p53 is a transcription factor involved in DNA repair, growth arrest and apoptosis [1-3]. It plays a critical role in cell responses to many cancer-causing events. Upon activation of certain oncogenes, p53 can stop cell cycle and induce apoptosis [1-3]. It has been reported that 50% of human cancers contain mutations in p53 [4], among which 95% are in the DNA-binding core (DBC) domain [4]. More interestingly 75% of DBC mutations occur as single missense mutations [4], i.e. most cancer mutants

are full-length proteins. This finding holds the promise of a new therapy, which intends to restore apoptosis in cancer cells by activating the full-length p53 [5].

The crystal structure of Cho et al offers the first hints for the molecular mechanisms of cancer mutations in the DBC domain [6]. Their structure shows that residues most frequently mutated in cancer are at or near the protein-DNA interface [6]; the two most frequently mutated residues (R248 and R273) directly contact DNA [6]; the remaining four hot spot residues (R175, R249, R282 and G245) appear to play a critical role stabilizing the structure of the DNA binding interface [6]. Based on Cho et al's structure, Bullock et al analyzed a wide range of missense mutants focusing on protein folding stability and DNA binding affinity [7,8]. They found that mutation sites causing loss of protein stability are mostly located in the beta-sandwich and Zinc-binding region, and mutation sites causing loss of DNA binding are at or near the DNAbinding interface [7,8]. Their analysis shows that a large fraction of cancer mutations may reduce stability of the DBC domain, resulting in loss of p53 functions at the body temperature. Thus, it is crucial to understand stability changes upon mutations in the DBC domain. To date only about 30 cancer mutants have been measured and analyzed [7-10], but about 1,300 different mutations have been reported for the DBC domain in cancer patients [4], among which about 1,000 mutations are missense.

In this work, we intend to analyze the stabilities and molecular contacts of all missense cancer mutations in the DBC domain to infer their likely molecular mechanisms. We have first validated the accuracy of three independent theoretical methods for protein stability prediction, PBSA [11], DFIRE [12], and FOLDX [13] with respect to measured relative stabilities for the DBC domain [7-10] before a comprehensive stability prediction is made for missense cancer mutants in the IARC database [4]. Based on the comprehensive stability prediction, we have analyzed the stability correlations with sequence, structure, and molecular contacts, and the correlations with clinical and functional data.

2. Results and Discussion

2.1. Overview of Relative Stabilities by Missense Mutations All IARC missense cancer mutations in the DBC domain, codons 96 to 289, are subject to three independent analyses by PBSA [11], DFIRE [12], and FOLDX [13]. Among a total of 1,006 missense mutations, 642 can be analyzed quantitatively, 152 mutations result in structural clash in homology models (see Method) or loss of Zinc binding, and 212 mutations involves PRO (from or to) and GLY (from). The last two groups of mutants can only be analyzed qualitatively. All computed relative stabilities are given in Additional file 1. In our analysis, a mutation is predicted to be stable (or unstable) only when at least two of the three predictions are consistent with each other. Among the 642 mutations analyzed quantitatively, 373 mutations are stable, i.e. $\Delta\Delta G \leq 1 \text{ kT}$ – at least 98% folded at 37°C (1 kT \approx 0.6 kcal/mol, k is Boltzmann constant, and T is thermodynamic temperature); 136 are unstable, i.e. $\Delta\Delta G > 3 \text{ kT}$ – at most 85% folded at 37°C; 97 mutations are with relative stabilities between 1 kT and 3 kT, i.e. mutations in the grey zone; and 36 mutants have inconsistent predictions by the three methods. Overall, we find 373 (36.3%) out of 794 non-PRO/GLY mutants (47.0%) are stable, 288 mutants are unstable (including Zinc-binding and clash mutants, see Method), 97 mutants (12.2%) are in the grey zone, and 36 mutants (4.5%) are without consistent predictions. Following the works of Fersht and co-workers [9,10], highly stable mutants ($\Delta\Delta G$ \leq -3 kT) are also identified in the hope to use these as suppressor mutations (Table 1). Nevertheless, a more logical approach in search of suppressor mutations would be to analyze the mutation space of the DBC domain systematically, which is actively under way in this group. In the following, only single missense mutations are analyzed (17 multiple mutations are excluded).

