

Protein Structure Prediction and Analysis Tools

Jianlin Cheng, PhD

Assistant Professor

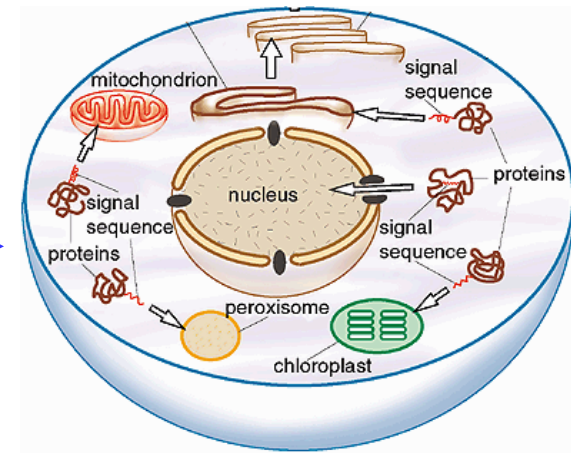
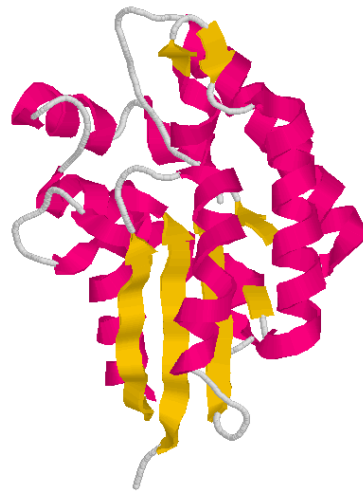
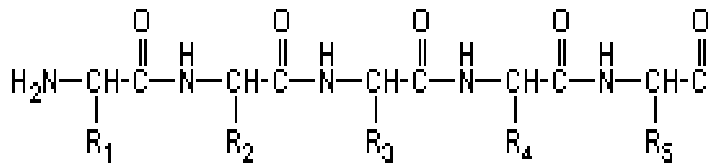
Department of Computer Science & Informatics Institute

University of Missouri, Columbia

2011

Sequence, Structure and Function

AGCWY.....

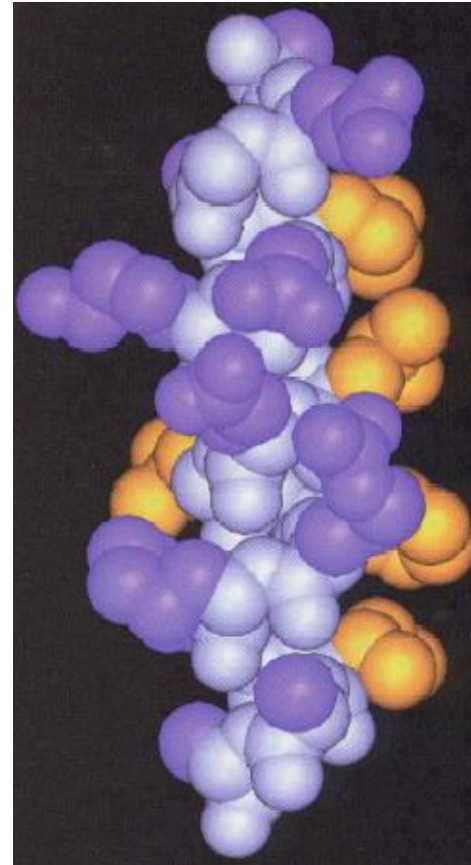
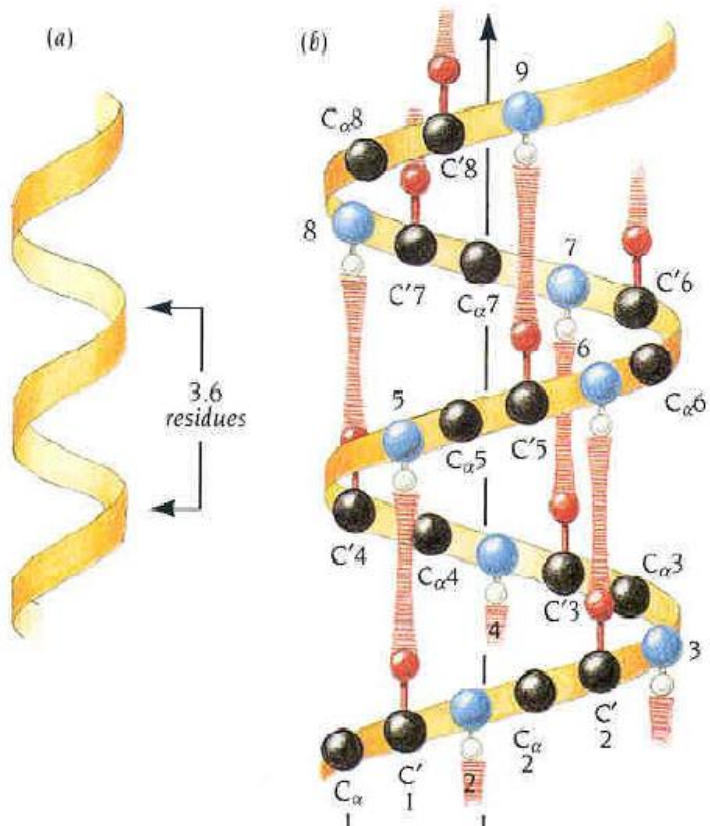


