

# Perturbation by sequence variation: impacts of coding mutations on protein fitness

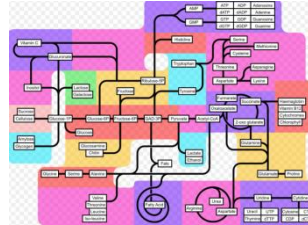
Anna Panchenko

National Center for Biotechnology Information, NIH

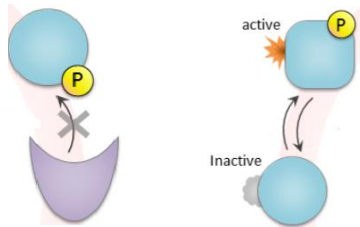


# What do we study?

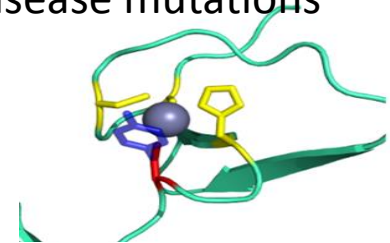
Biochemical pathways  
and interactomes



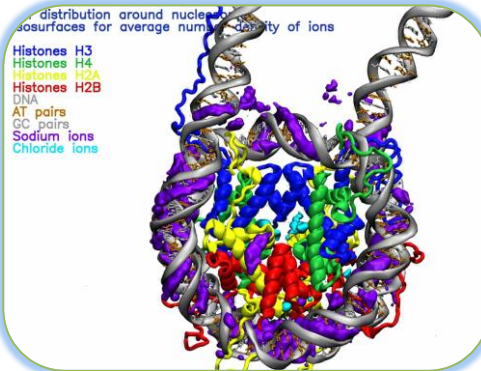
PTMs, phosphorylation



Genetic variants,  
disease mutations



Nucleosomes



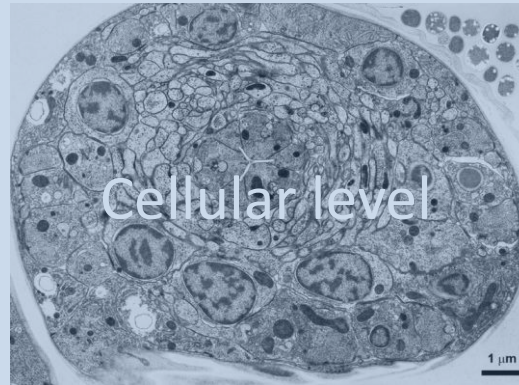
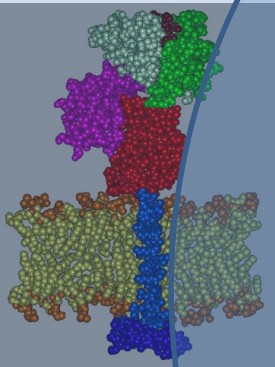
# Methods

<b>Bioinformatics</b>	Sequence alignment, structure superposition, annotation of domains, intrinsically disordered regions, functional sites Programs: Blast, Vast, Muscle, IBIS, SPEER, CDD, ...
<b>Structural Modeling</b>	Homology modeling of protein structures and structural complexes Programs: VMD, NAMD
<b>Energy calculations</b>	Empirical and statistical energy potentials, Molecular Mechanics Poisson-Boltzmann approach Force fields and programs: CHARMM27, CHARMM36, FoldX, BeatMusic and PopMusic,...
<b>Dynamics</b>	All-atom Molecular Dynamics simulations Programs: NAMD
<b>Evolutionary analysis</b>	Evolutionary conservation, phylogenetic analysis Programs: Mega, PAML, FastTree, ...

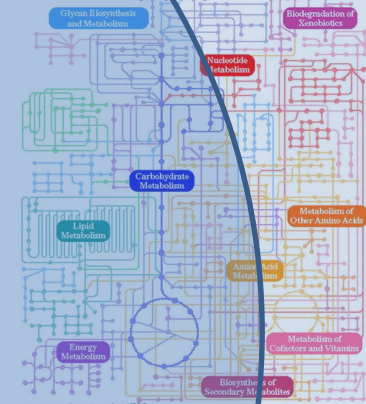
# Organismal level



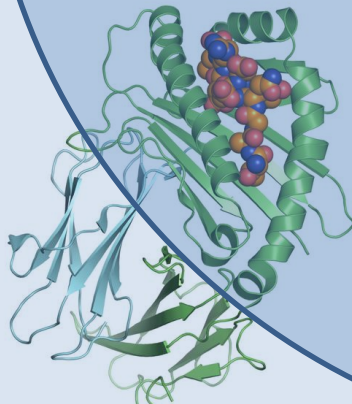
## Interactions



## Pathways



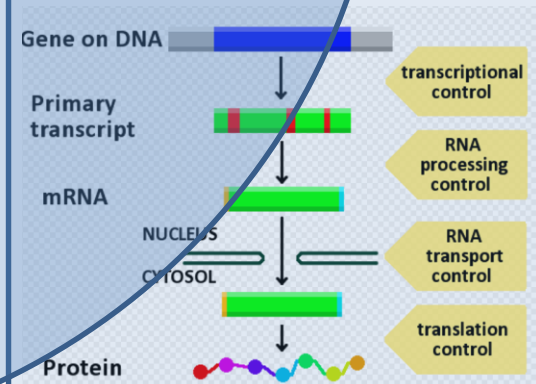
## Foldability & stability



AMKLPYTTTRDAAGHKLITR

# Molecular level

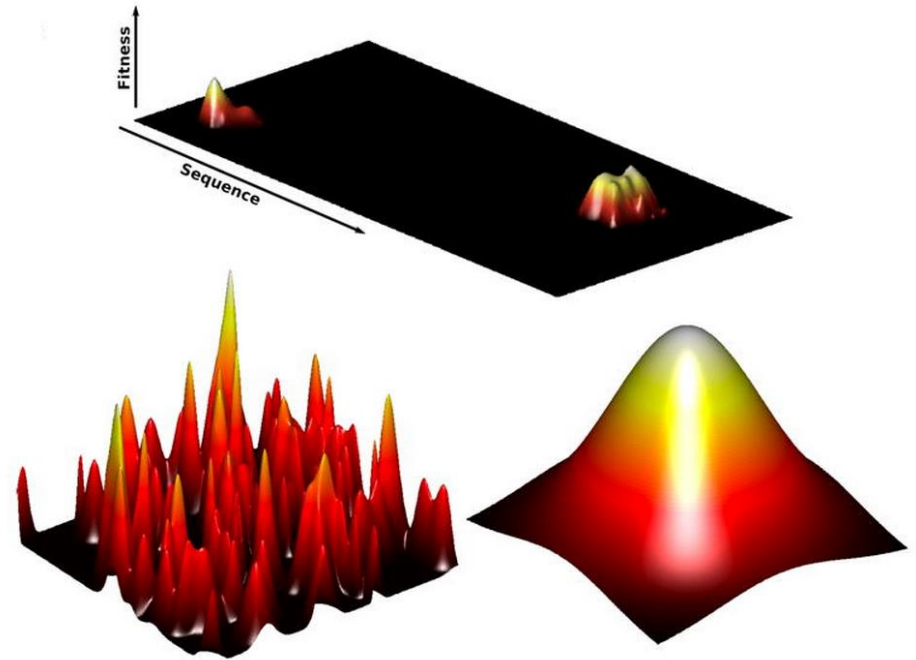
## Expression & regulation



# Variation may arise through genetic mutations and rearrangements

## Types of variations:

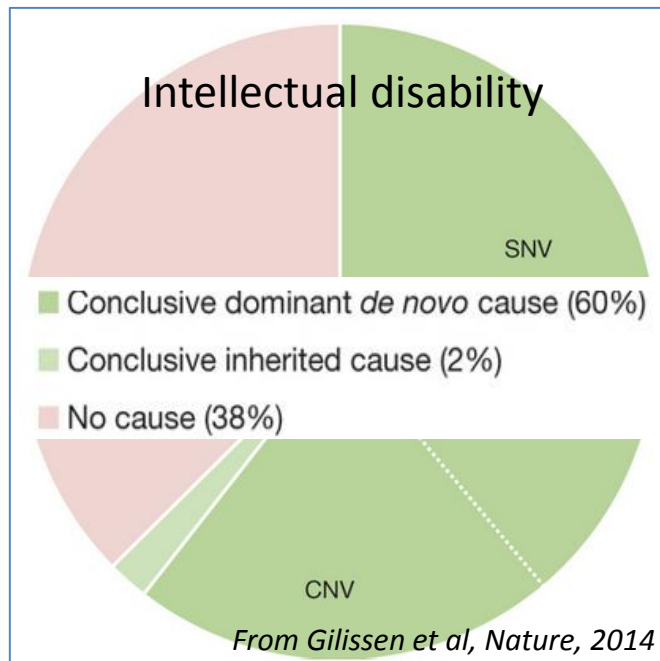
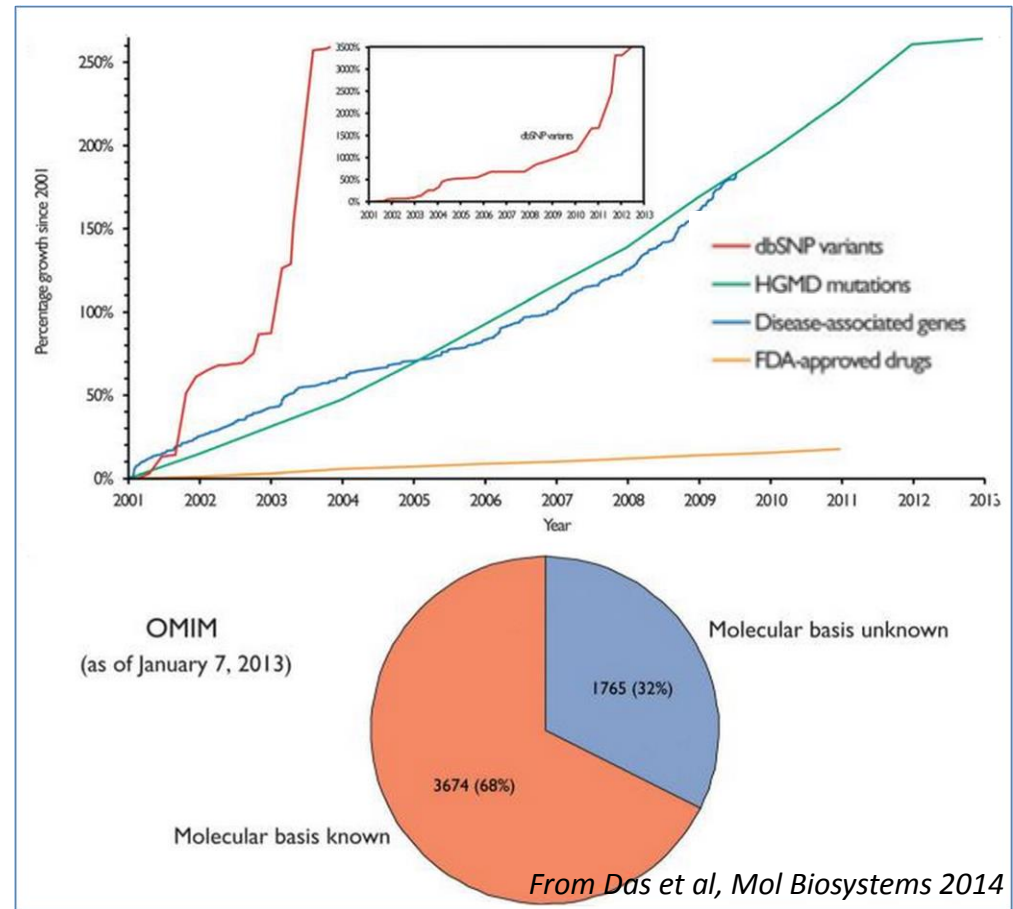
- missense mutations – change in amino acid type;
- Insertion and deletions;
- Polypeptide chain truncations;
- Domain shuffling;
- Post-translational modifications



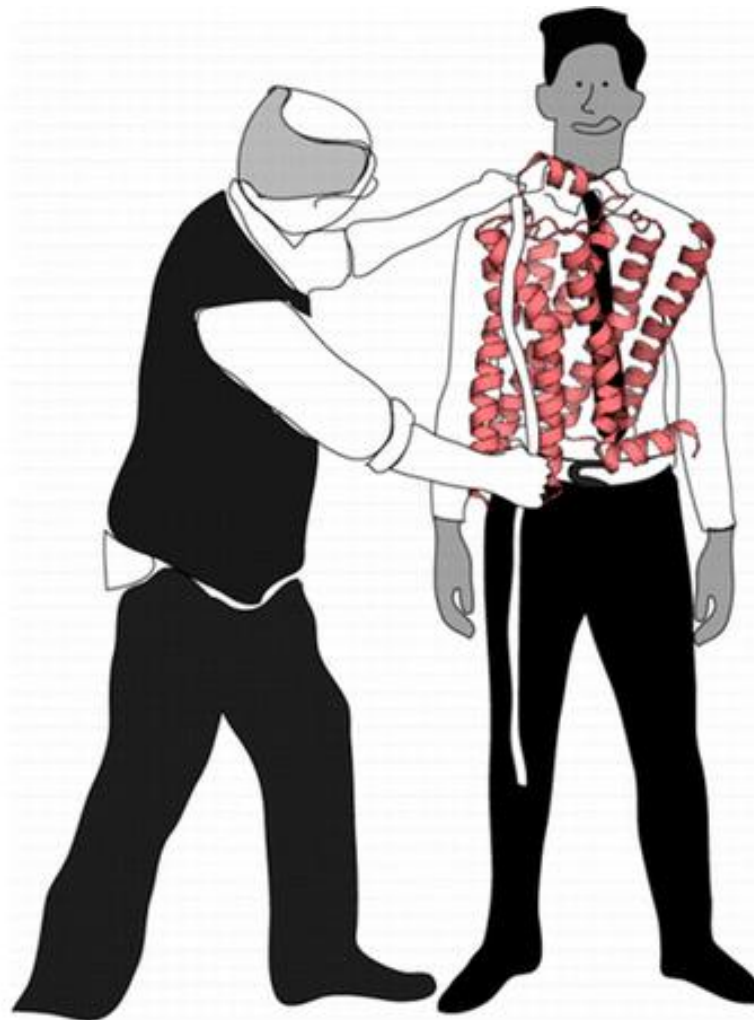
From Romero & Arnold, *Nat Rev Mol Cell Biol.* 2009

# Hundreds of mutations with unknown molecular mechanisms

Type of variation	Number in human
<i>De novo</i> variants	50-100
SNV	3.5 million
nsSNV, sSNV	10,000-15,000
Protein loss-of-function nsSNV (from HGMD)	50-500 (50-100)



# Personal genomics meets biophysics



*From Kroncke et al, Biochemistry 2015*

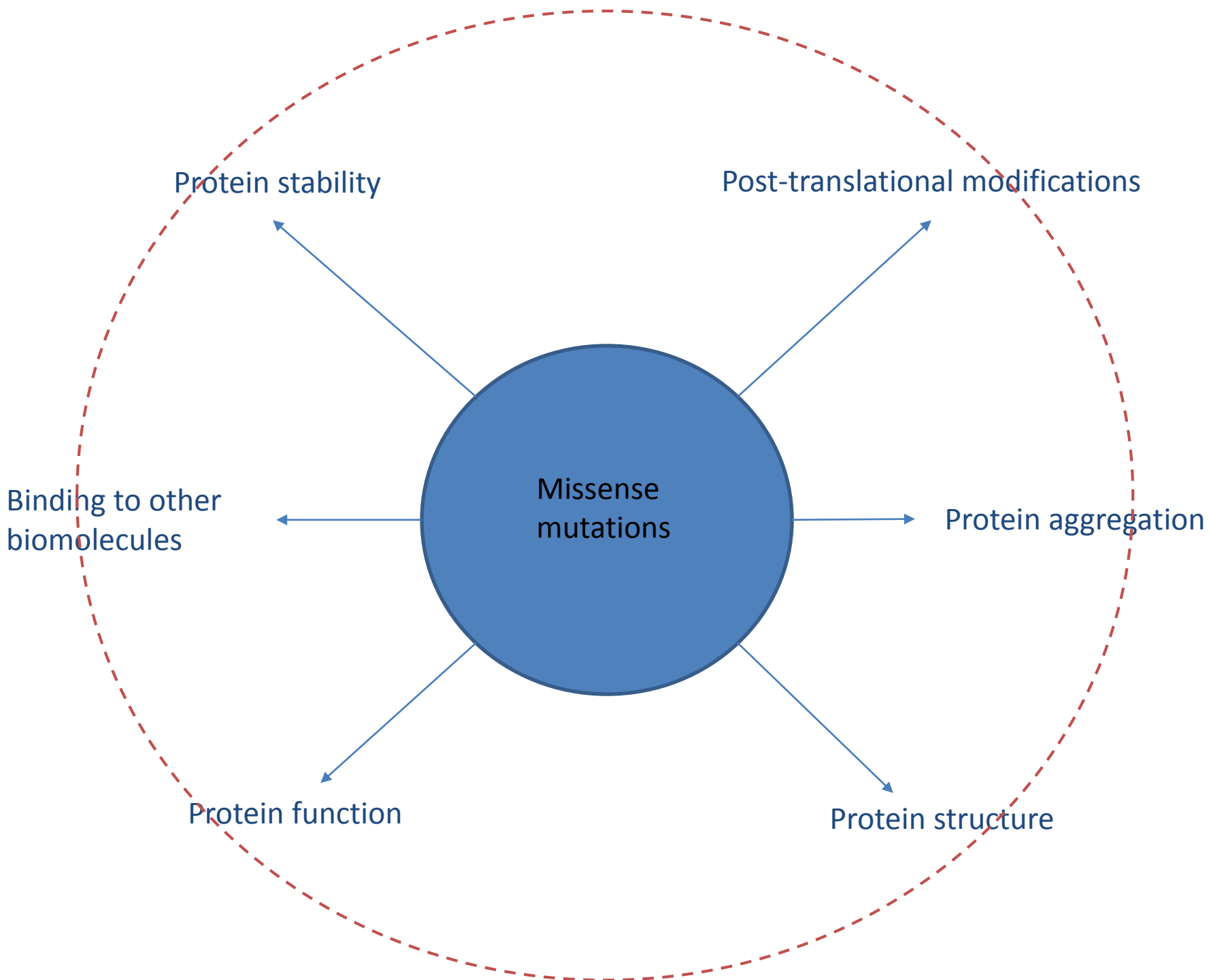
# Talk synopsis

- Impact of missense mutations on proteins: stability, binding and activity
- Deciphering the human protein interactome with interactions with resolved binding interfaces



# Why do we need to learn about the mechanisms of effect of mutations on proteins?

- To decipher how proteins evolved.
- To predict which mutations are damaging.
- To distinguish functionally important mutations, distinguish driver from passenger mutations.
- Prioritize mutations for experimental research.
- Drug design.