2.2. Missense Mutation Matrix

Fractions of unstable and stable mutations are shown in Table 2 and Table 3, respectively. In Table 2, the mutation matrix shows that 13 mutation patterns (V \rightarrow W, V \rightarrow E, I \rightarrow S, I \rightarrow T, I \rightarrow F, I \rightarrow N, F \rightarrow C, Y \rightarrow S, Y \rightarrow C, Y \rightarrow H, Y \rightarrow D, Y \rightarrow N, and R \rightarrow W) are always unstable, indicating their loss of stability is less likely due to local structural environments. Finally, Y (87.5%), I (72%), C (70%), V (67%), F (62.5%) and W (60%) are the most common amino acids before mutation in unstable mutants. W (94%), R (58%) and F (56%) are the most common amino acids after mutation in unstable mutants.

Table 3 shows the fractions of stable mutations. We found that 10 types of mutations ($S \rightarrow G$, $S \rightarrow A$, $S \rightarrow C$, $S \rightarrow T$, $F \rightarrow Y$, $Y \rightarrow F$, $K \rightarrow T$, $D \rightarrow A$, $N \rightarrow T$, and $Q \rightarrow E$) are always stable. These stable mutations do not depend on their environ-

Table 1: Highly stable mutations ($\Delta\Delta G \leq$ -3 kT) and their predicted relative stabilities by PBSA, DFIRE, and FOLDX, respectively.

MUTANTS	SEC.	$\Delta\Delta G$ (PBSA)	$\Delta\Delta G$ (DFIRE)	$\Delta\Delta G$ (FOLDX)	No. CASE.
A159V	S4	-2.27	-1.83	-1.49	41
AI6IV	S4	-3.51	-1.98	-1.14	12
N235I	S8	-2.85	-1.68	-2.04	5
N239Y	L3	-2.36	-2.41	-2.17	8
T256I	S9	-1.22	-1.96	-1.84	3
S269I	S10	-1.95	-1.99	-0.91	1

The unit of $\Delta\Delta G$ is in kcal/mol. Second column shows secondary structure elements. The last column gives the number of reported cases in the IARC database.

	G	Α	۷	L	Ι	S	С	Т	Μ	F	Y	W	Н	К	R	D	E	Ν	Q	ALL
Α	0/4		1/6			0/4		I/7		1/1						3/7				0.21
V	10/12	7/12		6/12	1/6				6/8	3/5		3/3				3/5	9/9			0.67
L			0/9		0/3	0/1			0/5	3/6			2/4		6/8				4/6	0.36
I			0/6	1/4		6/6		6/6	1/3	6/6	1/1					1/1		6/6		0.72
S	0/7	0/3		1/2	1/5		0/11	0/10		1/7	2/3			1/1	2/6			1/5		0.15
С	5/8	0/1				4/10		0/1		6/7	7/9	7/8	1/1		9/10			0/1		0.70
т		0/10			2/11	0/8			1/3					1/5	1/2			0/6		0.11
М			0/6	1/5	1/6			2/5						4/6	3/4					0.34
F	1/1		2/4	3/5	1/3	4/5	4/4				0/2									0.625
Y	1/1					6/6	6/6			0/5			7/7	1/1		7/7		6/6	1/1	0.875
W	1/1			0/1		1/1	0/1								1/1					0.60
н			0/1	2/5							2/7				4/6	1/6		1/4	2/6	0.34
к				0/1				0/3	0/3			1/1			1/5		1/4	0/4	1/3	0.17
R	3/15			0/12	1/3	2/13	0/9	2/4	0/2	1/1	1/2	6/6	3/10	1/4				1/2	3/8	0.26
D	0/7	0/3	1/8		0/1						2/8		2/7		1/1		0/7	0/8		0.12
E	2/11	0/9	0/10	0/2										5/11	1/1	1/10			1/8	0.16
Ν					1/8	0/9		0/7	0/1	1/1	3/6		2/6	0/6		1/8				0.15
Q				2/5									1/6	0/4	0/6		0/2			0.13
ALL	0.34	0.18	0.08	0.30	0.17	0.365	0.32	0.26	0.32	0.56	0.47	0.94	0.44	0.34	0.58	0.39	0.45	0.36	0.375	

Table 2: Fractions of unstable mutants (> 3 kT) from codons 96 to 289 before and after substitution.