Cell

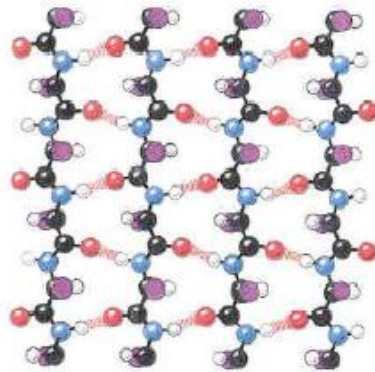
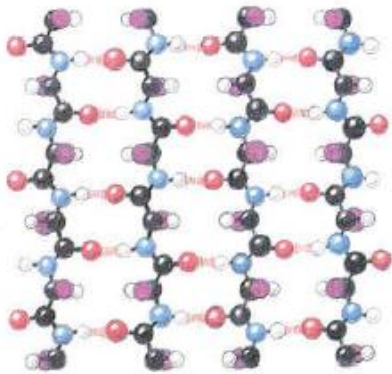
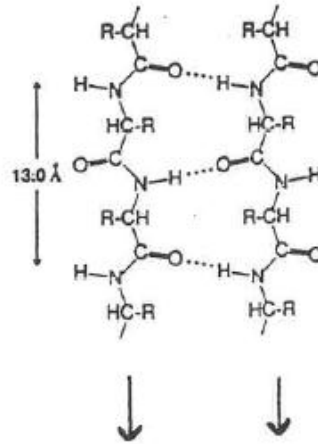
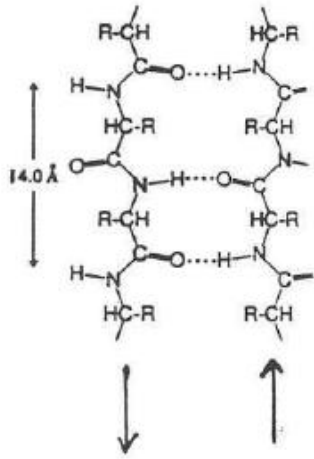
Protein Folding Movie