Protein stability

Post-translational modifications

Missense mutations

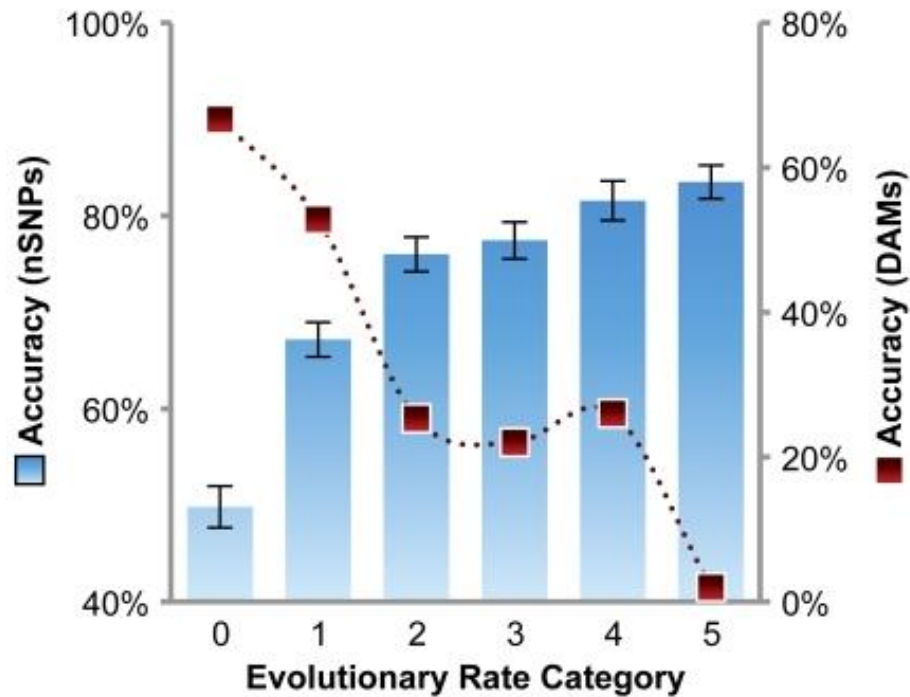
Protein aggregation

Binding to other biomolecules

Protein function

Protein structure

# Evolutionary conservation is related to functional importance

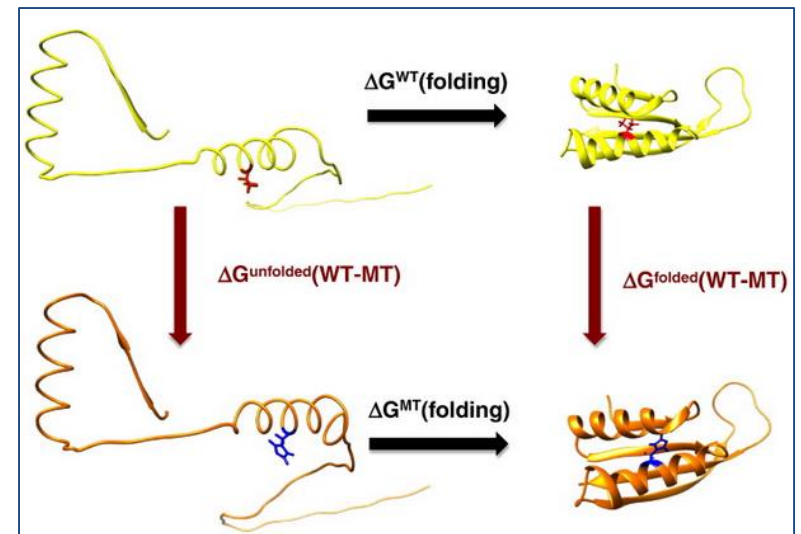
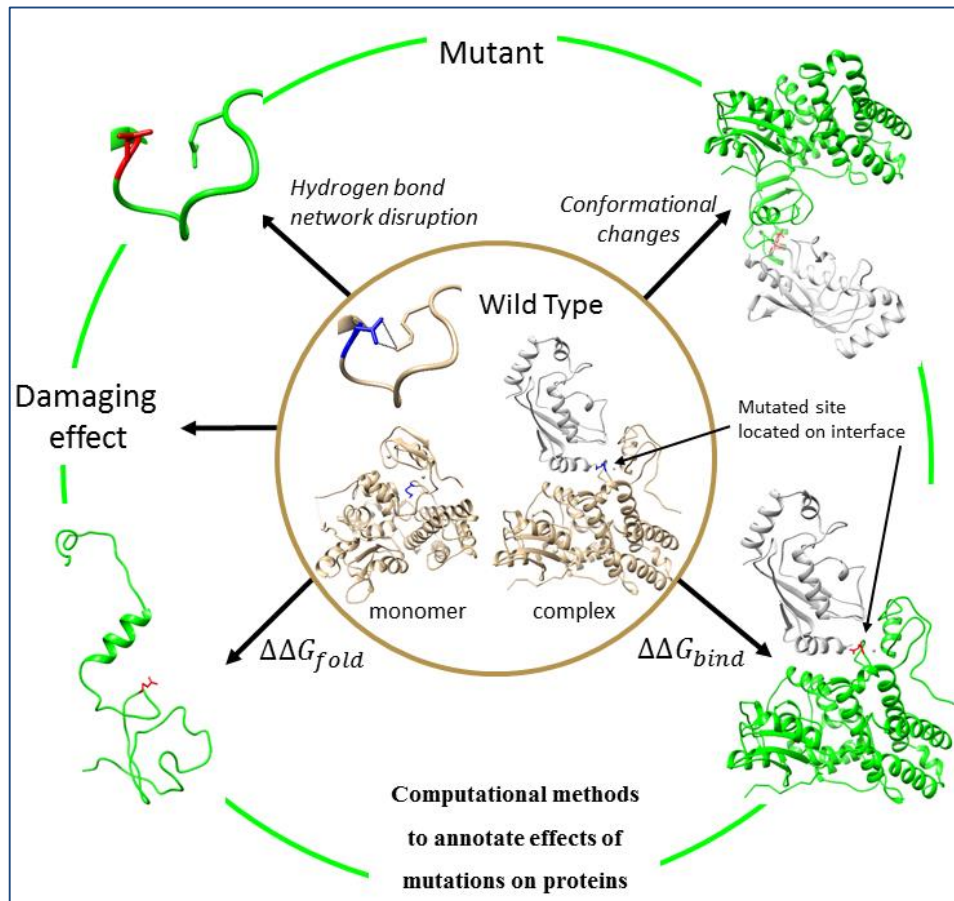


From Kumar et al, *Genome Research*, 2009

Mutations in functionally relevant sites might be damaging. Many methods exploit this observation.

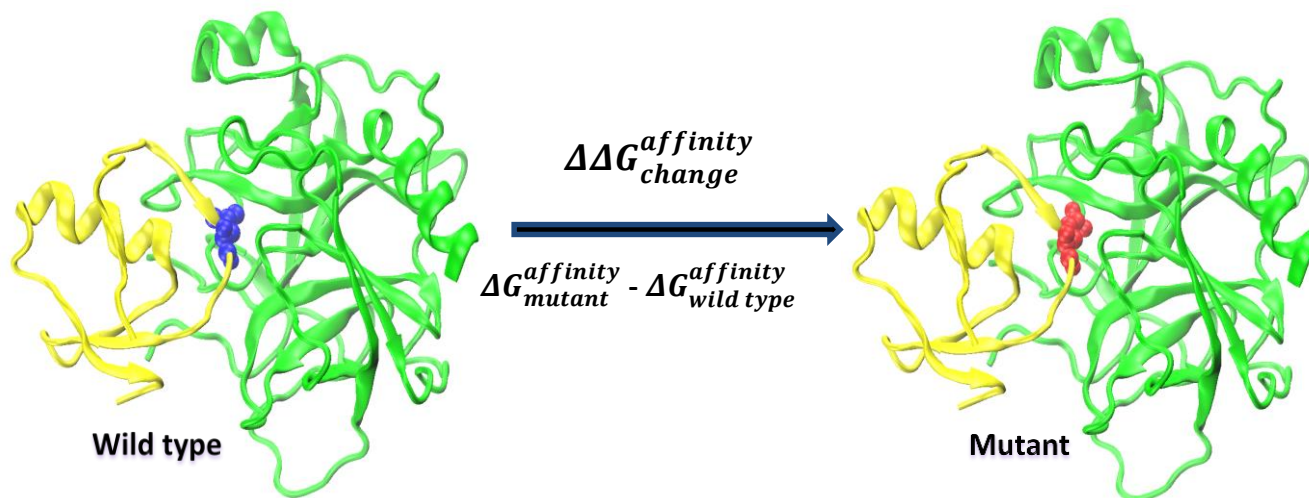
Methods that predict functional effects of mutations on proteins use evolutionary conservation

# Deciphering of molecular basis of mutational impacts



# Effects of missense mutations on protein-protein binding affinity

$$\Delta\Delta G_{bind} = \Delta G_{wt} - \Delta G_{mut}$$

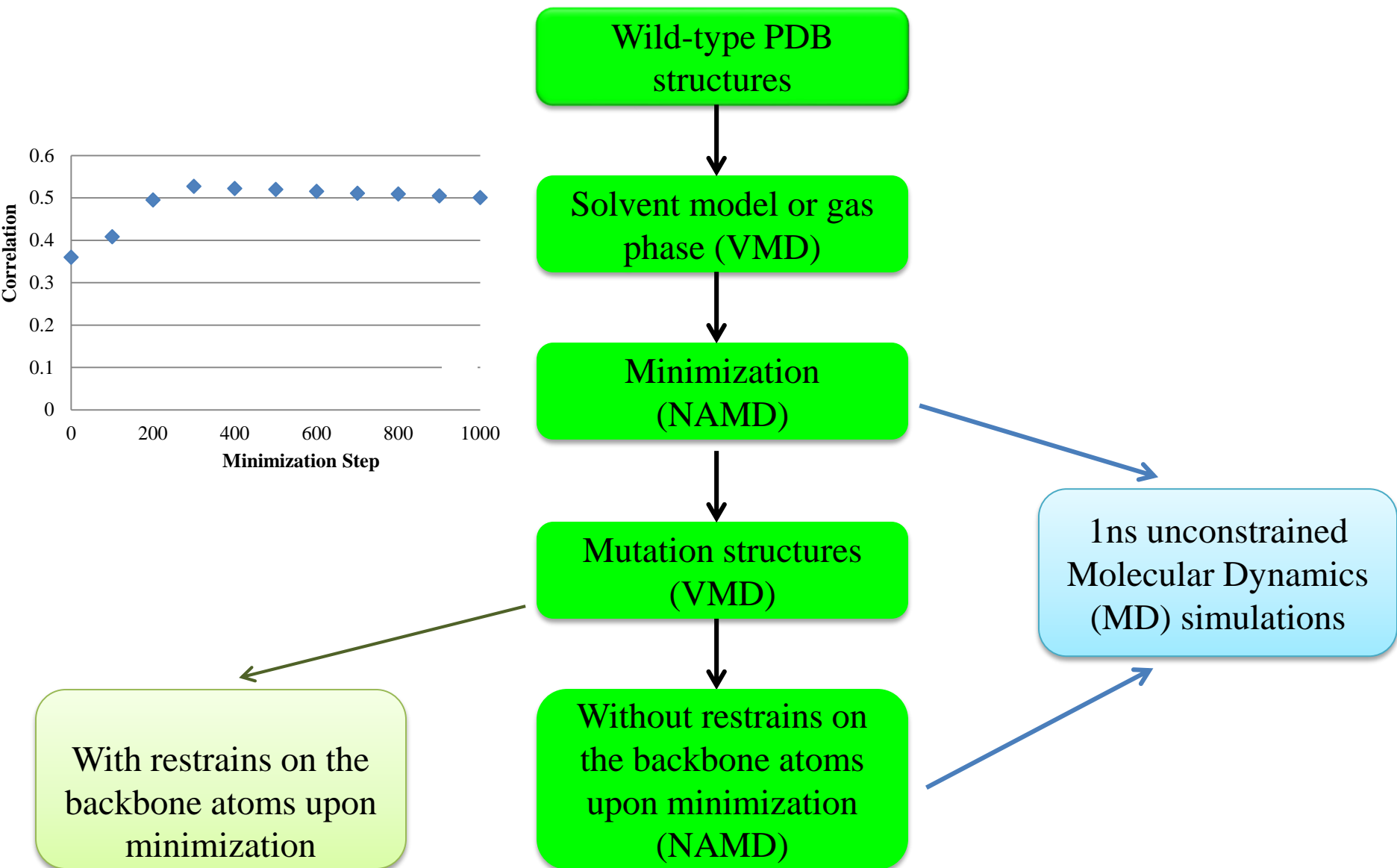


$$G = E_{gas}^{MM} + G_{solv}^p + G_{solv}^{np} - TS$$

Modified Molecular Mechanics Poisson-Boltzmann Surface Area approach

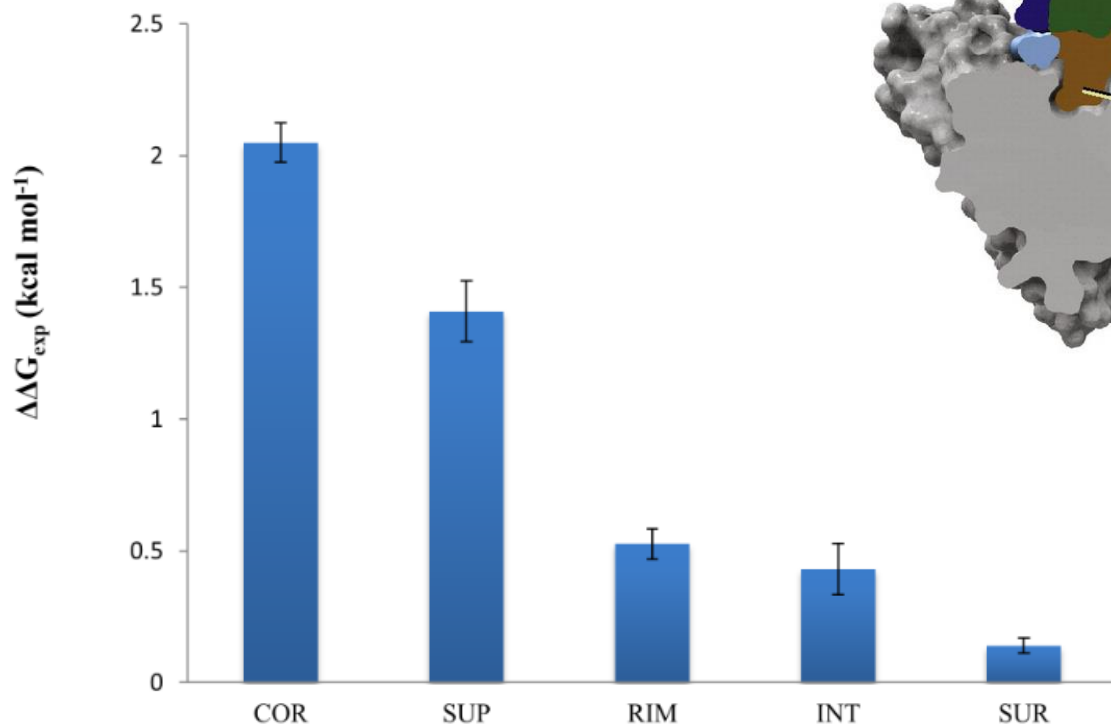
$$\Delta\Delta G_{Pred1}^{bind} = \alpha\Delta\Delta E_{vdw} + \beta\Delta\Delta G_{solv} + \gamma\Delta SA_{mut} + \delta$$