First column shows residues before mutation and first row shows residues after mutation. Each element means fraction of unstable mutations. For example, there are a total of 3 mutations from I to M, only one of them is unstable. There are a total of 6 mutations from Y to S. All of them are unstable. Elements with 100% unstable mutations are highlighted in bold. The last column shows the percentage of total unstable mutations from WT V residue result in 67% unstable core domain. The last row shows the percentage of total unstable mutations to a given amino acid. For example, 94% mutations to W are unstable.

ment. Finally, Q (78%), N (73%), K (71%), D (70%), S (67%), T (67%) and A (66%) are the most common amino acids before mutation in stable mutants. I (76%), T (72%), L (63%) and V (62%) are the most common amino acids after mutation in stable mutants.

2.3. Correlations between Sequence/Structure and Stabilities

Stability distribution over sequence is shown in Fig. 1. It can be found that certain mutation sites are always stable or unstable no matter what amino acids are involved. Sites with more than 50% unstable mutations are defined as unstable below. Stable sites are also defined similarly. Fig. 2A shows surface distributions of stable and unstable sites, colored by blue and red, respectively. It is interesting to note that many stable sites are far from the DNA binding region. These mutations are less likely to disrupt DNA binding. Thus folding stability and binding affinity are probably not the only mechanisms for p53 inactivation by missense mutations. The likely causes of inactivation by these stable missense mutations will further be discussed below. Distributions of stable and unstable sites

	G	Α	V	L	Ι	S	С	Т	М	F	Y	W	Н	К	R	D	Е	Ν	Q	ALL
А	2/4		5/6			3/4		6/7		0/1						3/7				0.655
۷	1/12	1/12		6/12	5/6				1/8	1/5		0/3				2/5	0/9			0.24
L			5/9		2/3	1/1			1/5	3/6			1/4		1/8				0/6	0.33
I			0/6	3/4		0/6		0/6	1/3	0/6	0/1					0/1		0/6		0.10
S	7/7	3/3		1/2	4/5		11/11	10/10		6/7	1/3			0/1	4/6			3/5		0.67
С	1/8	0/1				3/10		1/1		0/7	2/9	0/8	0/1		1/10			0/1		0.14
Т		9/10			9/11	3/8			2/3					3/5	1/2			3/6		0.67
М			5/6	4/5	5/6			2/5						1/6	1/4					0.56
F	0/1		1/4	1/5	1/3	1/5	0/4				2/2									0.25
Y	0/1					0/6	0/6			5/5			0/7	0/1		0/7		0/6	0/1	0.125
W	0/1			0/1		0/1	0/1								0/1					0.0
н			1/1	3/5							5/7				2/6	2/6		2/4	3/6	0.51
к				1/1				3/3	2/3			0/ I			4/5		2/4	3/4	2/3	0.71
R	3/15			11/12	1/3	4/13	5/9	2/4	1/2	0/1	1/2	0/6	5/10	3/4				0/2	3/8	0.43
D	3/7	3/3	6/8		1/1						5/8		4/7		0/1		6/7	7/8		0.70
E	4/11	6/9	8/10	1/2										4/11	0/1	6/10			4/8	0.53
N					7/8	8/9		7/7	1/1	0/1	2/6		3/6	4/6		6/8				0.73
Q				3/5									5/6	3/4	5/6		2/2			0.78
ALL	0.31	0.58	0.62	0.63	0.76	0.365	0.52	0.74	0.36	0.38	0.47	0.0	0.44	0.47	0.38	0.43	0.45	0.43	0.375	

Table 3: Fractions of stable mutants (< I kT) from codons 96 to 289 before and after substitution.

First column shows residues before mutation and first row shows residues after mutation. Each element means fraction of stable mutations. Elements with 100% stable mutations are highlighted in bold. The last column shows the percentage of total stable mutations from a given WT amino acid. The last row shows the percentage of total stable mutations to a given amino acid. See Table 2 for more detail.

over secondary structures are shown in Fig. 2B. Visual inspection does not lead to noticeable correlation between secondary structures and stabilities. Finally Fig. 3 shows side chain exposures of unstable and stable sites, respectively. Not surprisingly almost all unstable sites are buried. This is consistent with the fact that most buried sites are hydrophobic so that mutations at these sites mostly disrupt hydrophobic packing. Indeed 93% unstable sites are buried by more than 80%, and 86% are buried by more than 90%. In contrast, most stable sites are exposed and 82% stable sites are exposed by more than 20%.