<http://www.youtube.com/watch?v=fvBO3TqJ6FE&feature=fvw>

Alpha-Helix

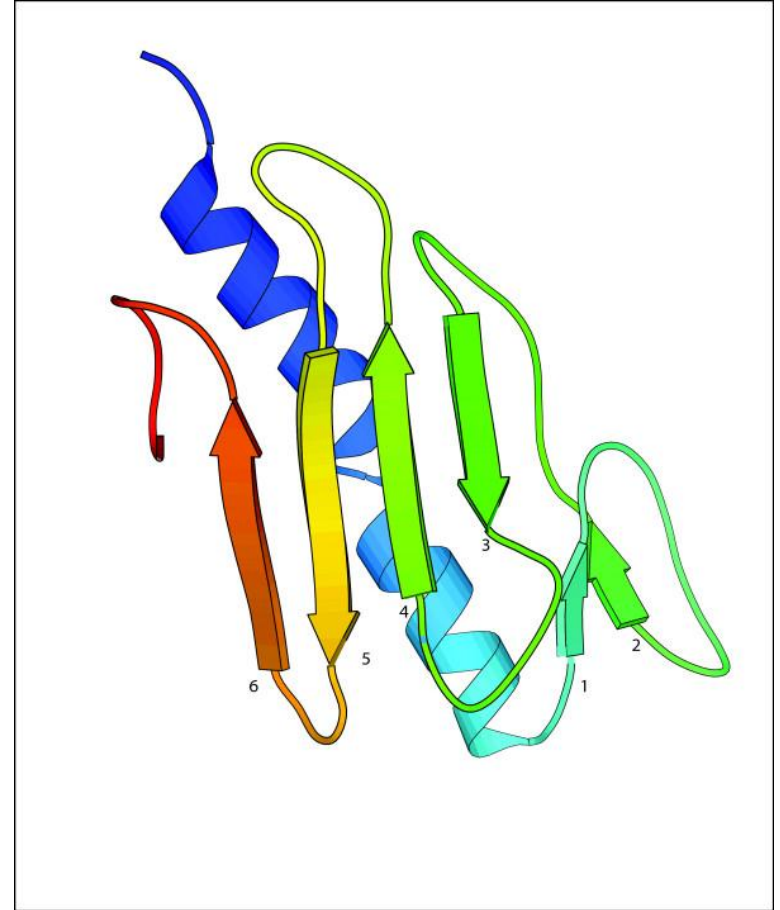


Beta-Sheet

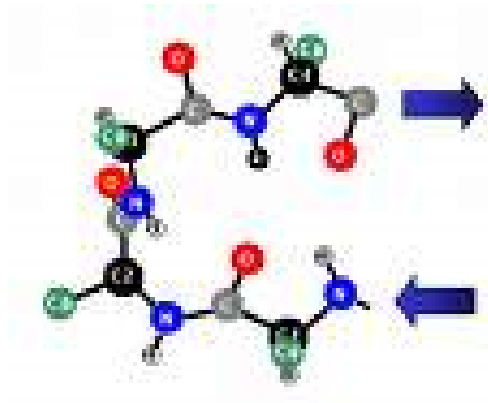


Anti-Parallel

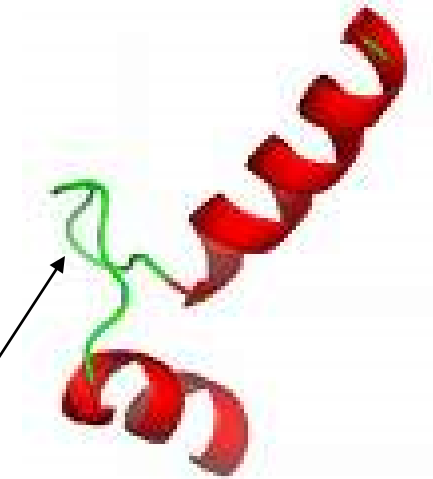
Parallel



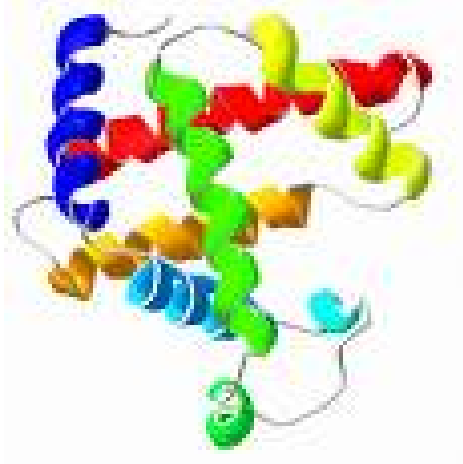
Non-Repetitive Secondary Structure



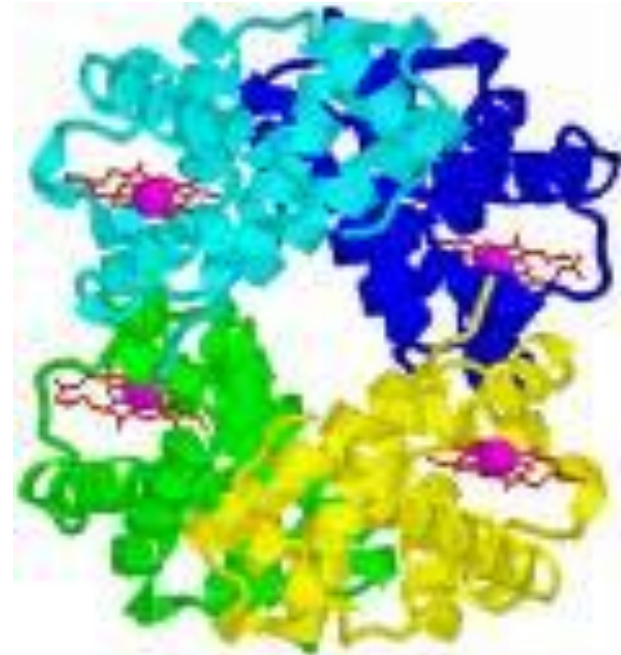
Beta-Turn



Loop



myoglobin



haemoglobin

Quaternary Structure: Complex



G-Protein Complex

Protein Structure Determination

- X-ray crystallography
- Nuclear Magnetic Resonance (NMR) Spectroscopy
- X-ray: any size, accurate (1-3 Angstrom (10^{-10} m)), sometime hard to grow crystal
- NMR: small to medium size, moderate accuracy, structure in solution



[Pacific Northwest National Laboratory](#)'s high magnetic field (800 MHz, 18.8 T) NMR spectrometer being loaded with a sample.

[Wikipedia, the free encyclopedia](#)

Storage in Protein Data Bank

- Home
- Tutorial About This Site
- Getting Started
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- Dictionaries & File Formats
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- BioSync
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- Frequently Asked Questions
- Known Problems
- Report Bugs/Comments

Welcome to the RCSB PDB

The **RCSB PDB** provides a variety of tools and resources for studying the structures of biological macromolecules and their relationships to sequence, function, and disease.

The RCSB is a member of the **wwPDB** whose mission is to ensure that the PDB archive remains an international resource with uniform data.

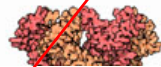
This site offers tools for browsing, searching, and reporting that utilize the data resulting from ongoing efforts to create a more consistent and comprehensive archive.

Information about compatible browsers can be found [here](#).

A **narrated tutorial** illustrates how to search, navigate, browse, generate reports and visualize structures using this **NEW SITE**. [This requires the Macromedia Flash player download.]

Comments? info@rcsb.org

Molecule of the Month: AAA+ Proteases



How would you make a protein cutting machine that would be safe to use inside a cell? Digestive proteases like trypsin and pepsin are small and efficient—they diffuse up to proteins and start cutting. This would never work inside a cell. The cell needs to have more control, so that only obsolete or damaged proteins are destroyed. The

NEWS

- Complete News
- Newsletter
- Discussion Forum

29-August-2006
New RCSB PDB Flyer Available in Print and Online

Two new brochures are available for RCSB PDB users: The General Information trifold & The Easy Steps for Structure Deposition.