# How to minimize mutant and wild type structures

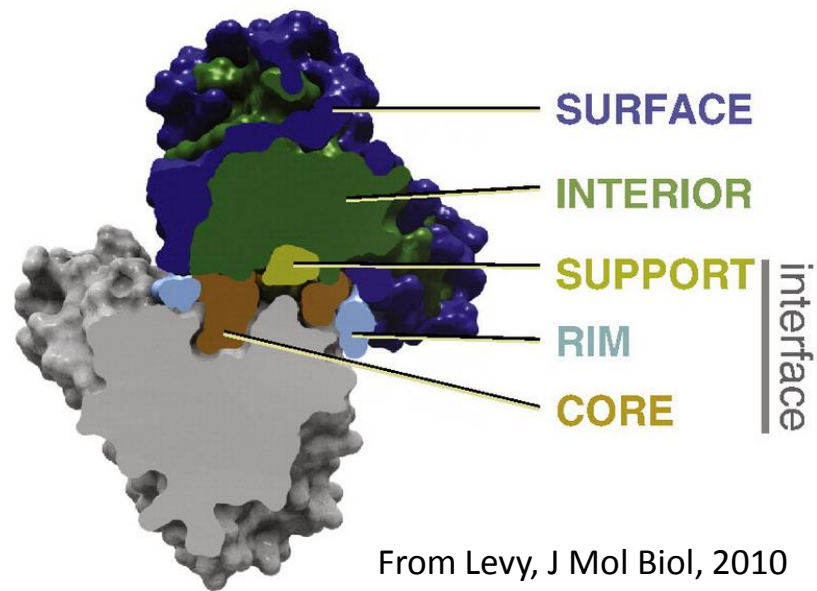


# Factors influencing prediction quality

## 1. Locations of mutations



Cross-section of a protein complex



# Factors influencing prediction quality

## 2. Volume and charge of substituted amino acids

<b>Wild-type</b>	<b>Mutant</b>		
	<b>Small</b>	<b>Medium</b>	<b>Large</b>
	R/# mutations	R/# mutations	R/# mutations
<b>Small</b>	0.52/97	0.51/123	0.67/39
<b>Medium</b>	0.61/590	0.58/450	0.34/130
<b>Large</b>	0.63/210	0.64/142	0.58/63

<b>Wild-type</b>	<b>Mutant</b>		
	<b>Negative</b>	<b>Neutral</b>	<b>Positive</b>
	R/# mutations	R/# mutations	R/# mutations
<b>Negative</b>	-	0.33/232	-
<b>Neutral</b>	0.72/86	0.58/1042	0.48/89
<b>Positive</b>	0.81/33	0.67/300	-



# Factors influencing prediction quality

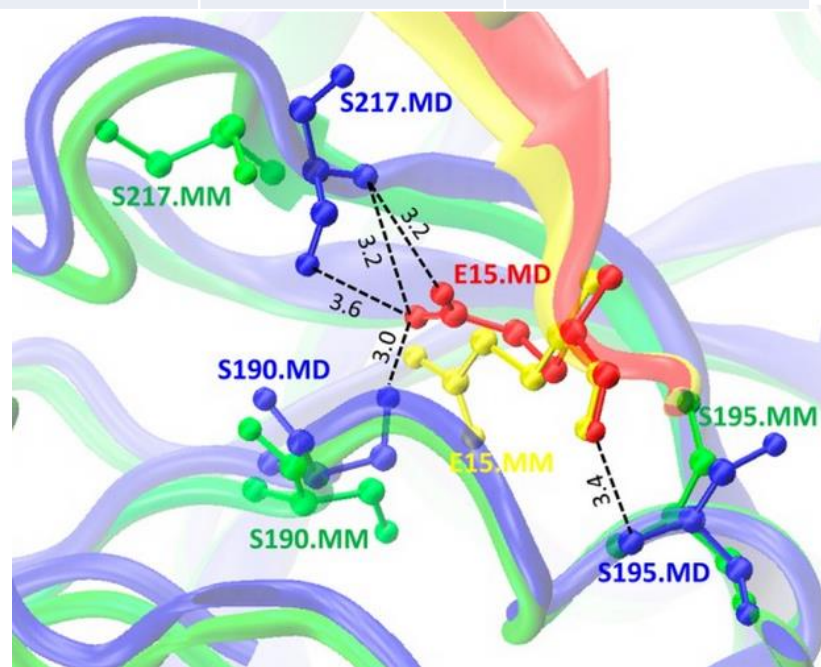
3. Low accuracy of predicting stabilizing mutations, probably due to lack of stabilizing mutants in experimental data sets

Method	Training/testing	Destabilizing	Stabilizing
Pred2	SKEMPI/NM	100%	7%
CC/PBSA	NM/NM	99%	32%
FoldX	test: NM	72%	48%
	test: SKEMPI	67%	41%
BeatMusic	test: NM	95%	23%
	Test: SKEMPI	90%	18%

# Factors influencing prediction quality

4. Difficult to adequately account for the flexibility of proteins.

Simulation method	Flexibility	Cross-validated R	RMSE (kcal/mol)
Minimization	Flexible backbone	0.61	1.22
1ns MD simulations	Flexible backbone	0.26	1.48





Home



Results



Help

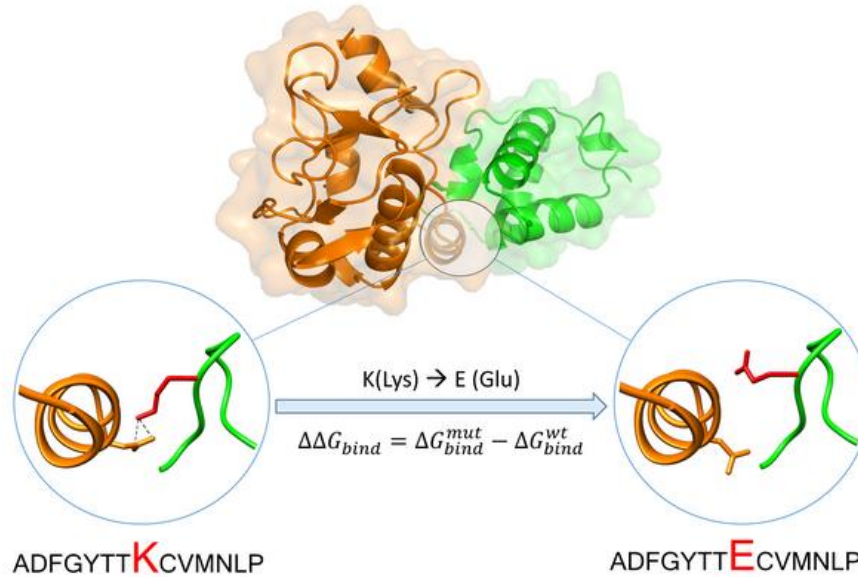


Method



References

**MutaBind** evaluates the effects of variations and disease mutations on protein-protein interactions. It predicts if a mutation may disrupt an interaction and calculates the changes in binding affinity. Structure of protein-protein complex is required for this method.



## Step 1 - Select Protein Complex

Input PDB code:

Example: 1CSE



Bioassembly



Asymmetric Unit

Upload PDB file:

No fil...osen

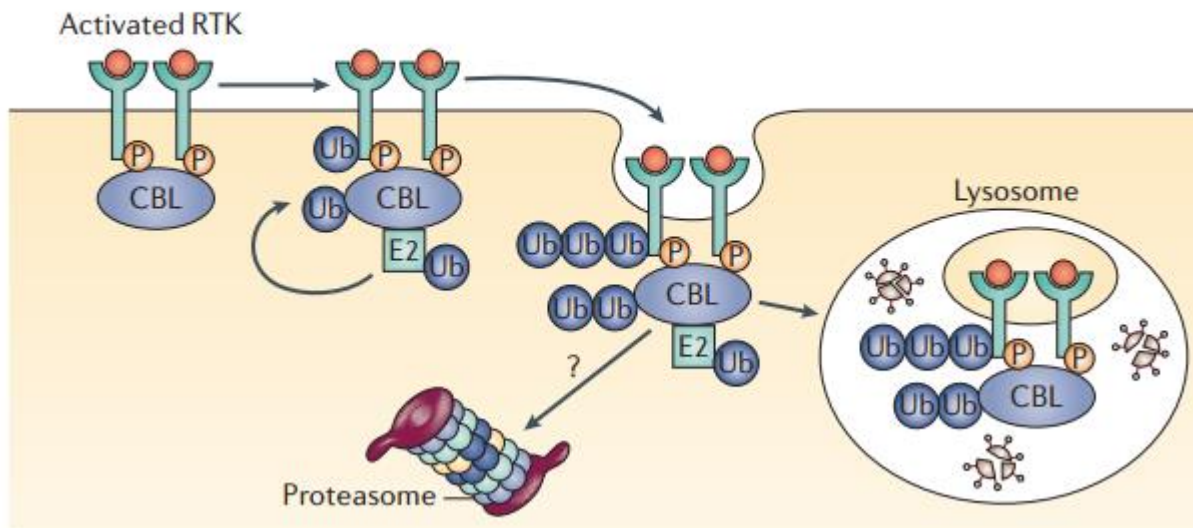
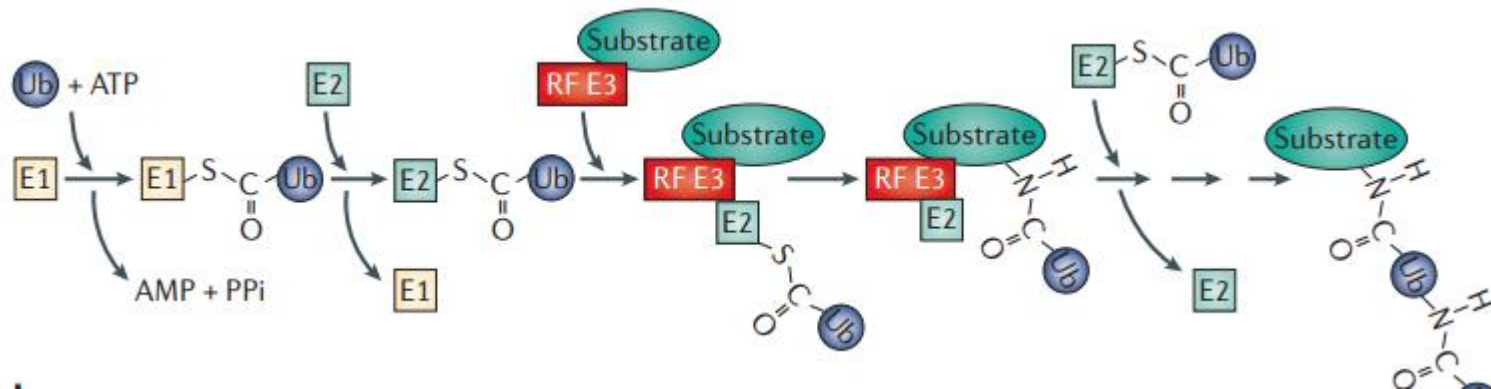
Additional Options

Is protease-inhibitor complex?