2.4. Correlation between Molecular Contacts and Stabilities

As shown above many stable cancer mutations are far from the DNA binding region. One possible reason that these DBC mutations inactivate p53 may be loss of protein-protein contacts with other p53 domains, with its tetramer subunits, or with its binding partners. It has been pointed out that specific p53-DNA binding is highly cooperative and involves interaction of p53 not only with the DNA, but also with other tetramer subunits [14]. Thus protein-protein interactions are critical in the formation of a stable p53-DNA complex, and have been implicated



Figure I

Stability distribution of IARC missense mutations from codons 96 to 289. Symbols in Italic Times: wild type amino acids; those in Normal Times: mutant amino acids; and those in Bold Times: secondary structures. Red grids: mutations with relative stabilities larger than 3 kT (including clash and Zinc-binding). Blue grids: mutations with relative stabilities less than 1 kT. Grey grids: mutations with relative stabilities between 1 kT and 3 kT. Mauve grids: mutations from or to PRO and from GLY. Yellow grids: mutations without consistent predictions.

in DNA binding and the architectural accommodation of four DBC domains to a single recognition element [14,15]. In addition, the tetramer DBC domain is stabilized both by interactions within each dimer and by interactions between dimers [16]. Such interactions are critical in stabilizing functional p53-DNA complexes in cases where specific DNA interactions are diminished as a result of truncated DBC domains [16]. Fig. 4 shows molecular contact (DNA contacts also listed) distributions of the p53 DBC domain based on crystal structures [16-19]. Here two residues are defined to be in contact when any atom in one residue is within 6 Å of any atom in the other residue. Many molecules are know to bind p53 but no complex structures are available [20-25]. This limits our molecular contact analysis. Thus only contacts observed in available structures are analyzed and shown in Fig. 4. Nevertheless, 57.3% stable sites are for protein or DNA contacts (25% for protein-only contacts, 19.1% for DNA-



Figure 2

Surface distribution of stable sites (in blue) and unstable sites (in red) (A, left). Secondary structure distribution of stable sites (in blue) and unstable sites (in red) (B, right).

only contacts, 13.2% for both protein and DNA contacts). These data imply that loss of protein contact may be a reason for p53 to lose its function upon stable mutations far from the DNA binding region.

All PRO/GLY sites are highly conserved in all mammalian p53 DBC domains so that their roles in structural integrity of the DBC domain can be assumed to be highly important. Existing DBC domain structures [6,16] are consistent with this assumption. Indeed measurements by the Fersht group show that P151S is 4.49 kcal/mol and G245S is 1.21 kcal/mol less stable than the WT DBC domain [7]. From Fig. 4, it can be found that many PRO and GLY sites, especially GLY sites (53.8%), are also involved in molecular contacts.

We found that most protein-protein contact sites (44.1%) are stable (Fig. 4). Consistent with previous analysis from the Fersht group [7,8], most DNA contact mutations



Figure 3 Side chain exposures of unstable sites (A, left) and stable sites (B, right).

(53.7%) are also stable (Fig. 1 &4). However, a few mutations are both in contact with DNA and unstable. Combining Fig. 1 and Fig. 4, it can be concluded that p53 inactivation may be due to: 1) loss of stability; 2) loss of DNA contact, protein-protein contact, or Zinc contact; 3) loss of stability and loss of DNA/protein-protein contact. Note that the above classification still cannot cover all missense cancer mutants. This results from our limited accuracy in stability prediction and limited structural data in molecular contact analysis. Nevertheless, it is clear that to activate p53 missense cancer mutants, we need to restore not only stability but also molecular contacts, especially protein-protein contacts.

2.5. Correlations between Cancer Types/Functions and Stabilities

Finally we present the distributions of unstable or stable mutations in different cancers in Fig. 5. It can be found that most reported clinical cases contain unstable mutations. Since many stable mutations are in contact with DNA (see Fig. 4) (32.3%), loss of protein folding stability and DNA binding affinity are the main mechanisms in cancers involving missense mutations, consistent with the previous experimental annotation efforts by the Fersht group. Fig. 6 shows the percentages of unstable or stable mutations in functional analysis of missense mutations. Not surprisingly, most mutations that retain functions are stable (35.7%). Interestingly most mutations that gain functions are unstable (55.4%), indicating destabilized and deformed p53 proteins are more likely to find new binding partners. It is worth to point out that the interplay between stability and function is quite general in biomolecular recognition as analyzed in a recent theoretical study [26].