Search database

RCSB PDB : Structure Explorer - Mozilla Firefox

File Edit View Go Bookmarks Tools Help

http://www.rcsb.org/pdb/navbsearch.do?newSearch=yes&isAuthorSearch=no&radioSet=All&inputQuickSearch=1vjg&image.x=0&image.y=0&image=Search

Google pdb

RCSB PDB
PROTEIN DATA BANK

A MEMBER OF THE **PDDB**

An Information Portal to Biological Macromolecular Structures

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1VJG

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1VJG

Title Crystal structure of putative lipase from the G-D-S-L family from Nostoc sp. at 2.01 Å resolution

Authors Joint Center for Structural Genomics (JCSG)

Primary Citation Joint Center for Structural Genomics (JCSG) Crystal structure of putative lipase from the G-D-S-L family from Nostoc sp. at 2.01 Å resolution. *To be published*

History Deposition 2004-02-19 Release 2004-03-16

Experimental Method Type X-RAY DIFFRACTION Data [EDS]

Parameters	Resolution[Å]	R-Value	R-Free	Space Group
	2.01	0.175 (obs.)	0.218	P 3 ₂ 2 1

Unit Cell	Length [Å]	a	56.19	b	56.19	c	129.32
	Angles [°]	alpha	90.00	beta	90.00	gamma	120.00

Molecular Description Asymmetric Unit Polymer: 1 Molecule: putative lipase from the G-D-S-L family Chains: A

Functional Class Structural Genomics Unknown Function

Source Polymer: 1 Scientific Name: **Nostoc sp. pcc 7120** Common Name: **Bacteria** Expression system: **Nostoc sp. pcc 7120**

Images and Visualization

Biological Molecule



Display Options

- KING
- Jmol
- WebMol
- Protein Workshop
- QuickPDB
- All Images

Done

Start | Inboxes - Outlook Express | CAP5937 | slides13 | slides1 | RCSB PDB : Structure ... | 10:42 AM Monday

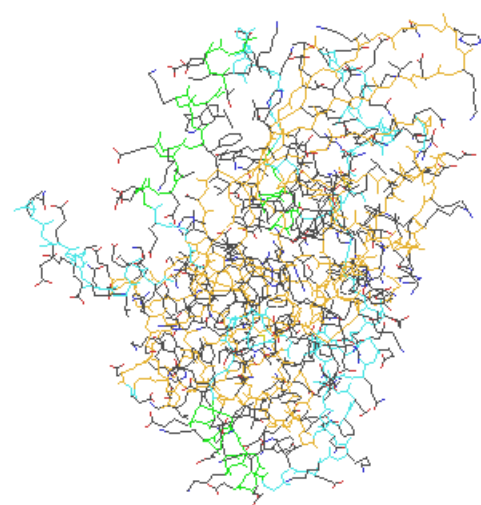
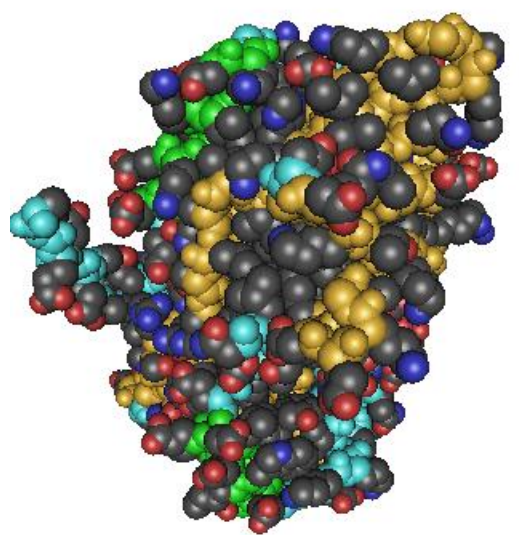
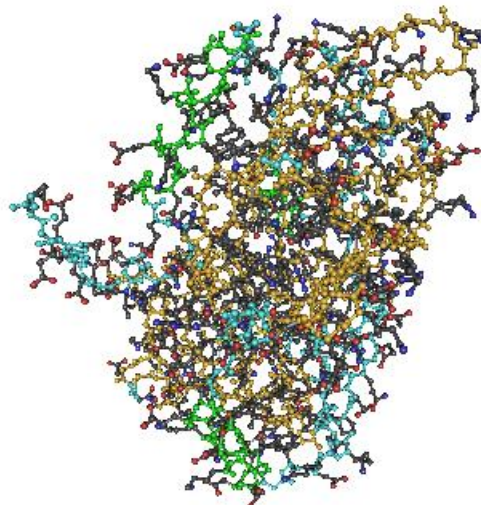
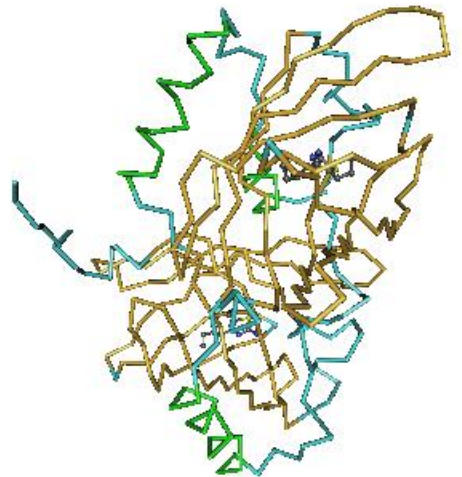
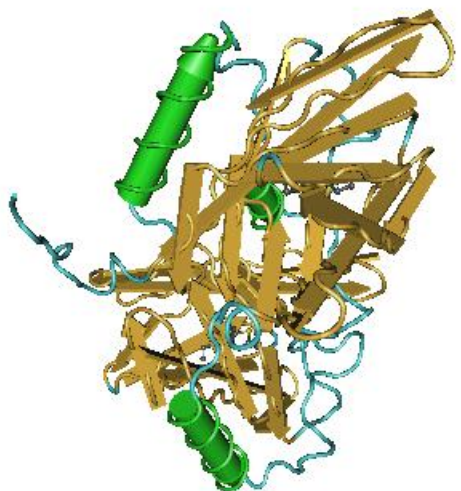
Search protein 1VJG