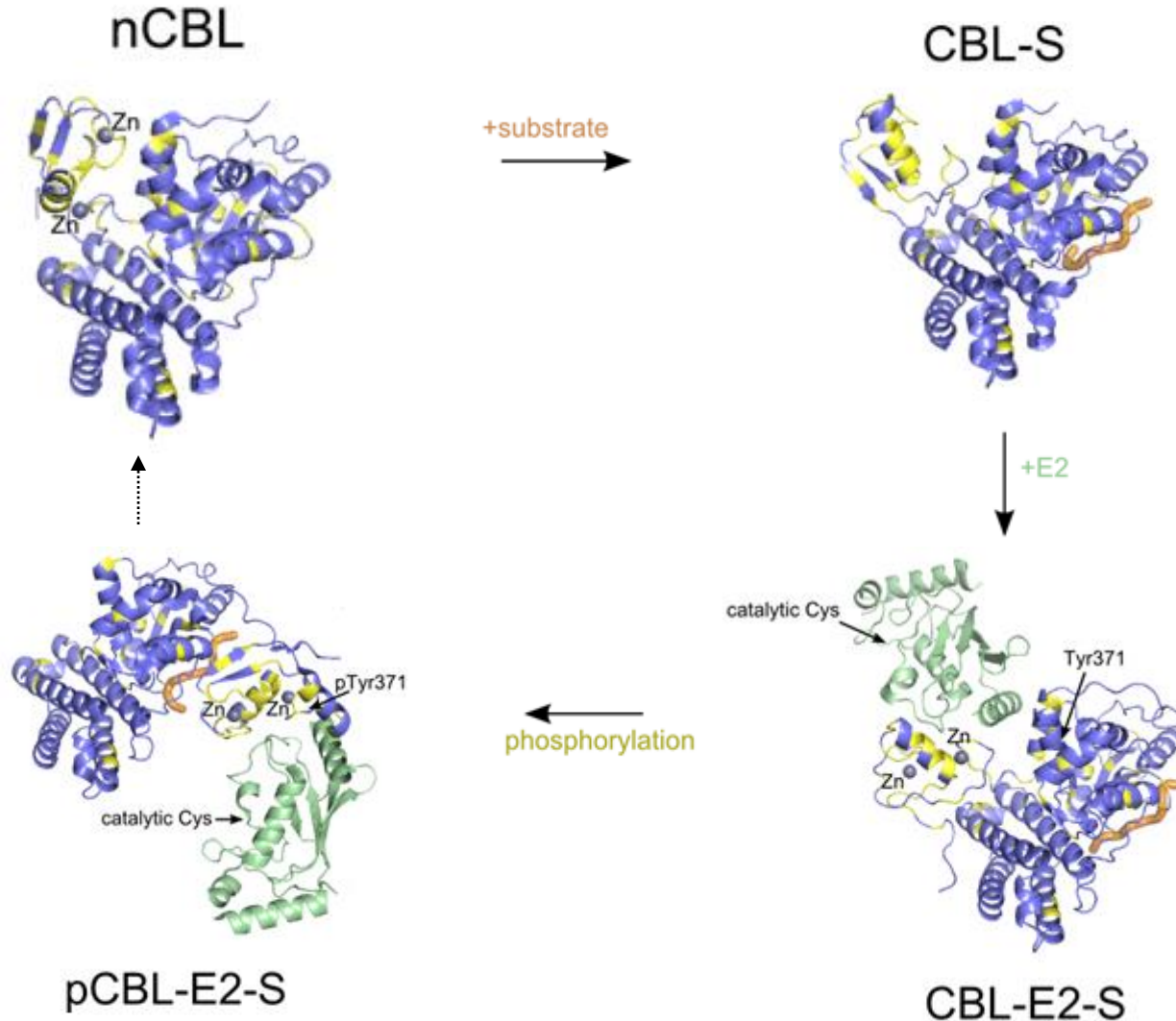
Next

# Mechanisms of action of cancer mutations: CBL case

# CBL ubiquitin ligase - CBL

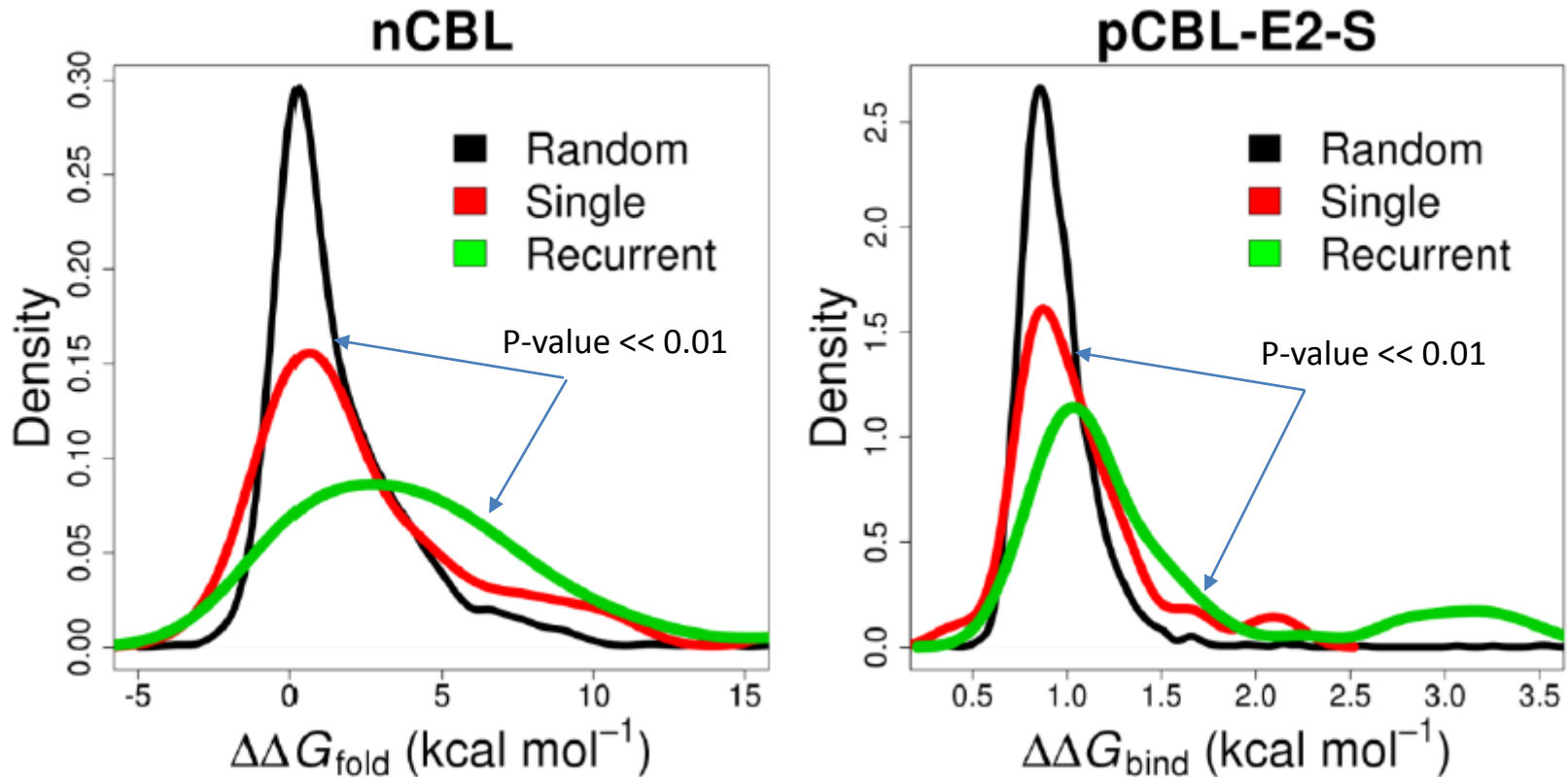


# CBL ubiquitin ligase activation cycle

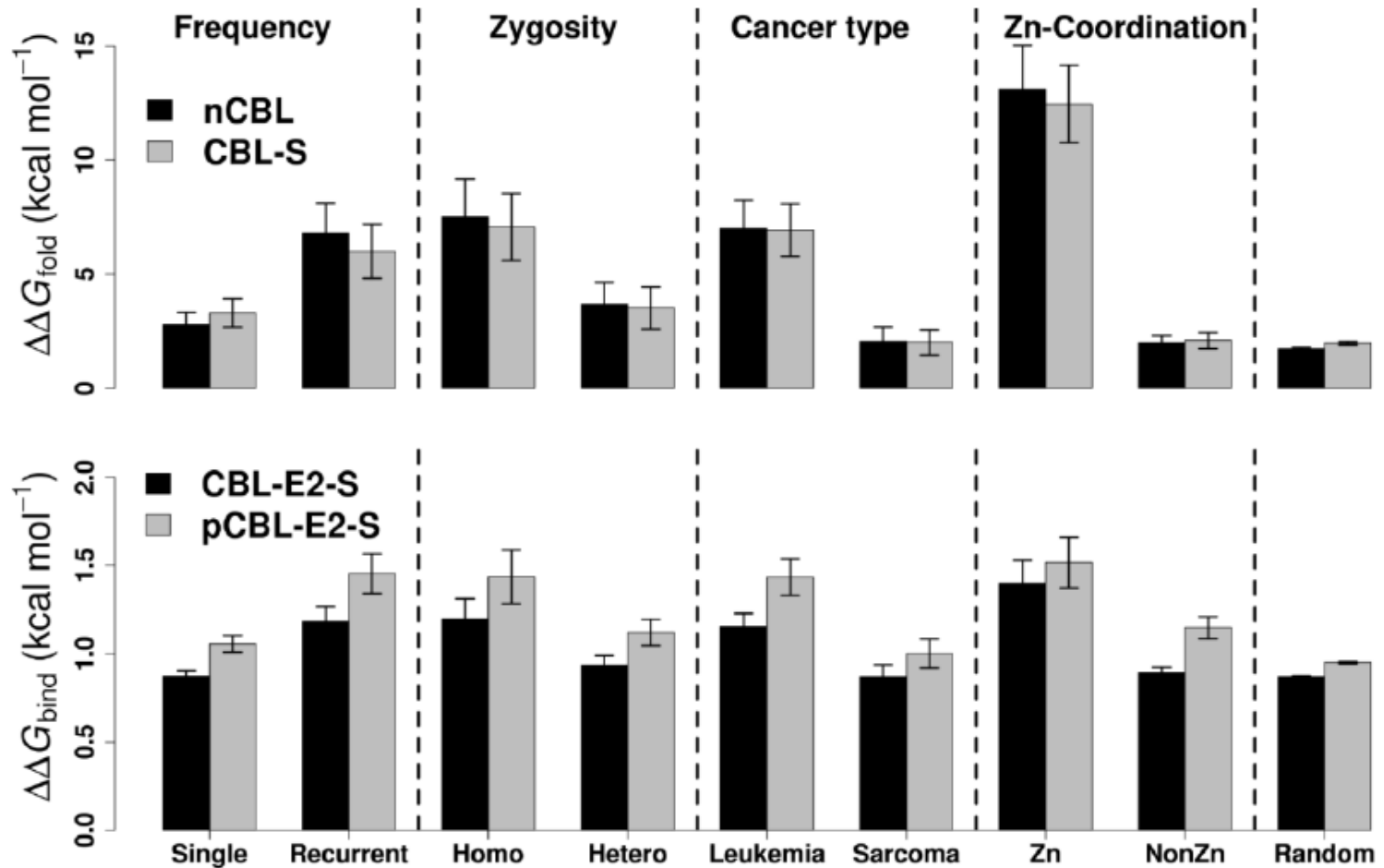


# Cancer mutations impact CBL stability and CBL-E2 binding

~110 cancer and 2100 random missense mutations



# Homozygous mutations and mutations found in Zn-clusters and leukemia patients have largest effects

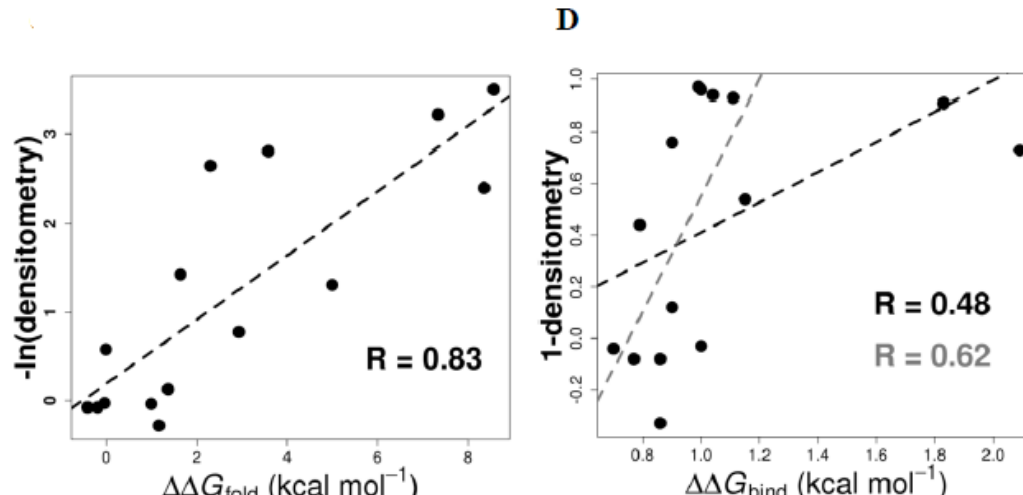
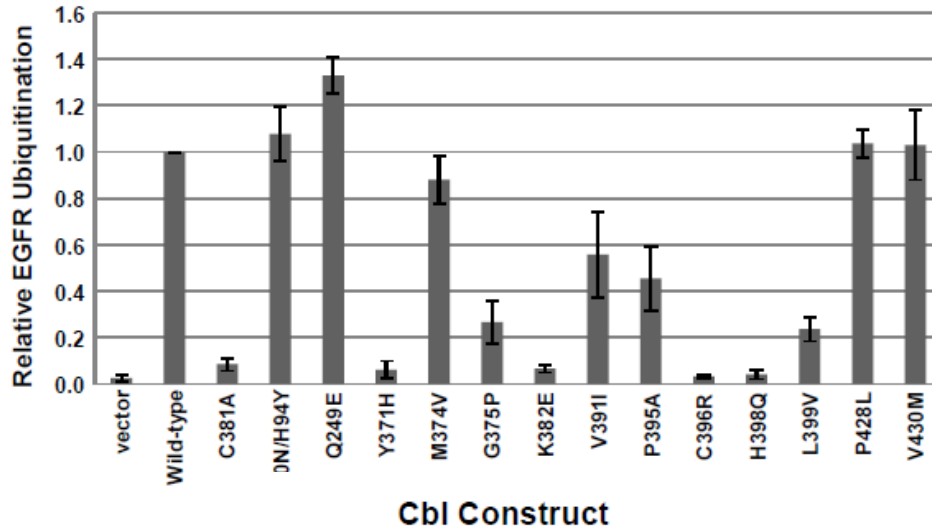




# Comparing experiments with computational models

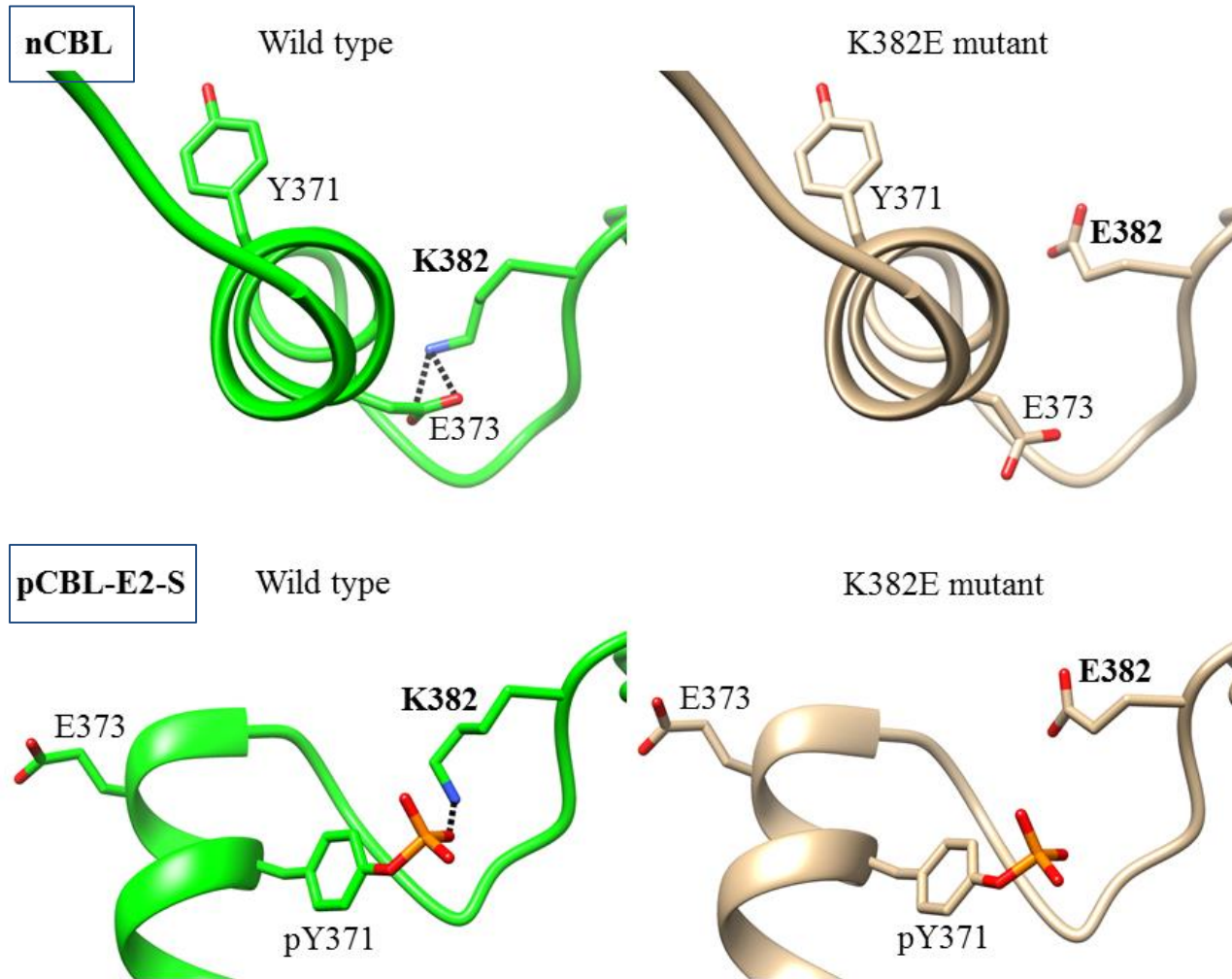
Mutations	Densitometry	Stability		Binding affinity		Methods to predict phenotypic effects			
		nCBL	CBL-S	CBL-E2-S	pCBL-E2-S				
<b>C396R</b>	<b>0.03±0.007</b>	<b>4.65</b>	<b>8.57</b>	<b>0.98</b>	<b>0.99</b>	<b>-11.39</b>	<b>0.99</b>	<b>4.69</b>	<b>0.88</b>
<b>H398Q</b>	<b>0.04±0.019</b>	<b>7.28</b>	<b>7.34</b>	0.79	<b>1.00</b>	<b>-7.57</b>	<b>1</b>	<b>4.34</b>	<b>0.73</b>
<b>Y371H</b>	<b>0.06±0.038</b>	<b>3.58</b>	<b>3.43</b>	<b>0.98</b>	<b>1.04</b>	-4.70	<b>1</b>	<b>2.35</b>	<b>0.92</b>
<b>K382E+</b>	<b>0.07±0.016</b>	<b>2.30</b>	0.56	0.82	<b>1.11</b>	-3.74	<b>1</b>	<b>2.56</b>	<b>0.77</b>
<b>C381A</b>	<b>0.09±0.028</b>	<b>8.36</b>	<b>7.55</b>	<b>2.29</b>	<b>1.83</b>	<b>-8.48</b>	<b>1</b>	<b>4.72</b>	<b>0.59</b>
<i>L399V</i>	<i>0.24±0.051</i>	1.64	0.22	0.80	0.90	-2.83	<b>1</b>	1.70	0.34
<i>G375P+</i>	<i>0.27±0.094</i>	-0.19	<b>5.00</b>	0.67	<b>2.09</b>	<b>-7.56</b>	<b>1</b>	<b>2.09</b>	<b>0.94</b>
<i>P395A</i>	<i>0.46±0.138</i>	<b>1.97</b>	<b>2.93</b>	0.66	<b>1.15</b>	<b>-7.54</b>	<b>1</b>	<b>2.54</b>	0.34
<i>V391I</i>	<i>0.56±0.185</i>	-0.14	-0.01	0.81	0.79	-0.37	0.13	0.45	<b>0.78</b>
<b>M374V+</b>	<b>0.88±0.104</b>	1.36	0.87	0.81	0.90	-3.56	0.83	<b>2.28</b>	0.35
<b>V430M</b>	<b>1.03±0.150</b>	-0.04	-1.15	0.87	<b>1.00</b>	-2.19	<b>1</b>	<b>2.16</b>	0.35
<b>P428L</b>	<b>1.04±0.059</b>	0.80	0.99	0.85	0.70	-4.29	<b>0.98</b>	<b>2.13</b>	<b>0.71</b>
<b>S80N</b>	<b>1.08±0.115</b>	-0.42	-0.5	0.71	0.86	-2.69	<b>1</b>	<b>2.56</b>	<b>0.84</b>
<b>H94Y</b>	<b>1.08±0.115</b>	-1.00	-0.2	0.80	0.77	-4.26	<b>1</b>	<b>2.22</b>	<b>0.78</b>
<b>Q249E</b>	<b>1.33±0.077</b>	0.88	1.16	0.79	0.86	-2.81	<b>1</b>	<b>2.78</b>	<b>0.84</b>
<b>Cutoff</b>		1.80	2.04	0.87	0.95	-4.75	0.87	2.07	0.49

# Stability-activity tradeoff



$$\frac{D_{\text{mut}}}{D_{\text{WT}}} \sim e^{-\Delta\Delta G_{\text{fold}}}$$

# K382E mutations significantly destabilizes the closed and active CBL states



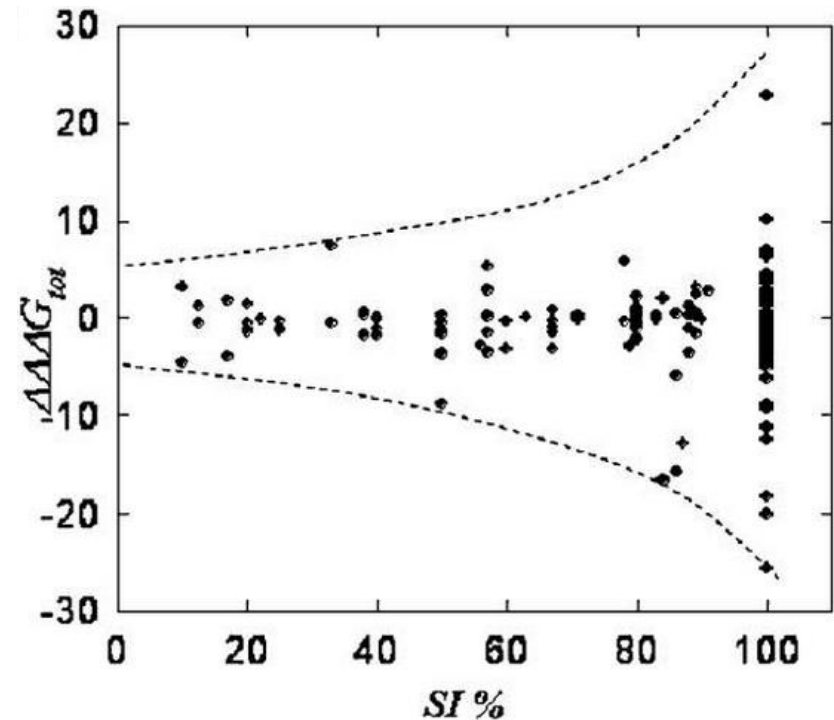
# Effect of OMIM nsSNPs on protein complex stability