2.6. Limitations

Finally, it is necessary to discuss the limitations of the pure thermodynamics analysis that was utilized here for efficiency. Indeed, the effect of mutations on folding kinetics cannot be captured with thermodynamics analysis alone. Recent studies by Mahanty et al. [27,28] suggest mutations may interfere with folding kinetics even if they can increase stability. Thus whether stabilized mutations eventually lead to functionally more robust protein cannot be understood with the thermodynamics analysis alone [29-31].

3. Methods

Protein stability change upon mutation is defined as the folding free energy difference between a mutant protein (MT) and the wild type protein (WT):

$$\Delta \Delta G = \Delta G_{\rm N-D}^{\rm MT} - \Delta G_{\rm N-D}^{\rm WT} = \Delta G_{\rm N}^{\rm MT-WT} - \Delta G_{\rm D}^{\rm MT-WT}$$

			2																					p53	p53	p53
	53 p53	po.	3					_						DINA	DINA	DINA	DINA	DINA	DINA	DINA	DINA	DINA		DINA	DINA	DINA
96	97 98	99 10	0 101 10	2 103	104	105 1	<u>06</u> 10	<mark>7</mark> 108	109 11	0 111	112 1	<mark>13</mark> 114	115	116	_ 117 _	118	119	120	_121_	122	123	124	125 12	127	128	129
p53	p53	p53	p53																							
DNA	DNA	DNA	DNA		DNA			DNA	DNA																	BP1
130	131	132 1	133 134	135	136	137	138	139	140	41 14:	2 143	144 1	45 <mark>1</mark> 4	6 14	7 148	149 1	50 15	1 152	153 1	154 15	5 156	157	158 159	160 1	61 16	2 163
I	BP2	BP2	BP2							BP2	BP2		В	P2												
I	BP1 BP	1 BP1	BP1		BP2				BF	1 BP1	BP1	BP2	В	P1												
	p53 p5	3 p53	p53 p53	3 p53	BP1	BP1		BP1	p5	3 p53	p53	p53	o53 p	53 B	P2 BF	P2 BF	2 p53	p53	p53 p	53						p53
164	165 16	6 167	168 169	9 170	171	172	173	174	175 17	6 177	178	179	180 1	81 1	82 18	33 18	4 185	186	187 1	88 18	9 190	191	192 193	194	95 19	6 197
p53	p53 p5	53 p53																			p53 p	53 p	53			
p53 198	p53 p5	53 p53	202 203	8 204	205	206	207	208 2	209 210) 211	212	213 21	4 215	5 216	6 217	218	219 2	20 22	1 222	223	p53 224 2	053 p	53 26 227	228 2	29 23	0 231
p53 198	p53 p5 199 20	53 p53 00 201	202 203	204	205	206	207	208 2	209 210) 211	212	213 21	4 <mark>215</mark>	5 216	6 217	218	219 22	20 22	1 222	223	p53 224 2	053 p 2 <mark>25</mark> 2	53 26 227	228 2	29 23	0 231
p53 198	p53 p5 199 20	53 p53 00 201	202 203	204	205	206	207	208 2 BP2	09 210) 211	212	213 21 BF	4 215 22	5 216	6 217	218	219 2	20 22	1 222	223	p53 224 2	053 p 225 2	53 26 227	228 2	29 23	0 231
p53 198	p53 p5 199 20	53 p53 00 201	202 203	3 <mark>204</mark>	205 3P2	206	207 : BP2	208 2 BP2 BP1	09 210 BP2 BP1) 211 BP2 E	212 2 3P2	213 21 BF BF	4 <mark>215</mark> 22 21 BF	5 216 P2	6 217	218	219 2	20 22	1 222	223	p53 224 2	053 p! 225 21	53 26 227	228 2	29 23	0 231
p53 198	p53 p5 199 20	53 p53 00 201	202 203	204 E	205 3P2	206	207 BP2 p53	208 2 BP2 BP1 p53	809 210 BP2 BP1 p53) 211 BP2 E BP1 E	212 3P2 3P1 B	213 21 BF BF P2 p5	4 215 22 21 BF 3 BF	5 216 P2 P1 B	217 P2 BP	218	219 22	20 22	1 222	223	p53 224 2	053 p 225 2	53 26 227	228 2	29 23	0 231