PDB Format (2C8Q, insulin)

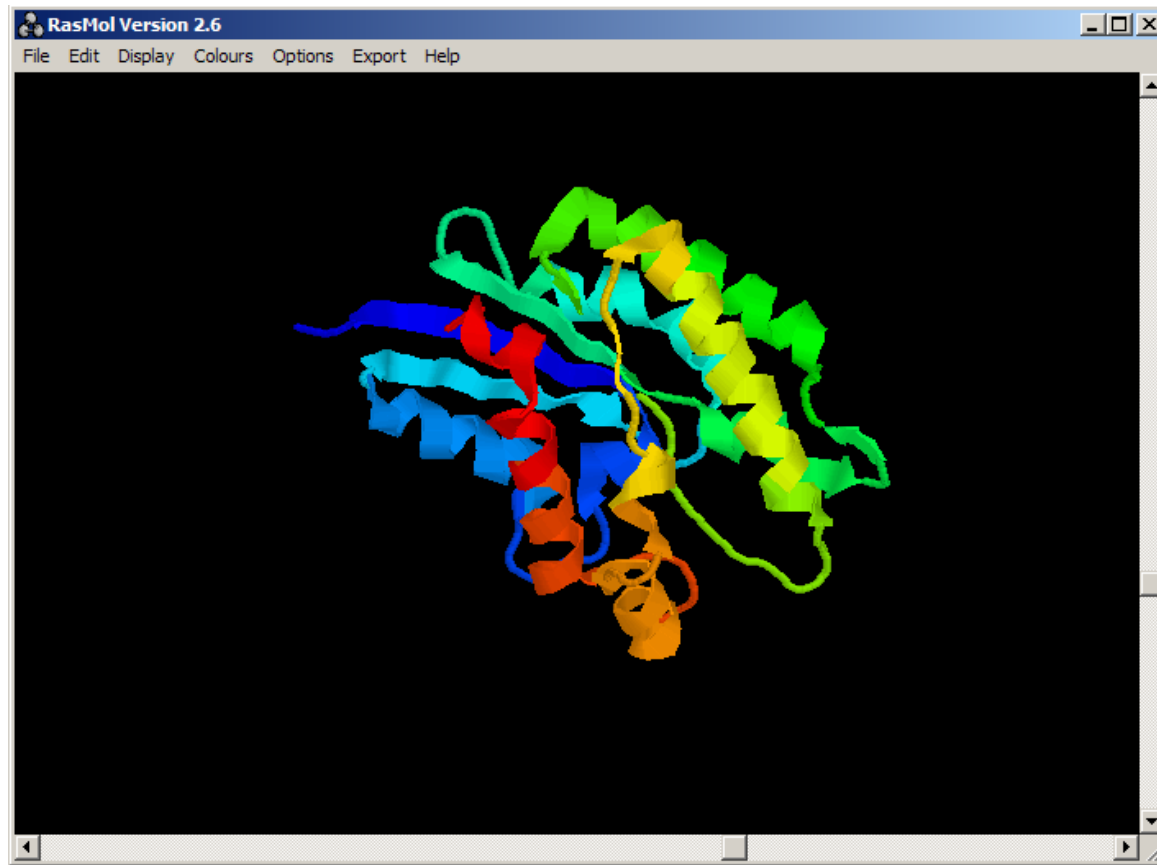
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COMPND     2 MOLECULE: INSULIN A CHAIN;
COMPND     3 CHAIN: A;
COMPND     4 MOL_ID: 2;
COMPND     5 MOLECULE: INSULIN B CHAIN;
COMPND     6 CHAIN: B
SOURCE     MOL_ID: 1;
SOURCE     2 ORGANISM_SCIENTIFIC: HOMO SAPIENS;
SOURCE     3 ORGANISM_COMMON: HUMAN;
SOURCE     4 ORGAN: PANCREAS;
SOURCE     5 MOL_ID: 2;
SOURCE     6 ORGANISM_SCIENTIFIC: HOMO SAPIENS;
SOURCE     7 ORGANISM_COMMON: HUMAN;
SOURCE     8 ORGAN: PANCREAS
KEYWDS     LASER, UV, CARBOHYDRATE METABOLISM, HORMONE, DIABETES
KEYWDS     2 MELLITUS, GLUCOSE METABOLISM
EXPDTA     X-RAY DIFFRACTION
AUTHOR     X.VERNEDE,B.LAVAUTL,J.OHANA,D.NURIZZO,J.JOLY,L.JACQUAMET,
AUTHOR     2 F.FELISAZ,F.CIPRIANI,D.BOURGEOIS
REVDTA     1   08-MAR-06 2C8Q   0
JRNL       AUTH   X.VERNEDE,B.LAVAUTL,J.OHANA,D.NURIZZO,J.JOLY,
JRNL       AUTH 2 L.JACQUAMET,F.FELISAZ,F.CIPRIANI,D.BOURGEOIS
JRNL       TITL   UV LASER-EXCITED FLUORESCENCE AS A TOOL FOR THE
JRNL       TITL 2 VISUALIZATION OF PROTEIN CRYSTALS MOUNTED IN
JRNL       TITL 3 LOOPS.
JRNL       REF   ACTA CRYSTALLOGR.,SECT.D           V.   62   253 2006
JRNL       REFN  ASTM ABCRE6  DK ISSN 0907-4449
REMARK     2
REMARK     2 RESOLUTION. 1.95 ANGSTROMS.
REMARK     3
REMARK     3 REFINEMENT.
REMARK     3   PROGRAM       : REFMAC 5.2.0005
REMARK     3   AUTHORS        : MURSHUDOV,VAGIN,DODSON
REMARK     3
REMARK     3   REFINEMENT TARGET : MAXIMUM LIKELIHOOD
```


Structure Visualization

- Rasmol
(<http://www.umass.edu/microbio/rasmol/getras.htm>)
- MDL Chime (plug-in)
(<http://www.mdl.com/products/framework/chime/>)