Mean values of  $\Delta\Delta\Delta G$  distributions

	$\Delta\Delta\Delta G(\text{total})$ kcal/mol	$\Delta\Delta\Delta G(\text{van der}$ $\text{Waals})$ kcal/mol	$G(\text{electrostatic})$ kcal/mol
OMIM	-1.65	-1.03	-2.35
Non-OMIM	-0.70	0.14	-0.45

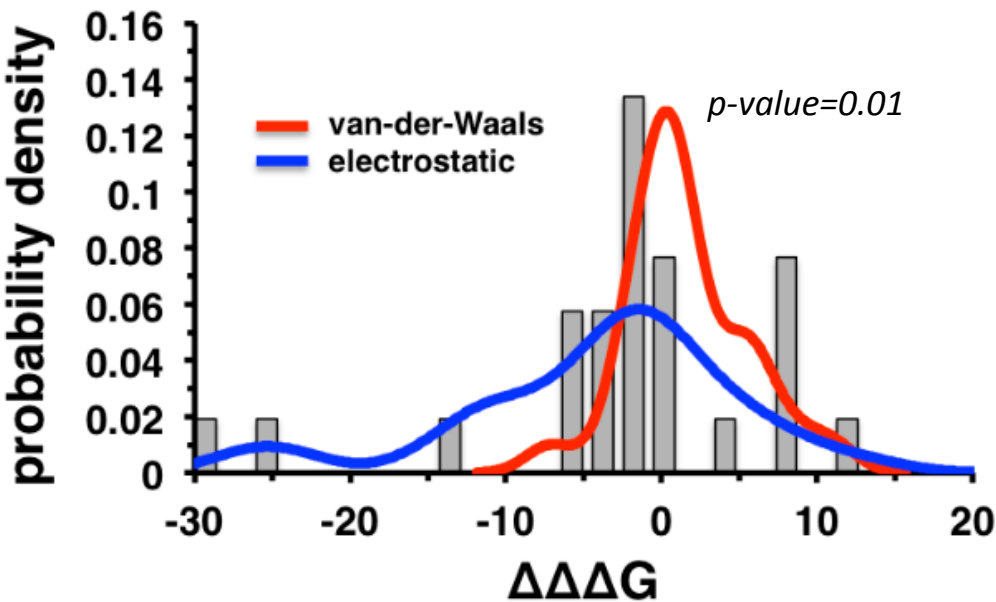
p-value > 0.01 (for total and van der Waals components)  
p-value = 0.006 (for electrostatic component)

- OMIM mutations destabilize electrostatic components of binding energy;
- Largest effect of mutations is observed at evolutionary conserved sites.

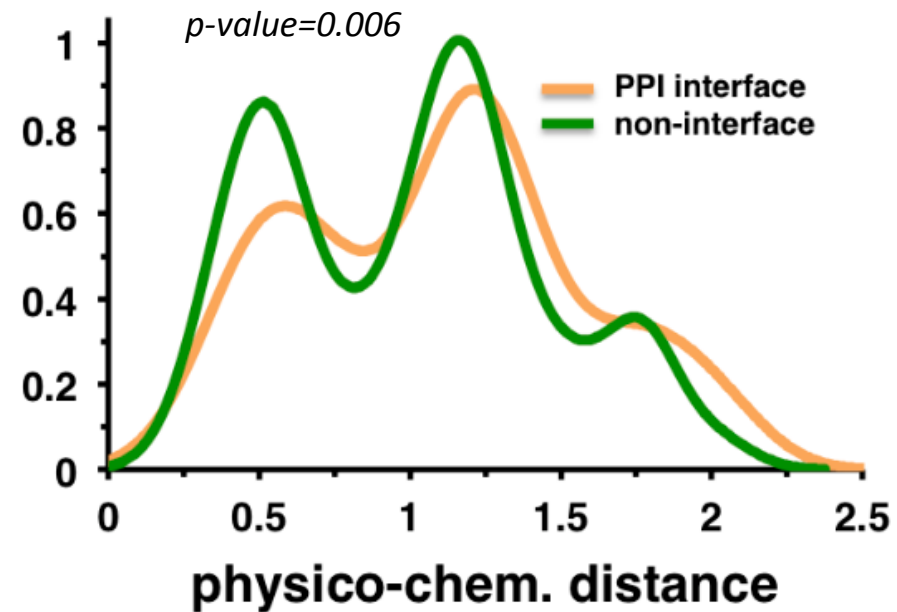


# Effect of glioblastoma mutations on protein binding

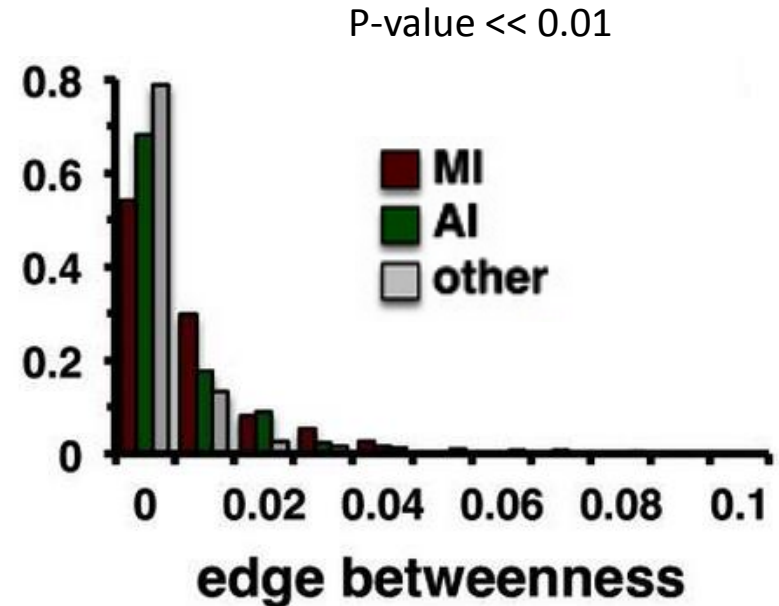
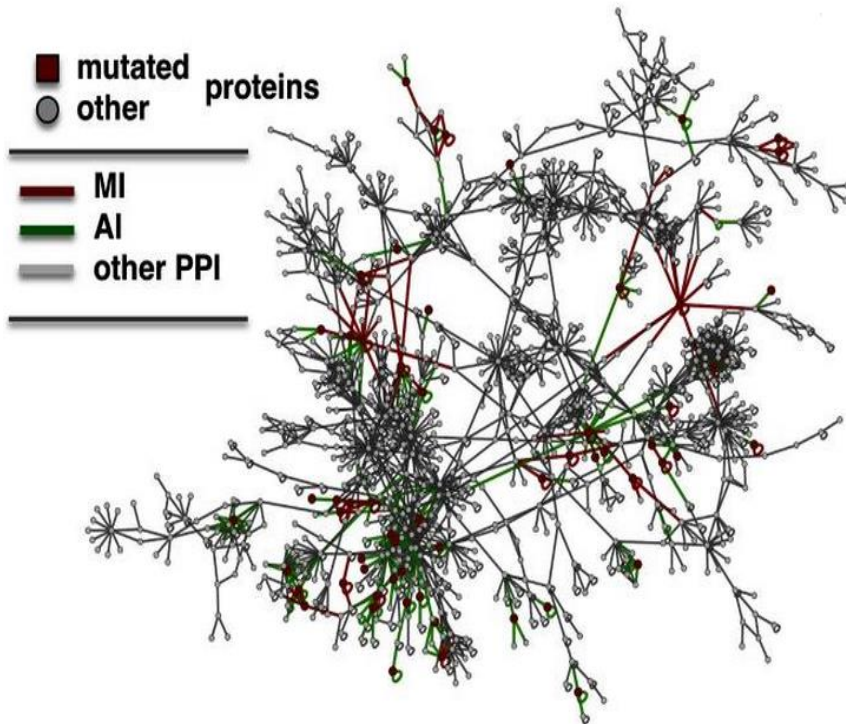
Binding energy difference upon mutation for electrostatic and van-der-Waals components



Physico-chemical distances between mutations on protein-protein interfaces and non-interface regions



# Topological properties of mutated gene network



number of shortest paths going through a node

**AI** – 444 interactions between proteins with mutation anywhere in protein

**MI** – 160 interactions between proteins with mutation on interface

Interactions with mutations occur in central network positions!

# Predicted driver mutations

## Protein-protein

ABL2  
ARL1  
EPHA2  
IDH1  
NLGN2  
NRAS  
RAB3C  
RAC2  
RAD52  
TP53

## Protein-DNA

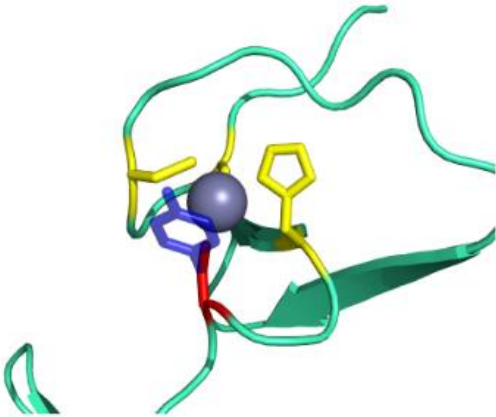
BCL11A  
PAX9  
TP53  
ZIK1  
ZNF339

## Protein-RNA

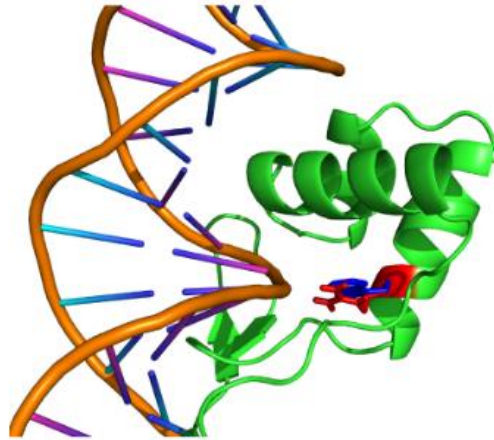
ELAVL2  
KLK9  
RBMS3  
RPL11

## Protein-ion

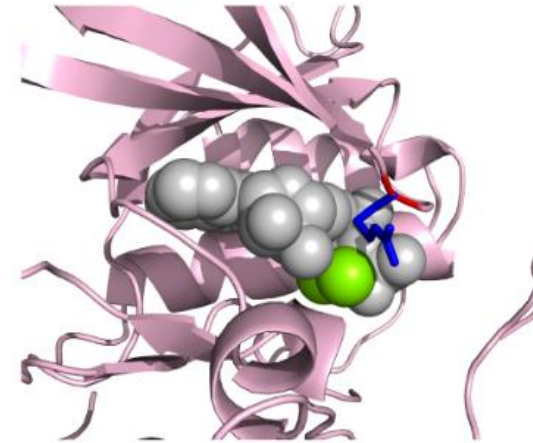
ADAMTS17  
DSG4  
GZMH  
HPCAL4  
LCT  
LMX1A  
MAPK9  
NELL2  
SGK2  
TP53  
ZIK1  
ZNF497



Zinc binding motif of LMO-2 (homolog of LMX1A), C→Y

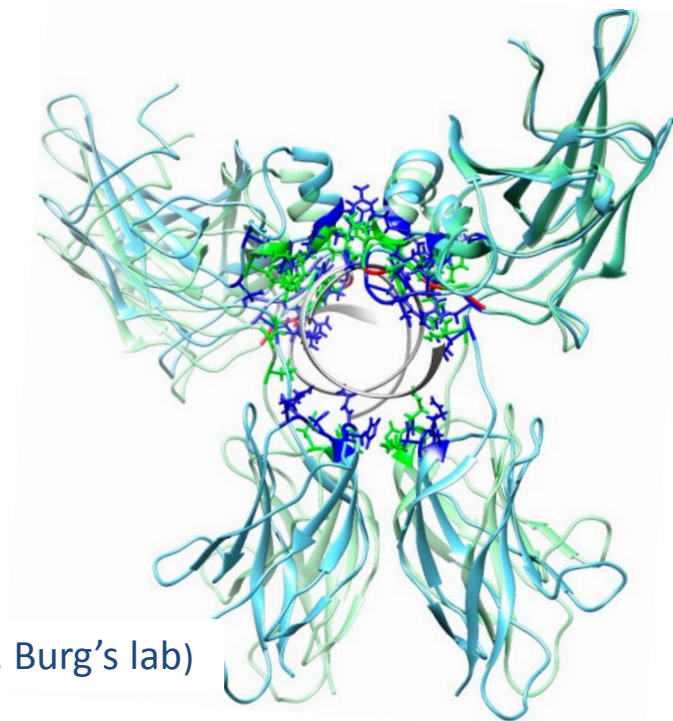
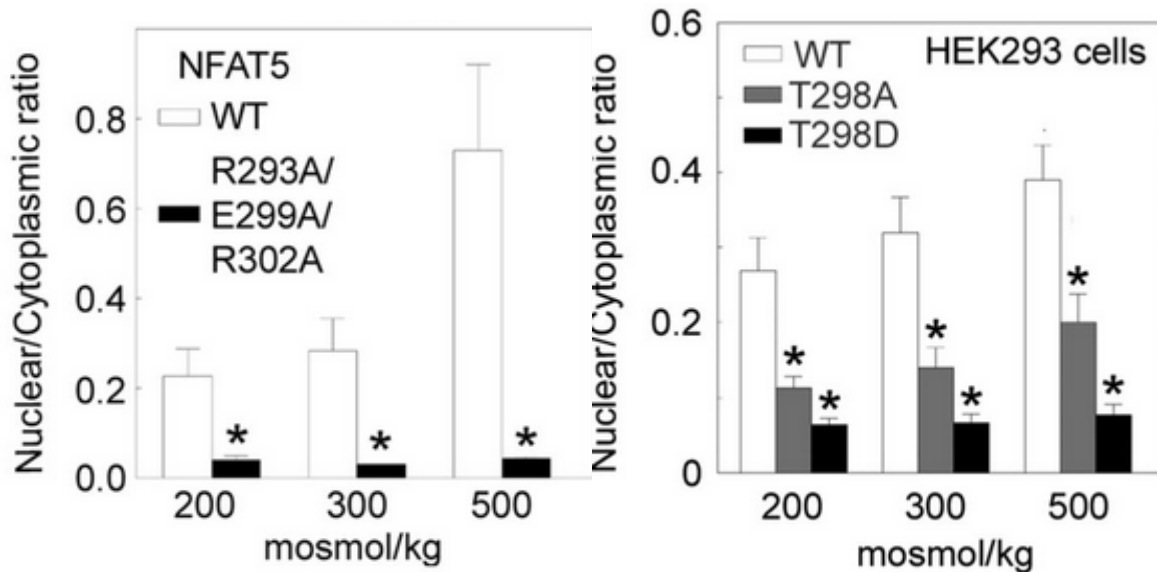


DNA binding site of Pax-6, R→W



Protein-ion binding site of MAPK10, G→R

# Mutations in DNA-binding loop of NFAT5 produce unique outcomes on binding and dynamics



Izumi et al, *Am J Physiol Cell Physiol*. 2012 (experimental data from M. Burg's lab)