p53 198	p53 p5 199 20	53 p53 10 201	202 203	204 E	205 3P2 553	206 DNA	207 BP2 p53 DNA	208 2 BP2 BP1 p53 DNA	BP2 BP1 p53	BP2 E BP1 E p53 r	212 3P2 3P1 B 553 B	213 21 BF BF P2 p5 P1 DN	4 215 22 21 BF 3 BF	216 22 21 B	217 P2 BP P1 BP	218 2 2	219 2	20 22	1 222	223	p53 224 2	553 p 225 2	53 26 227	228 2	29 23	0 231
p53 198	p53 p5 199 20 DNA	53 p53 00 201	202 203	E E 238 2	205 3P2 553 2NA	206 DNA	207 BP2 p53 DNA 241	208 2 BP2 BP1 p53 DNA	809 210 BP2 BP1 p53 DNA 243	BP2 E BP1 E p53 p	212 3P2 3P1 B 53 B	213 21 BF BF P2 p5 P1 DN	4 215 22 21 BF 3 BF IA DN	216 22 21 B NA B	217 P2 BP P1 BP	218 2 1 0 251	219 2	20 22	1 222	223	p53 p 224 2	225 2	53 26 227	228 2	29 23	<u>0 231</u>
p53 198 232	p53 p5 199 20 DNA 233 23	53 p53 50 201 84 235	202 203 236 237	204 E F D 238 2	205 3P2 553 0NA 1 239	206 DNA 240	207 BP2 p53 DNA 241	208 2 BP2 BP1 p53 DNA 242	809 210 BP2 BP1 p53 DNA 243	BP2 E BP1 E p53 p 244 2	212 2 3P2 3P1 B 53 B 245 2	213 21 BF BF P2 p5 P1 DN 46 24	4 218 22 21 BF 3 BF IA DN 7 24	216 22 21 B 1A B	217 P2 BP P1 BP 49 25	218 2 1 0 251	219 2	20 22 253 25	1 222 54 255	223	p53 p 224 2 257 25	53 pl 225 2 58 259	53 26 227 260 26	228 2	29 23 263 26	0 231 64 265
p53 198 232	p53 p5 199 20 DNA 233 23	3 p53 0 201 34 235	202 203 236 237	204 E P D 238 2 BP2	205 3P2 553 DNA 1 239	206 DNA 240	207 BP2 p53 DNA 241	208 2 BP2 BP1 p53 DNA 242	809 210 BP2 BP1 p53 DNA 243	BP2 E BP1 E p53 p 244 2 BP2	212 2 3P2 3P1 B 553 B 245 2 BP2	213 21 BF BF P2 p5 P1 DN 46 24	4 215 22 21 BF 3 BF IA DN 7 24	216 22 21 B 1A B 18 2 055	217 P2 BP P1 BP 49 25	218 2 1 0 <u>251</u>	219 2 252 2	20 22 253 25 53	1 222 54 255	223 256 2	p53 (224 2 257 25	553 pl 225 2 58 259	53 26 227 260 26	228 <mark>2</mark> 1 262	29 23 263 26	0 231 64 265
p53 198 232	p53 p5 199 20 DNA 233 28	3 p53 0 201 34 235	202 203 236 237	204 E F D 238 2 BP2 DNA	205 3P2 553 0NA 1 239	206 DNA 240	207 BP2 p53 DNA 241	208 2 BP2 BP1 p53 DNA 242	809 210 8 BP2 8 BP1 953 8 DNA 243	BP2 E BP1 E p53 p 244 2 BP2 BP2	212 2 3P2 3P1 B 553 B 245 2 BP2 DNA	213 21 BF BF P2 p5 P1 DN 46 24	4 215 22 21 BF 3 BF IA DN 7 24	22 22 21 B NA B 8 2 p53	217 P2 BP P1 BP 49 25 3 p53	218 2 1 0 251 E	219 2 252 2 3P1 p	20 22 253 25 53 3	1 222 54_255	223	p53 (224 2 257 25	553 pl 225 2 58 259	53 26 227 260 26	228 2 1 262	29 23 263 26	0 231 64 265
p53 198 232	p53 p5 199 20 DNA 233 23	3 p53 0 201 34 235	202 203 236 237	204 E F D 238 2 BP2 DNA	205 3P2 553 0NA 239 DNA	206 DNA 240 DNA	207 BP2 p53 DNA 241	208 2 BP2 BP1 p53 DNA 242 DNA	809 210 BP2 BP1 p53 DNA 243 DN/	BP2 E BP1 E p53 g 244 2 BP2 DNA	212 2 3P2 3P1 B 553 B 245 2 BP2 DNA	213 21 BF BF P2 p5 P1 DN 46 24 DN	4 215 22 21 BF 3 BF IA DN 7 24	22 22 21 B NA B 18 2 p53 A DNA	217 P2 BP P1 BP 49 25 3 p53 A DNA	218 2 1 0 251 E	219 22 252 2 3P1 p DNA DI	20 22 253 25 53 VA	1 222 54 255	223	p53 224 2 257 25	53 pt 225 2 58 259	53 26 227 _260 26	228 2 1 262	29 23 263 26	0 231 64 265