- Protein Explorer
(<http://molvis.sdsc.edu/protexpl/frntdoor.htm>)
- Jmol: <http://jmol.sourceforge.net/>
- Pymol: <http://pymol.sourceforge.net/>



Rasmol (1VJG)



Structure Analysis

- Assign secondary structure for amino acids from 3D structure
- Generate solvent accessible area for amino acids from 3D structure
- Most widely used tool: DSSP (Dictionary of Protein Secondary Structure: Pattern Recognition of Hydrogen-Bonded and Geometrical Features. **Kabsch and Sander, 1983**)

DSSP server: <http://bioweb.pasteur.fr/seqanal/interfaces/dssp-simple.html>

DSSP download: <http://swift.cmbi.ru.nl/gv/dssp/>

DSSP Code:

H = alpha helix

G = 3-helix (3/10 helix)

I = 5 helix (pi helix)

B = residue in isolated beta-bridge

E = extended strand, participates in beta ladder

T = hydrogen bonded turn

S = bend

Blank = loop

DSSP Web Service

**DSSP : Definition of secondary structure of proteins given a set of 3D coordinates
(W.Kabsch, C. Sander)**

your e-mail

PDB File

or you can instead enter a PDB id.

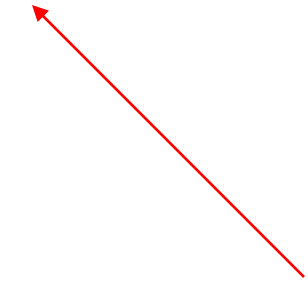
<http://bioweb.pasteur.fr/seqanal/interfaces/dssp-simple.html>

#	RESIDUE	AA	STRUCTURE	BP1	BP2	ACC	N-H-->O	O-->H-N	N-H-->O	O-->H-N	TCO	KAPPA ALPHA	PHI	PSI	X-CA	Y-CA	Z-CA		
1	5	A	S		0	0	179	0, 0.0	2, -0.0	0, 0.0	0, 0.0	0.000	360.0	360.0	360.0	125.7	-8.6	43.0	43.9
2	6	A	K	-	0	0	123	1, -0.1	2, -0.4	37, -0.1	37, -0.2	-0.235	360.0-108.7	-87.0	151.4	-7.5	41.4	40.6	
3	7	A	T E	-a	39	0A	75	35, -0.6	37, -2.5	1, -0.0	2, -0.3	-0.593	34.7-132.0	-72.2	128.3	-4.3	39.5	39.6	
4	8	A	Q E	+a	40	0A	91	-2, -0.4	69, -0.6	35, -0.2	2, -0.4	-0.639	26.0 179.8	-86.4	132.7	-2.0	41.5	37.4	