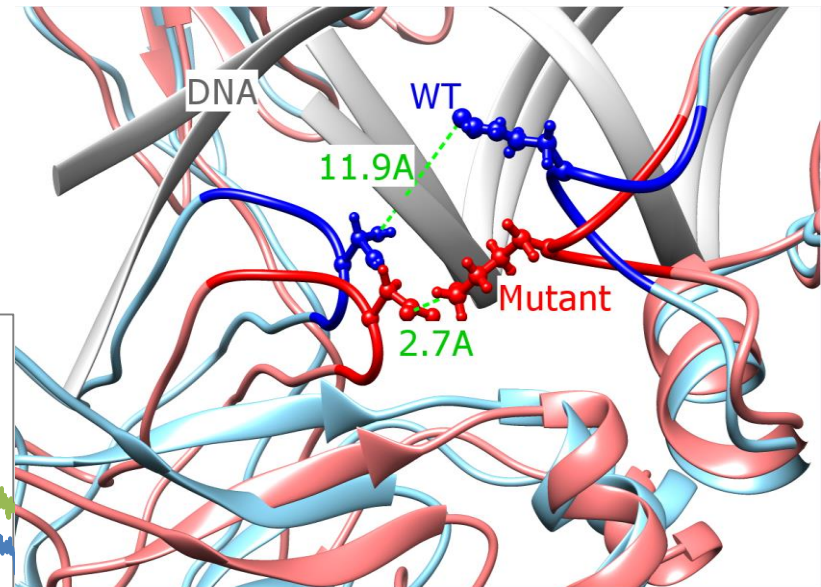
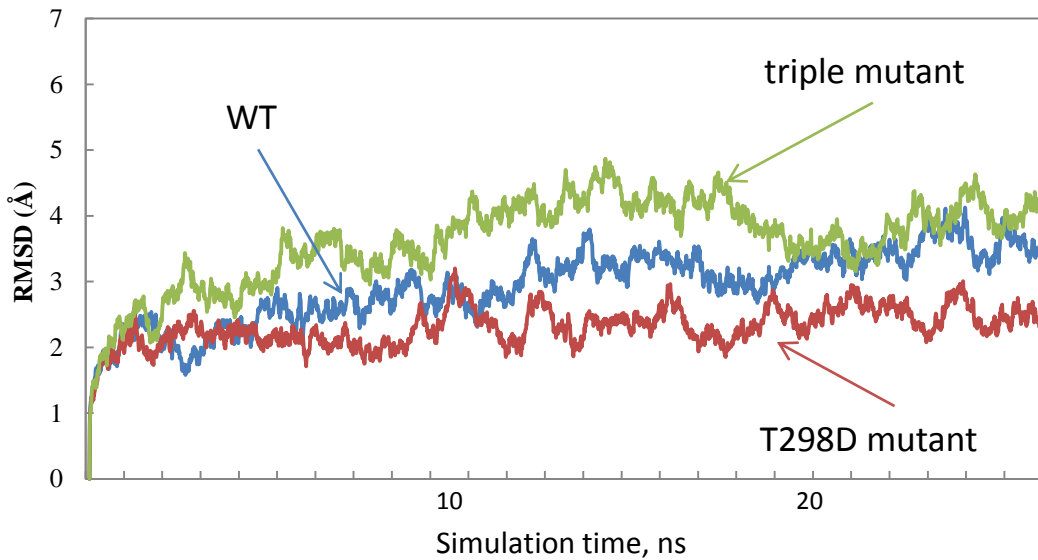
Binding energy, kcal/mol

	Complex - DNA	Chain C - DNA	Chain D - DNA	Chain C – Chain D
Native	10.00 (0.62)	3.61 (0.39)	2.89 (0.39)	2.73 (0.23)
T298D	9.24 (0.54)	3.79 (0.23)	2.76 (0.23)	2.92 (0.23)
R293, E299, R302	6.47 (0.62)	3.13 (0.23)	2.36 (0.23)	2.78 (0.31)



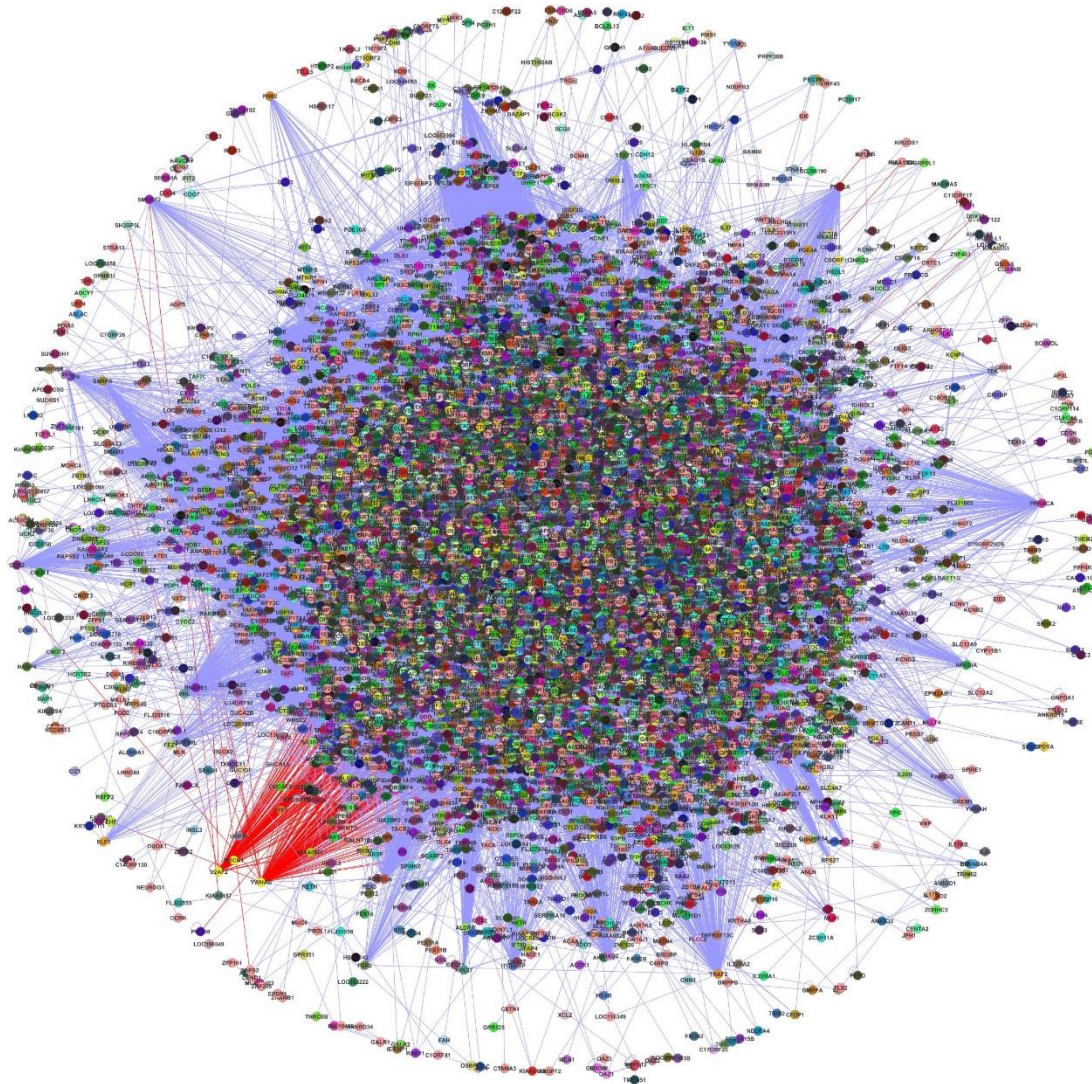
# Phosphomimetic mutation T298D constraints movements of two chains

Changes in conformation, Molecular Dynamics simulations (NAMD program, explicit solvent)



Effect of T298D on structure, formation of an extra salt bridge between two chains in a dimer

# Reality: PPI Interactomes



All hairball graphs look alike

# Protein interactions in the cell

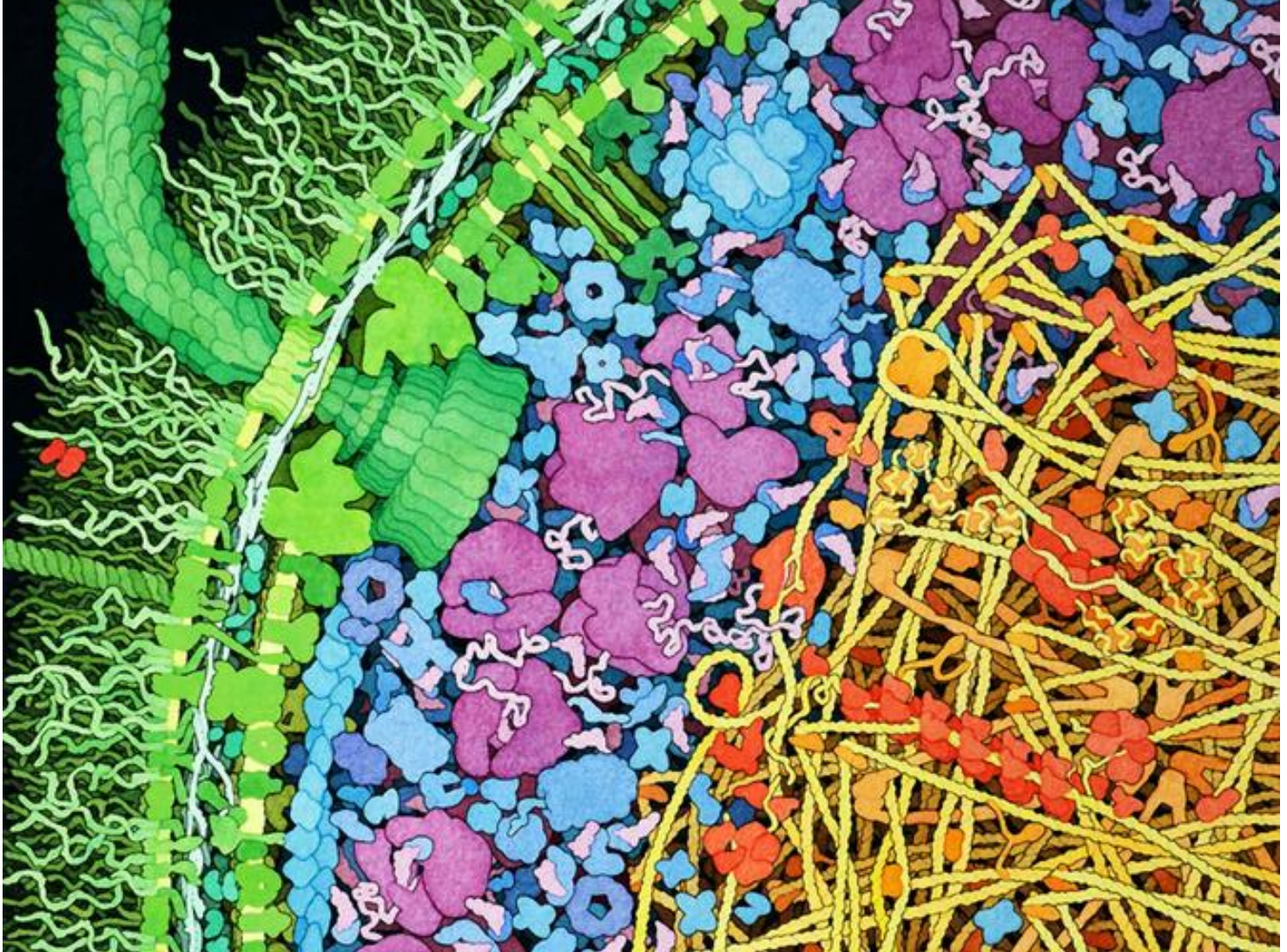
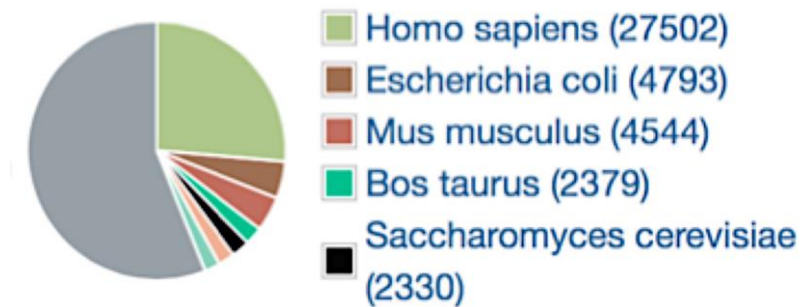
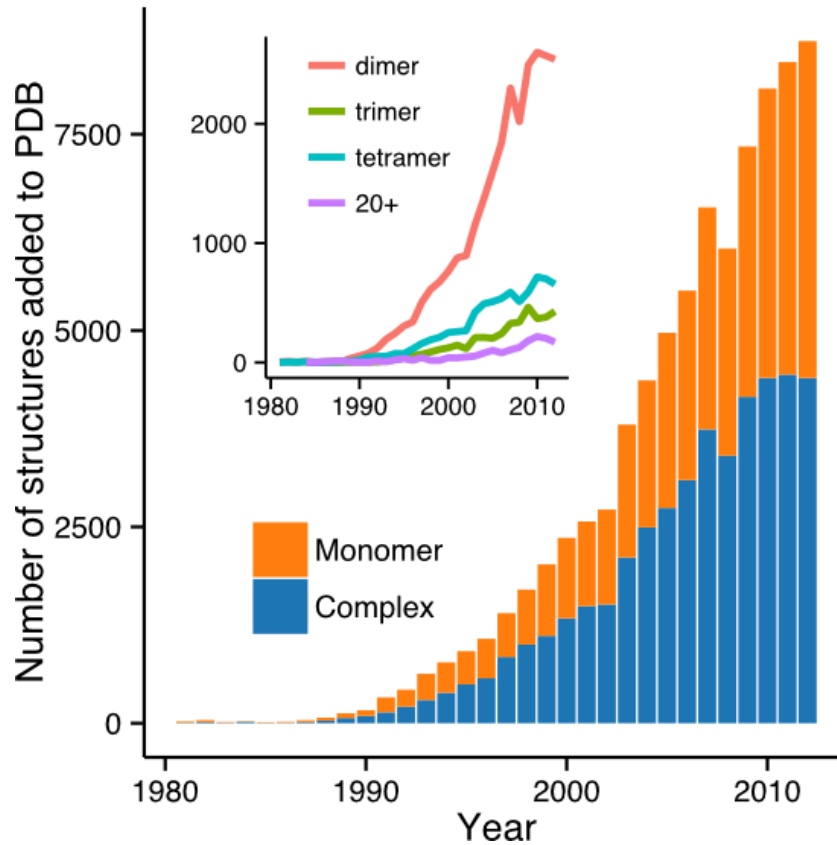


Illustration by David S Goodsell

# Structural interactomes are informative and useful

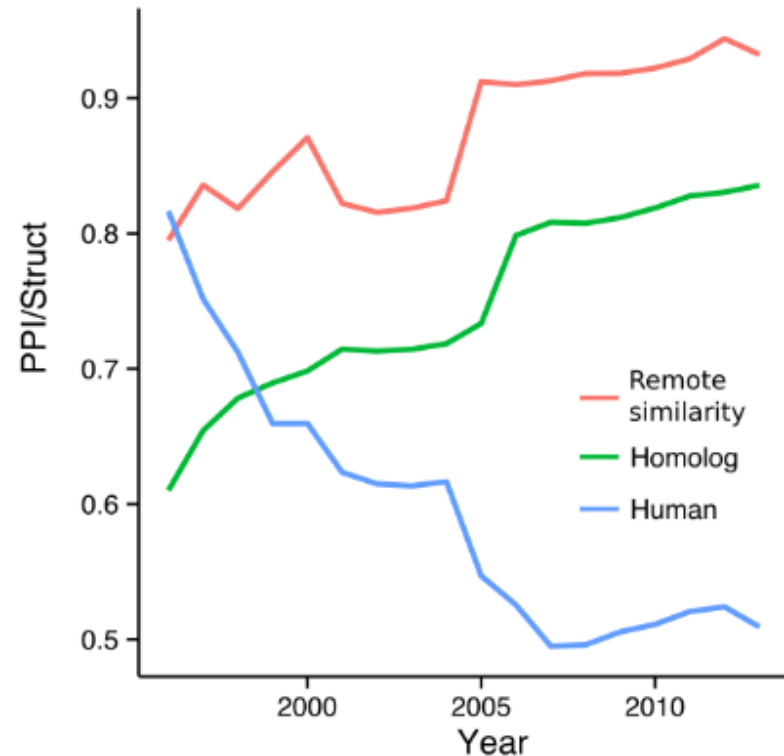
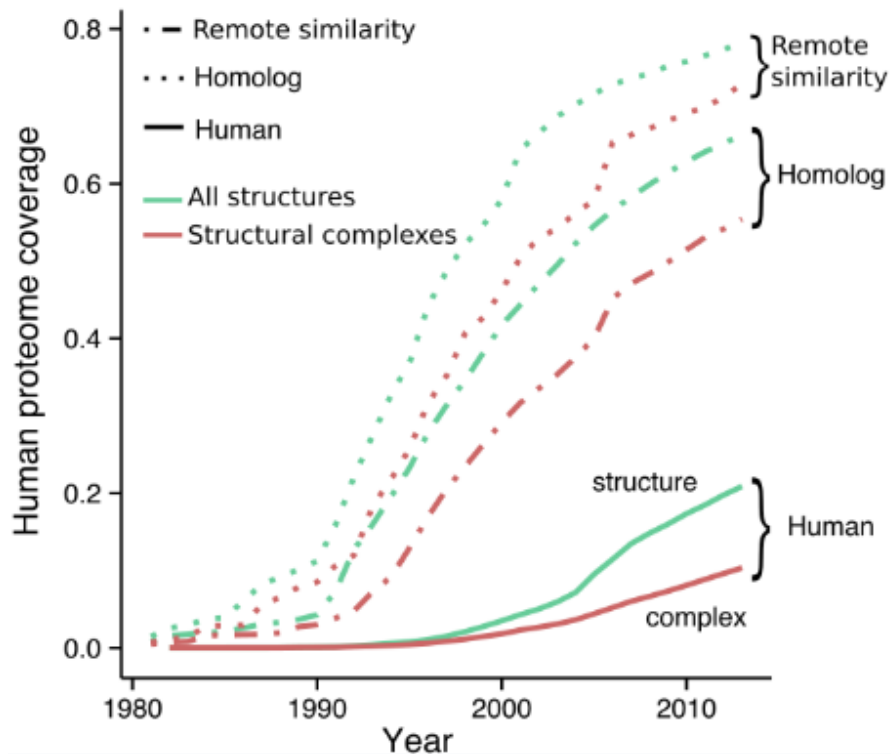
- Interactome with structural details:
  - Which proteins interact?
  - How they interact:
    - Which domains interact?
    - Which residues form binding sites?
- Atomic-resolution interfaces are needed to study:
  - The mechanisms of interactions.
  - The effects of mutations on stability of proteins and their complexes.
  - To modulate interactions (drugs)
- Strategies:
  - Use available structures of protein complexes.
  - Dock structural monomers if structural complex is not available.
  - Template-based modeling of protein complexes (or interfaces).