Figure 4

Distribution of molecular contacts. The DNA contacts (as in 2AC0, 2ADY, 2AHI, 2ATA (16)) are denoted by 'DNA', the tetramer subunit contacts (as in 2AC0, 2ADY, 2AHI, 2ATA (16)) are denoted by 'p53', the contacts with p53-binding protein I (as in IGZH (17) and IKZY (18)) or 2 (as in IYCS (19)) are denoted by 'BP1' or 'BP2', respectively. Unstable sites are shown by red grids; stable sites by blue grids; Pro/Gly residues on wild type by mauve grids.

Here, the folding free energy of a protein is the free energy difference between its native state (N) and denatured state (D): $\Delta G'_{N-D} = G'_{N}-G'_{D}$, with ' = MT or WT Prediction of protein stabilities has attracted much attention in computational biology with many computational models proposed [12,32-35] that take into account various factors important for protein stability [36-41].

In this work, we have chosen three well-established computational methods to predict protein stability change upon mutations. The first method is PBSA [11,42-44] that can be used to estimate stability if we assume hydrophobic and electrostatic (salt-bridge) interactions are the predominant components in stabilizing proteins. Here the hydrophobic free energy is estimated by a term linearly proportionally to the solvent accessible surface area (SA), and the electrostatic free energy is computed by solving the Poisson-Boltzmann (PB) equation [45]. Similar to protein pKa computations, the protein dielectric constant is set to be 20 to address electronic polarization, rotational polarization, and ionization effect. [46-48] The solvent dielectric constant is 80. The solvent ionic strength is set to be 150 mM. All calculations were performed at temperature 283 K. The second method is DFIRE by Zhou and Zhou [12] who have constructed a new residue-specific all-atom potential of mean force from nonhomologous protein structures with their proposed new reference state. The third method is FOLDX by Guerois et al [13] who

have developed an efficient all-atom free energy function with weighted free energy terms trained using empirical data from experimental stabilities. After a control study of three models with available experimental data for the DBC domain, we use all three methods to comprehensively analyze the missense cancer mutants in the IARC database.

Crystal structure (codons 96 to 289), b chain of 1TSR [6] is used for the native state of the p53 DBC domain. Homology models are obtained by SCWRL3 [49] for the native state of missense mutants. If necessary, the denatured state is modeled as a tripeptide centered at the mutation site.

In this work, mutations causing structural clash and disruption of Zinc binding-sites (176, 179, 238, and 242) are assumed to be unstable by more than 3 kT. Indeed, these mutants are generally more unstable as observed in experiment (with $\Delta\Delta G$ ranging from 2.75 to 4.78 kcal/mol for F134L, V157F, H168R, R175H, M237I, I255F, R282W, T123A, H168R, and C242S [7,9,50]. Mutations from PRO/GLY and mutations to PRO are not covered in quantitative analysis because such mutations generally cause significant backbone entropy changes that cannot be modeled without time-consuming molecular dynamics analysis. However, they are analyzed qualitative based on sequence comparison of all mammalian p53.