5	9	A	I E	-ab	41	73A	3	35, -1.9	37, -2.9	-2, -0.3	2, -0.5	-0.991	13.3-156.5-129.4	131.5	-0.7	39.9	34.2		
6	10	A	R E	-ab	42	74A	48	67, -2.8	69, -1.7	-2, -0.4	2, -0.4	-0.910	14.8-173.2-105.2	126.8	1.6	41.6	31.8		
7	11	A	I E	-ab	43	75A	0	35, -2.5	37, -2.6	-2, -0.5	2, -0.5	-0.983	11.9-162.4-124.9	124.4	1.7	40.3	28.2		
8	12	A	C E	-ab	44	76A	0	67, -2.3	69, -2.6	-2, -0.4	2, -0.6	-0.931	6.5-159.9-100.8	130.8	3.9	41.2	25.3		
9	13	A	F E	-ab	45	77A	0	35, -2.2	37, -3.0	-2, -0.5	2, -0.5	-0.955	13.2-169.0-109.5	117.1	2.7	40.2	21.8		
10	14	A	V E	+ab	46	78A	0	67, -3.1	69, -2.2	-2, -0.6	2, -0.3	-0.926	34.8 71.1-116.5	129.9	5.6	40.1	19.4		
11	15	A	G E	S-ab	47	79A	0	35, -0.9	37, -1.9	-2, -0.5	69, -0.2	-0.921	70.2 -50.2	169.0-146.4	5.3	39.9	15.6		
12	16	A	D S >> S-		0	0	4	67, -0.8	4, -2.2	-2, -0.3	3, -0.6	-0.023	78.2 -51.3-111.5-151.8	4.2	41.6	12.4			
13	17	A	S H 3>>S+		0	0	7	35, -0.3	5, -1.7	1, -0.2	4, -1.5	0.803	130.2 57.8 -67.3 -28.8	1.2	43.5	11.1			
14	18	A	F H 345S+		0	0	5	2, -0.2	12, -0.5	1, -0.2	-1, -0.2	0.884	108.5 46.5 -68.2 -33.2	-1.2	40.8	12.2			
15	19	A	V H <45S+		0	0	1	-3, -0.6	12, -0.3	64, -0.2	-2, -0.2	0.900	111.1 52.2 -68.9 -41.4	-0.0	41.1	15.7			
16	20	A	N H <5S-		0	0	71	-4, -2.2	-2, -0.2	30, -0.1	-1, -0.2	0.774	110.8-127.0 -62.6 -26.6	-0.3	45.0	15.4			
17	21	A	G T ><5 -		0	0	5	-4, -1.5	3, -2.2	-5, -0.2	8, -0.4	0.741	36.4-174.6 83.1 25.3	-3.9	44.5	14.2			
18	22	A	T T 3 < +		0	0	14	-5, -1.7	-1, -0.2	1, -0.3	-2, -0.0	-0.199	68.4 29.2 -54.0 135.4	-3.4	46.6	11.0			
19	23	A	G T 3 S+		0	0	28	1, -0.3	-1, -0.3	159, -0.1	162, -0.2	0.121	86.2 120.8 94.7 -21.4	-6.7	47.0	9.2			
20	24	A	D X -		0	0	9	-3, -2.2	3, -1.2	160, -0.2	-1, -0.3	-0.706	48.9-160.5 -79.7 117.6	-8.9	46.8	12.4			