# What do we have so far? Growth of structural and PPI data



# Structural complexes are available for less than 10% of human protein-coding genes!

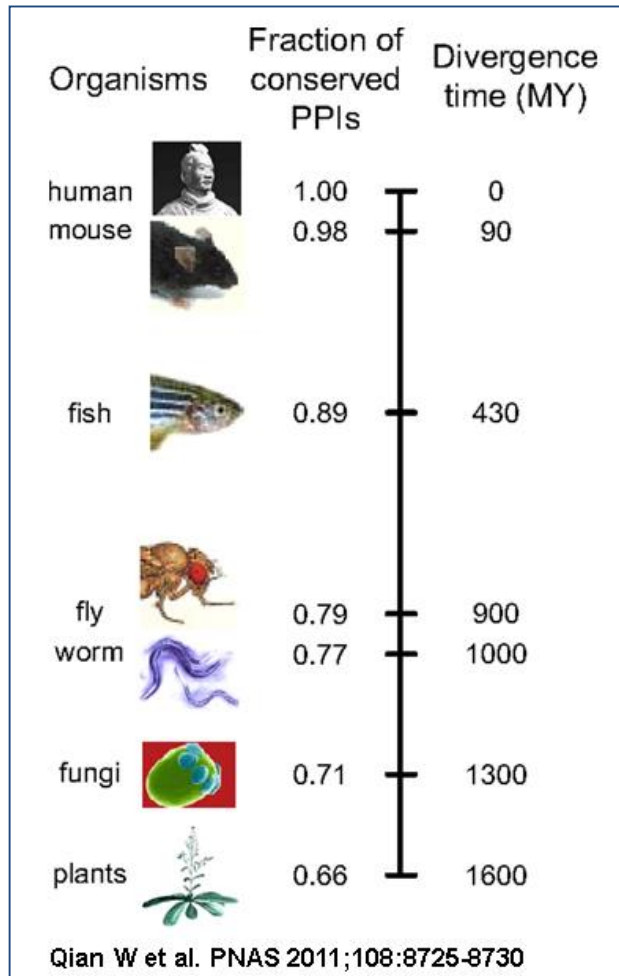
- Two-hybrid assay - 14,000 interactions between human proteins



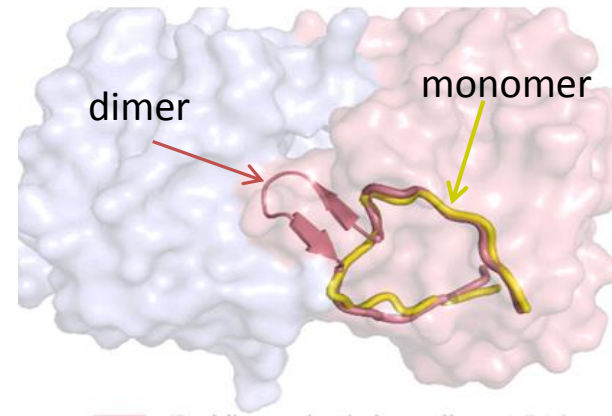
# Conservation of protein interactions and oligomeric states

## Conservation of interactions partners

Rate of PPI evolution =  $(2.6 \pm 1.6) \times 10^{-10}$   
per PPI per year

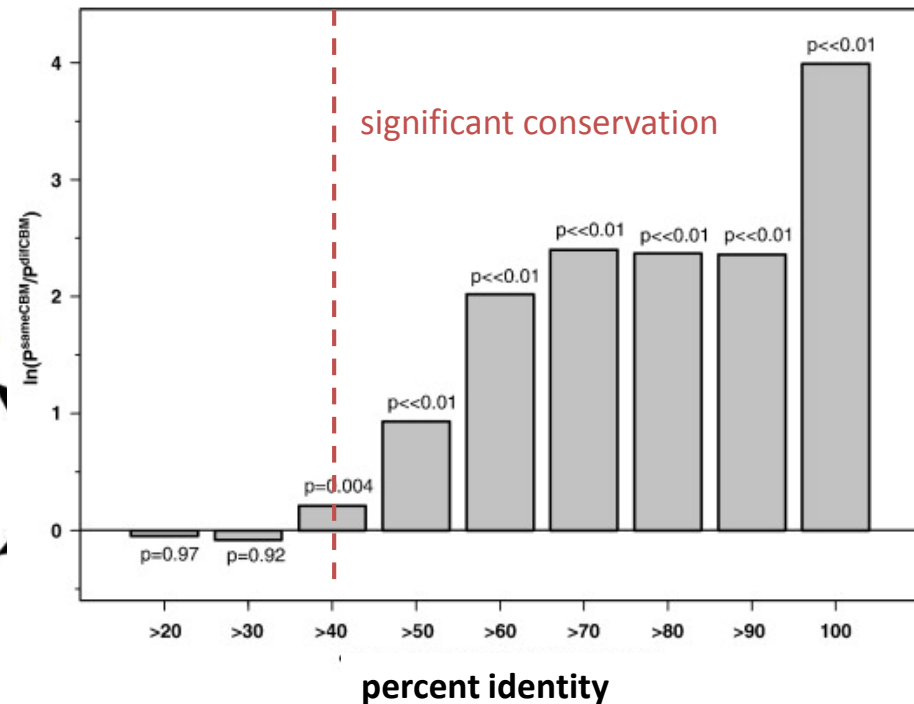
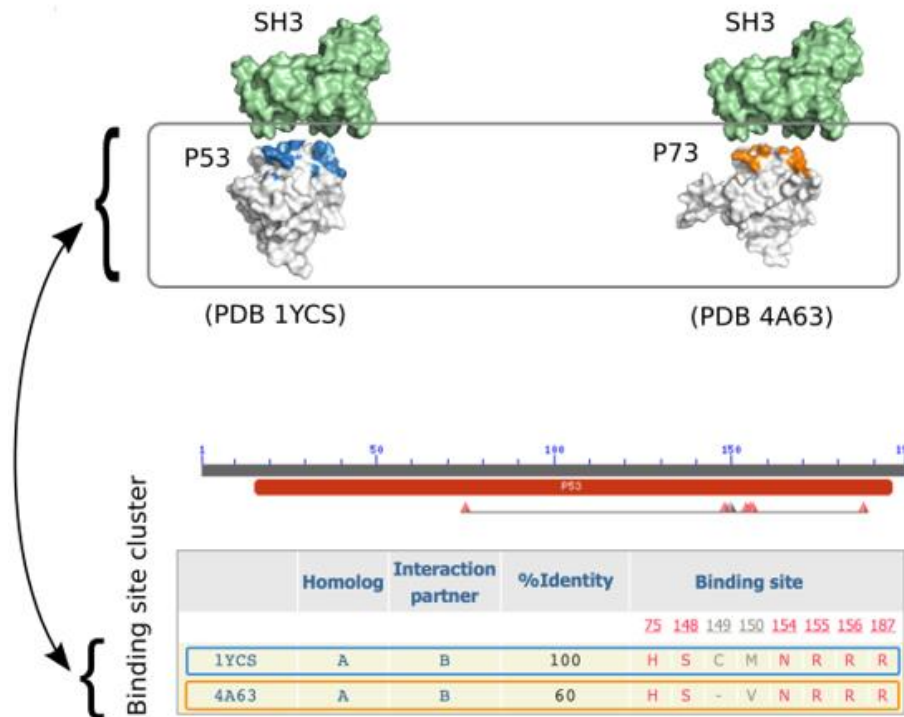


## Conservation of oligomeric states



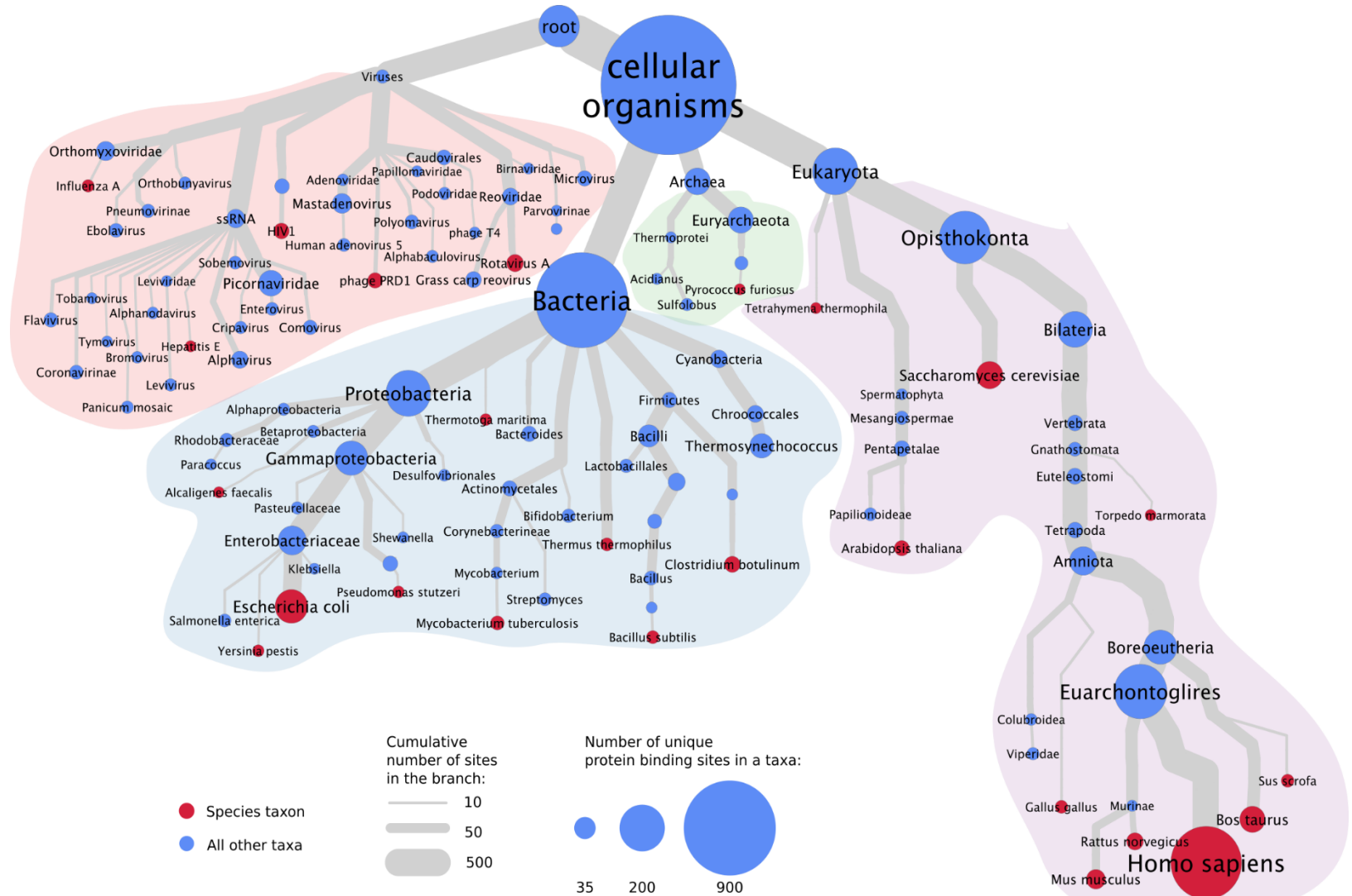
	Sensitivity TP/(TP + FN)	Specificity TN/(FP + TN)
Presence/absence of enabling and disabling features	0.70	0.74
Percent identity	0.71	0.62
RMSD	0.72	0.60

# Different protein complexes might have similar binding interfaces



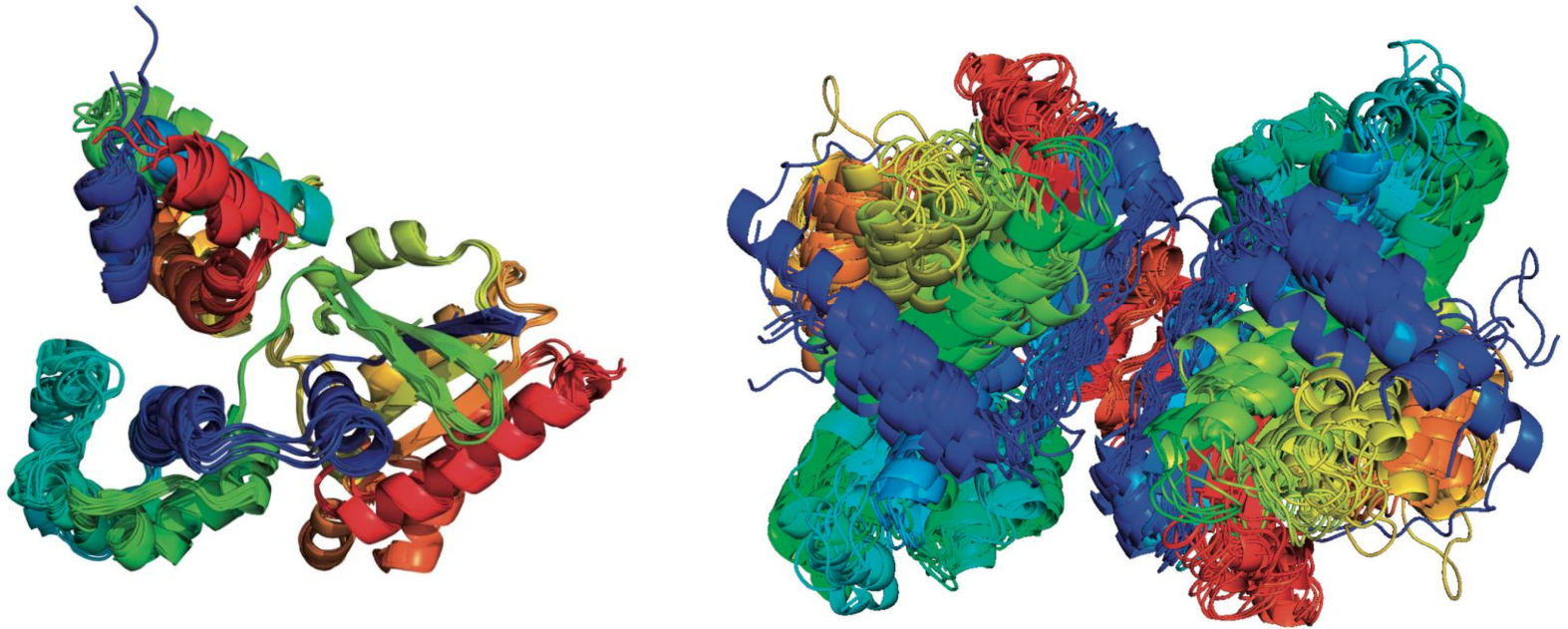


# Tracing back evolution of protein binding sites to the root of all organisms



# The underlying interolog hypothesis

- If proteins are similar they may interact in a similar way
- Homologs may have similar interfaces



# IBIS – NCBI server to analyze interactions and binding sites

<http://www.ncbi.nlm.nih.gov/Structure/ibis/ibis.cgi>

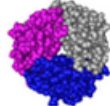
Observed interactions – from structural complexes

Inferred interactions – from homologous structures with observed interactions

Types of interactions: protein-protein, protein-nucleic acids, protein-small molecule, protein-peptide, protein-ion

Biological relevance of binding sites:

- occurs in several non-redundant homologs;
- structurally and sequence conserved;
- validated biounit

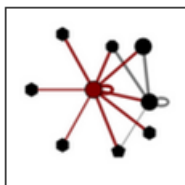


Protein PDB ID, Accession or GI

**Protein-Protein (8)** **Protein-Chemical (4)** **Protein-Peptide (1)** **Protein-Ion (1)**

### Query 1YCR\_A

Mdm2



All interactions for query sequence

EXCEL  XML

### Search 1YCR A interactions

Similarity to query

Sequence Identity:

Structure RMSD:

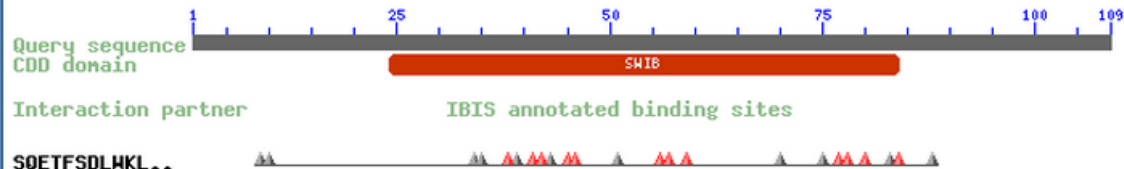
Interaction partner type

PDB Code:

Taxonomy:

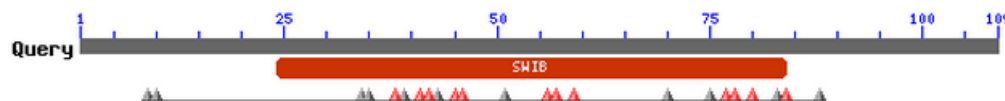
### Mdm2 (sequence A)

[1YCR sequence B](#)



List of peptide interaction partners and binding sites. Similar binding sites of homologs of the query are grouped into clusters. To view the cluster members click on the plus sign. "o" denoted observed interactios.