Percentages of unstable (red), stable and DNA contact (blue), and stable and no DNA contact (green) mutations at different tumor sites. Total number of mutations is also given by each tumor site.

Quantitative agreements between experimental data and theoretical data from PBSA, DFIRE, and FOLDX, respectively, are shown in Fig. 7. Corresponding linear correlation coefficients are 0.91, 0.89 and 0.79, respectively; root mean square deviations are 0.80, 0.88, 1.25 kcal/mol, respectively; and unsigned average errors are 0.62, 0.75, 0.98 kcal/mol, respectively. In this work, three qualitative predictions are used to analyze all missense mutants: 1) relative stabilities less than 1 kT; 2) relative stabilities larger than 3 kT; and 3) relative stabilities less than -3 kT - highly stable mutations. Success rates of these qualitative predictions are also given in Table 4. The overall accuracies for the 1 kT prediction are 93%, 85%, and 85% by PBSA, DFIRE, and FOLDX, respectively; those for the 3 kT prediction are 96%, 91%, 93%, respectively; and those for the -3 kT prediction are 98%, 96%, 91%, respectively. Overall, excellent accuracies can be achieved by the three theoretical methods.

4. Conclusion

Most human cancers contain mutations in the transcription factor p53 and majority of these are missense and located in the DNA binding core domain. In this study, the stabilities of all core domain missense mutations are predicted and are used to infer their likely inactivation mechanisms. Overall, 47.0% non-PRO/GLY mutants are



Figure 6

Percentages of unstable (red), stable and DNA contact (blue), and stable and no DNA contact (green) mutations in different functional properties. Retained: functional property of the mutant is similar to that of the WT; temp: the activity of the mutant is affected by the temperature at which the experiment is performed; domi-neg (no): dominant negative activity that the mutant does not counteract the effects of the WT under the control of a p53responsive element when the two proteins are co-expressed in human or yeast cells; domi-neg (yes): the mutant protein counteract (or has a partial inhibiting effect on) the activity of the WT; lost: the WT functional property is lost by the mutant; gain: functional properties displayed by the mutant is not displayed by the WT protein. The total number of mutations is given by each functional property.

stable ($\Delta\Delta G < 1.0$ kT) and 36.3% mutants are unstable ($\Delta\Delta G > 3.0$ kT), 12.2% mutants are with 1.0 kT < $\Delta\Delta G < 3.0$ kT. Only 4.5% mutants are with no conclusive predictions. Certain types of either stable or unstable mutations are found not to depend on their local structures. Y, I, C, V, F and W (W, R and F) are the most common residues before (after) mutation in unstable mutants. Q, N, K, D, A, S and T (I, T, L and V) are the most common residues before (after) mutation in stable mutants. The stability



Figure 7

Correlations between the three theoretical calculations and the experimental measurements by the Fersht group. From left to right, the figure is for PBSA, DFIRE and FOLDX, respectively.

		PBSA	DFIRE	FOLDX
≤ I kT	ТР	91%	83%	89%
	TN	96%	87%	81%
	ACCU.	93%	85%	85%
> 3 kT	ТР	100%	100%	90%
	TN	93%	87%	96%
	ACCU.	96%	91%	93%
≤ -3 kT	ТР	80%	75%	50%
	TN	100%	98%	95%
	ACCU.	98%	96%	91%

Table 4: Accuracies for qualitative predictions, by PBSA, DFIRE, and FOLDX, respectively.

TP: accuracy for true positive predictions; TN: accuracy for true negative predictions; ACCU: accuracy for overall accuracy combining TP and TN.

correlations with sequence, structure, and molecular contacts are also analyzed. No direct correlation between secondary structure and stability is apparent, but a strong correlation between solvent exposure and stability is noticeable. Our correlation analysis shows that loss of protein-protein contacts may be an alternative cause for p53 inactivation. Correlation with clinical data shows that loss of stability and loss of DNA contacts are the two main inactivation mechanisms. Finally, correlation with functional data shows that most mutations which retain functions are stable, and most mutations that gain functions are unstable, indicating destabilized and deformed p53 proteins are more likely to find new binding partners.

Additional material

Additional file 1

Quantitative stability predictions of all missense mutants. Click here for file [http://www.biomedcentral.com/content/supplementary/1757-5036-2-5-S1.dat]

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