Interaction Partner	Ranking Score	Number of Cluster Members	Average %Identity to Query	Number of Binding Site Residues	Taxonomic Diversity
<input type="checkbox"/> <input checked="" type="radio"/> SQETFSDLWKLLEN	n/a	24	83	23	Euteleostomi



Interaction Partner	%Identity to query	Binding site
-	9 10 34 35 38 39 41 42 43 45 46 51 56 57 59 70 75 77 78 80 83 84 88	E T M K L F L G Q I M Y Q H V F F V K H I Y Y
100	E T M K L - L G - I M - Q H V - - V K H - Y Y	
73	- - - - I - L G - I M Y Q - - - - V K P L Y -	
58	- - V K M - L G - I M Y Q H V - - V - - L Y -	
58	- - - - M - L G - I M Y Q H V - - V - - - Y -	

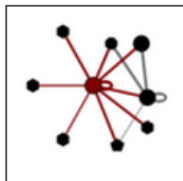
[See all members](#)

\* click structure accession or sequence letter to explore structure and sequence information.



### Query 1YCR\_A

Mdm2



All interactions for query sequence

Download data

EXCEL XML

#### Search 1YCR A interactions

##### Similarity to query

Sequence Identity:

Structure RMSD:

##### Interaction partner type

- Chemical: [Compounds list](#)
- PDB Code: [Compounds list](#)
- Taxonomy

##### Clusters with

Show:

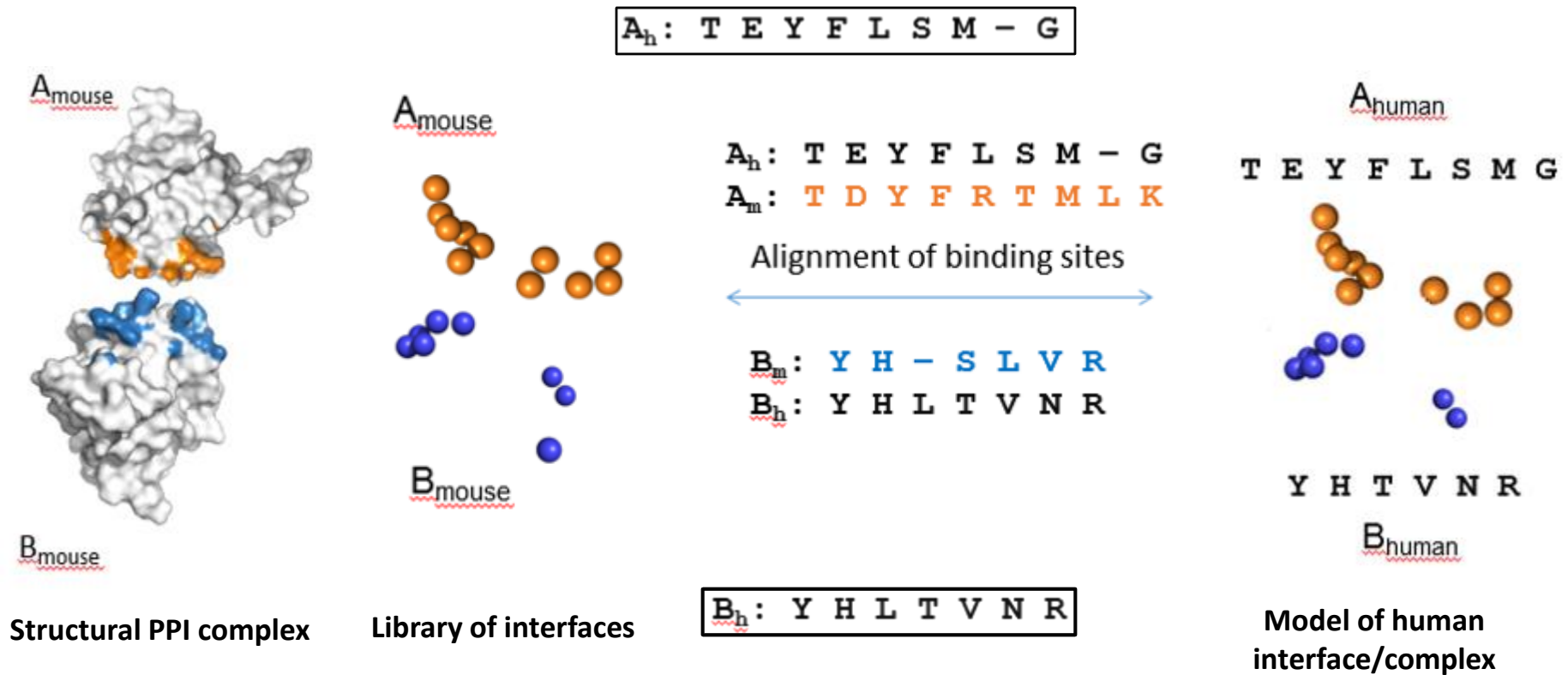
##### Referenc

- Shoem integrates AC1L9LXV
- Shoem predict pro AC1L9MCG
- CHEMBL2024323
- CHEMBL2059435
- CHEMBL2177187
- CHEMBL2177183

	Chemical	Homologous complex	Chain	%Identity to query	Number of binding site residues	Binding site	Taxonomic Diversity
Query	-	-	-	-	-	34 38 39 41 42 43 45 46 51 55 56 57 59 66 70 75 77 78 80 83 84 M L F L G Q I M Y Q Q H V L F F V K H I Y	-
<input checked="" type="checkbox"/>	DIZ	1T4E	A	100	12	- L - L G - I M Y - Q - - - F F V - H I Y	Homo sapiens
<input type="checkbox"/>	DIZ	1T4E	B	100	13	- L - L G - I M Y - Q - - - F F V - H I Y	Homo sapiens
<input type="checkbox"/>	MI6	3LBL	A	100	11	- L - L G - I M - - - - - F F V - H I Y	Homo sapiens
<input type="checkbox"/>	CHEMBL2177187	4ERE	A	100	11	- L F - G - I - Y - - - - F - V K H I Y	Homo sapiens
<input type="checkbox"/>	CHEMBL2059435	4ERF	A	100	11	- L - L G - I M Y - - - - - - - V K H I Y	Homo sapiens
<input type="checkbox"/>	SureCN9993627	4OAS	A	100	11	- L F - G Q I - - - - - F - V K H I Y	Homo sapiens
<input type="checkbox"/>	CHEMBL2347399	4JV7	A	100	12	- L F L G Q I M Y - Q - - - - F V - - I -	Homo sapiens
<input type="checkbox"/>	CHEMBL2347401	4JV9	A	100	11	- L F L G Q I M Y - Q - - - - - V - - I -	Homo sapiens
<input type="checkbox"/>	CHEMBL2347383	4JVE	A	100	12	- L F L G Q I M - Q - - - - - V - H I Y	Homo sapiens
	(2's,3r,4's,5'r)-N-(2-						
<input type="checkbox"/>	Aminoethyl)-6-Chloro-4'-(3-	4JVR	A	100	12	- L - L G - I M Y - - - - - F F V - H I Y	Homo sapiens
<input type="checkbox"/>	CHEMBL2347393	4JWR	A	100	10	- L - L G - I M - - - - - F - V - H I Y	Homo sapiens
	3-((1s)-2-(Tert-						
<input type="checkbox"/>	Butylamino)-1-[(4-Chlorobenzyl)(F	3TJ2	A	100	13	- L F - G - I M Y - - - V - F F V - H I Y	Homo sapiens
<input type="checkbox"/>	SureCN9993362	4HBM	A	100	10	- L - L G - I - - - - - F - V K H I Y	Homo sapiens
	3-((1s)-2-(Tert-						
<input type="checkbox"/>	Butylamino)-1-[(4-Chlorobenzyl	4MDN	A	100	12	M L F L G - I M - - - - - F - V - H I Y	Homo sapiens
<input type="checkbox"/>	28W	4MDQ	A	100	10	- L F - G - I - Y - Q - - - - F V - H I -	Homo sapiens
	1-(((5r,6s)-5,6-Bis(4-						
<input type="checkbox"/>	Chlorophenyl)-6-Methyl-3-(P	3VZV	A	99	11	- L - L G - I M Y - Q - - - - - V - H I Y	Homo sapiens
	(5r,6s)-2-(((2s,5r)-2-						
<input type="checkbox"/>	(((3r)-4-Acetyl-3-Methylpip	3W69	A	99	12	- L - L G - I M Y - Q - V - - - - V - H I Y	Homo sapiens
	(5z)-5-[(6-Chloro-7-						
<input type="checkbox"/>	Methyl-1h-Indol-3-	3VBG	A	99	6	- L - L G - I - - - - - - V - - I -	Homo sapiens

gi M, Fong JH, Marchler-Bauer A, Bryant SH, Madej T, Panchenko AR. (2010), *Inferred Biomolecular Interaction Server—a web server to analyze and sites. Nucleic Acids Res. 38(Database issue): D518-24.*

# Modeling of interactions and interfaces

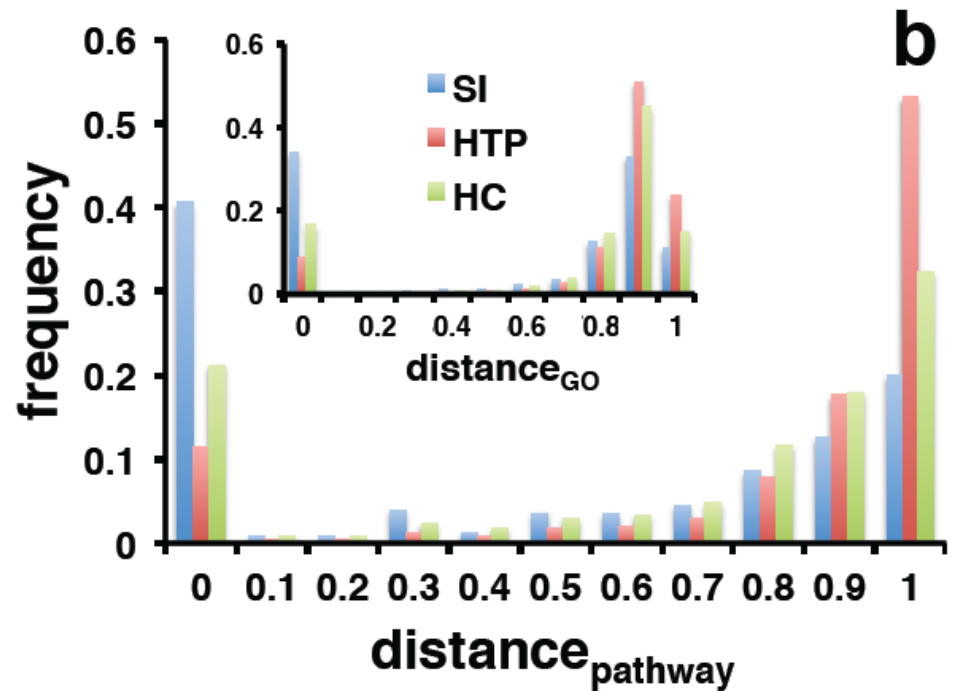
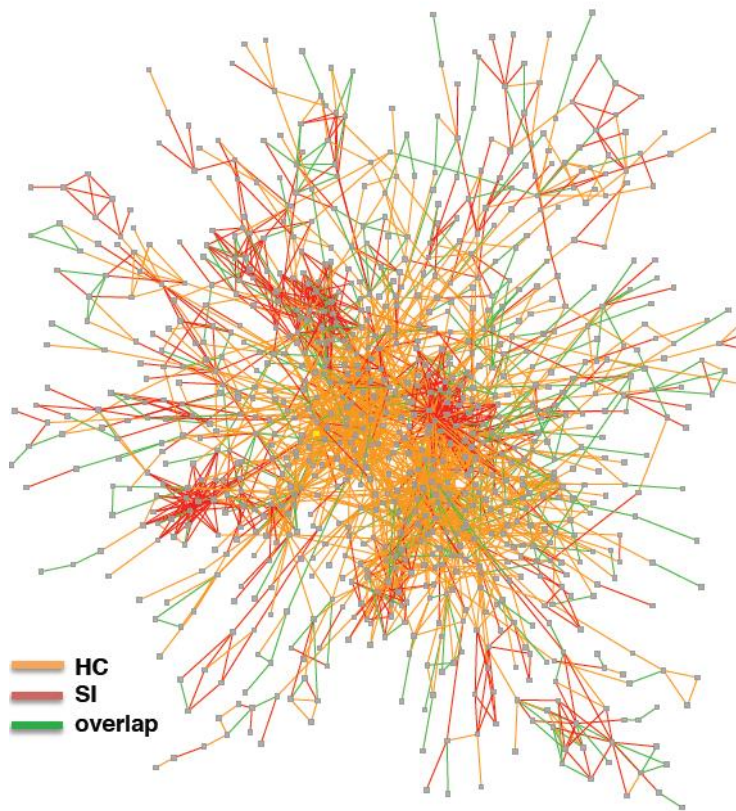


Verification:

- minimization of complex;
- interface complementarity;
- interface conservation;
- co-localization;
- co-expression.

# Mapping of human interactome using structural complexes

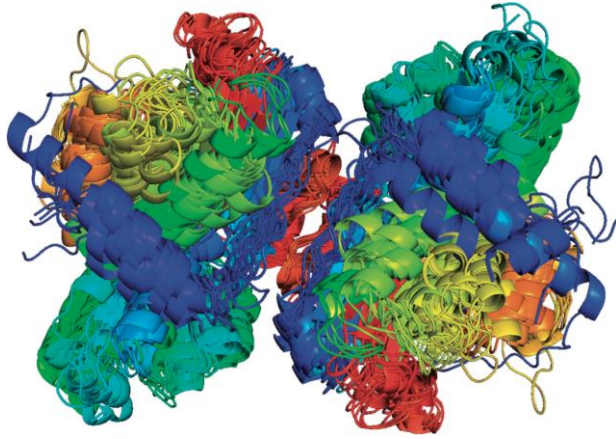
4400 human genes  
20000 interactions



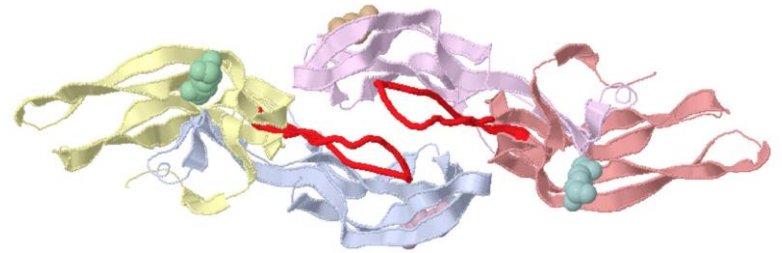
$$\Delta_{ij} = 1 - \frac{|\Gamma_i \cap \Gamma_j|}{|\Gamma_i \cup \Gamma_j|} \quad \Gamma_i - \text{list of attributes for protein } i \text{ (GO or pathway annotations),}$$

Structurally inferred networks (SI) are more functionally coherent than high-throughput networks (HTP, HC)

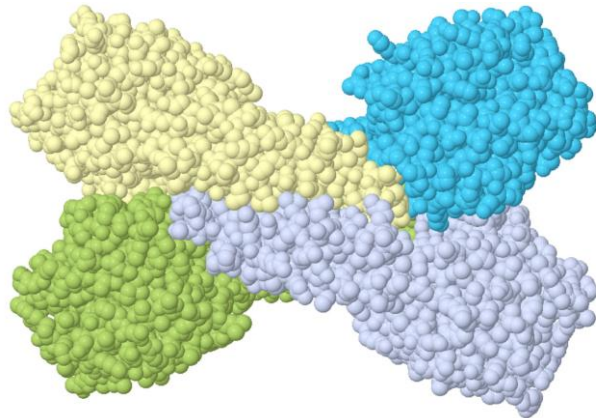
# Challenges in computational analysis and prediction of PPI



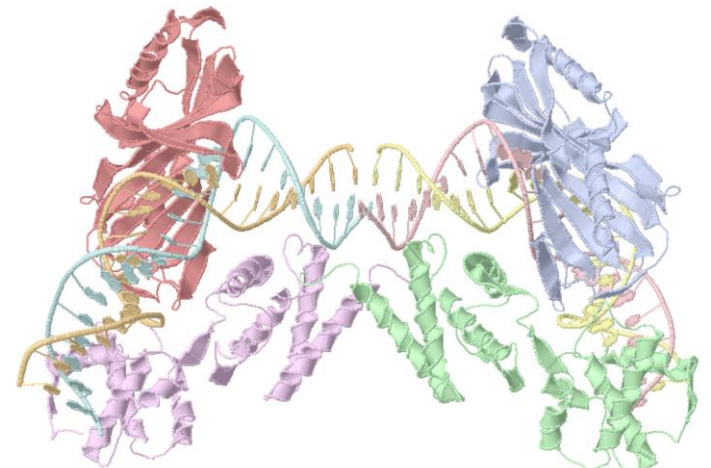
Substantial conformational changes



Highly variable regions



Higher oligomers



Interactions with other molecules



## Acknowledgements

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Kosuke Hashimoto (RIKEN, Japan)

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Stefan Wuchty (*University of Miami, USA*)

Stanley Lipkowitz (*NCI, NIH*)