21	25	A	P T 3 S+		0	0	91	0, 0.0	-1, -0.2	0, 0.0	159, -0.0	0.677	91.8 60.1 -70.9 -17.3	-10.9	50.1	12.6			
22	26	A	E T 3 S-		0	0	119	-3, -0.0	-2, -0.1	3, -0.0	158, -0.0	0.426	105.0-132.3 -87.9 -3.3	-11.4	49.4	16.3			
23	27	A	C S < S+		0	0	112	-3, -1.2	-5, -0.1	-6, -0.2	-6, -0.0	0.730	80.2 98.1 62.8 28.1	-7.6	49.4	16.9			

Amino
Acids

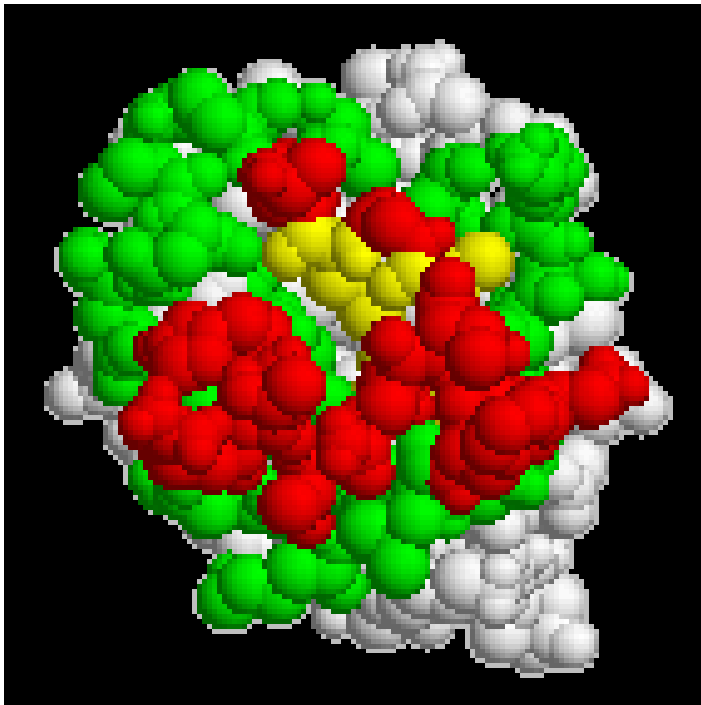
Secondary
Structure

Solvent
Accessibility



Solvent Accessibility

Size of the area of an amino acid that is exposed to solvent (water).



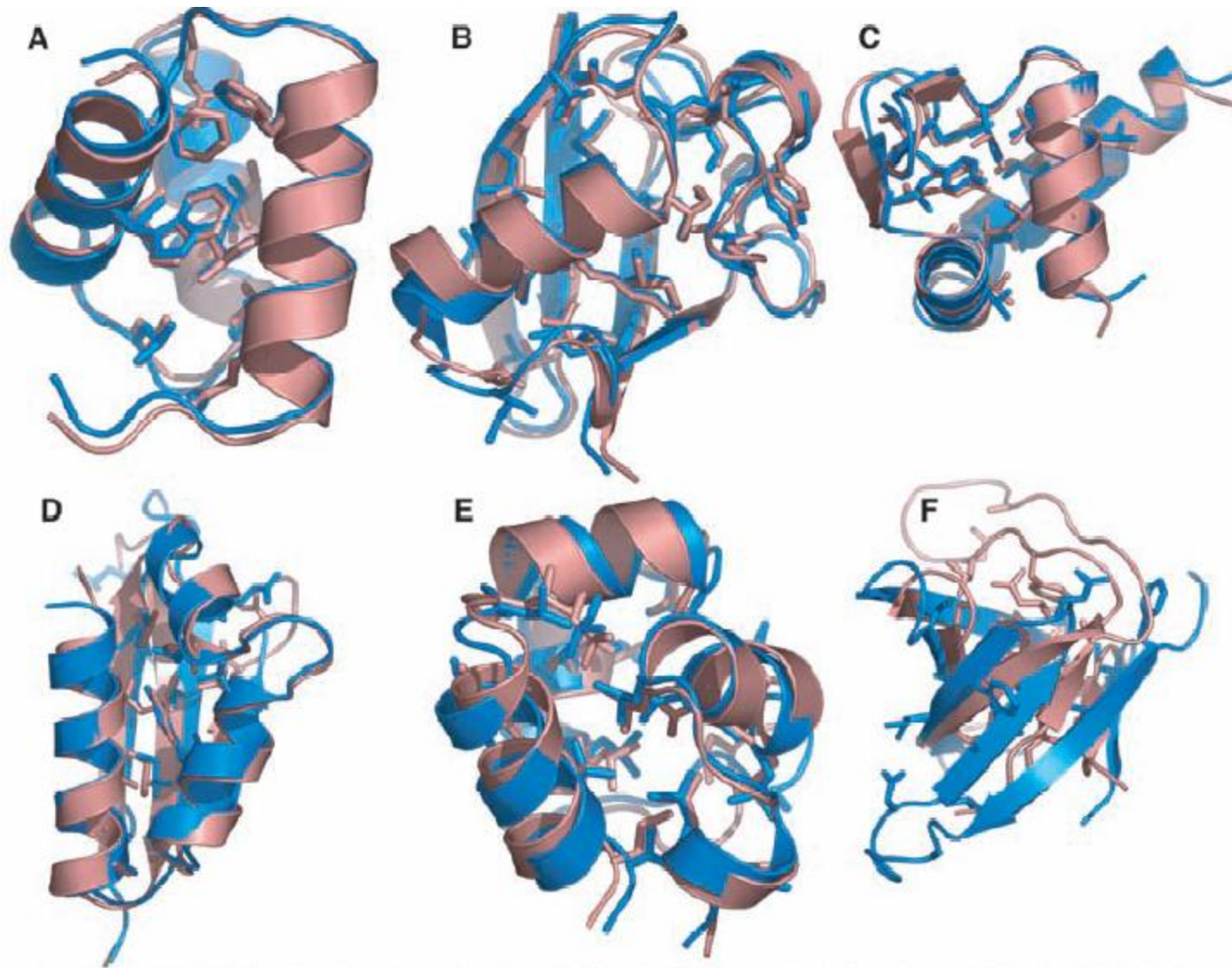
Maximum solvent accessible area for each amino acid is its whole surface area.

Hydrophobic residues like to be Buried inside (interior).

Hydrophilic residues like to be exposed on the surface.

Structure Comparison (Alignment)

- Are the structures of two protein similar?
- Are the two structure models of the same protein similar?
- Different measures (RMSD, GDT-TS (Zemla et al., 1999), MaxSub (Siew et al., 2000), TM score (Zhang and Skolnick, 2005))

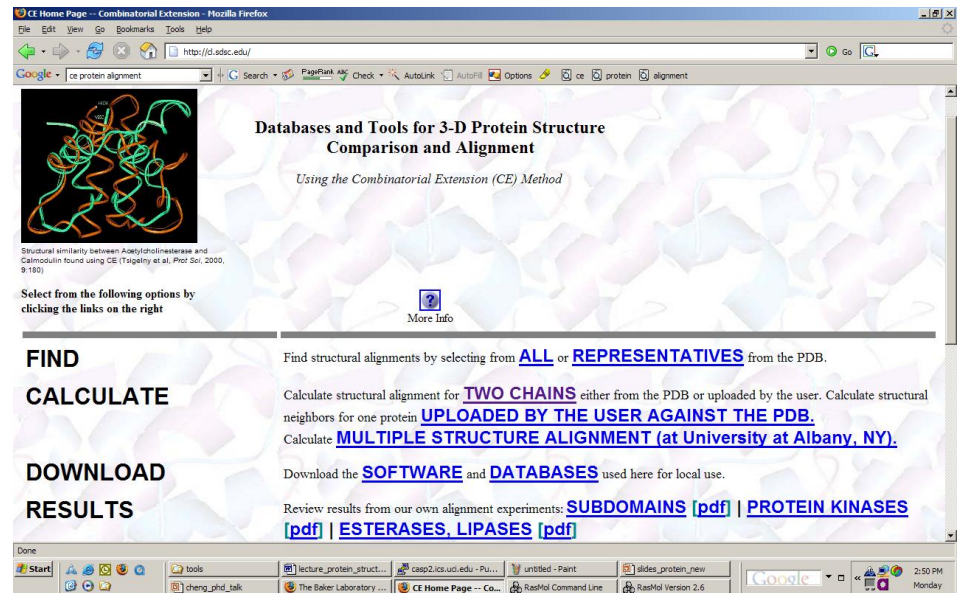


Superimposition

David Baker, 2005

Useful Structure Alignment Tools

- CE
(<http://cl.sdsc.edu/>)
- DALI
(<http://www.ebi.ac.uk/dali/>)
- TM-Align:
<http://zhang.bioinformatics.ku.edu/TM-align/>



CE CALCULATE TWO CHAINS Calculate structural alignment for two polypeptide chains either from the PDB or uploaded by the user.

Specify two polypeptide chains and optionally the similarity level and use of sequence information and then press the "Calculate Alignment" button. Selecting the appropriate ? will provide help on that spe

Calculate Alignment Reset Form

Select Similarity Level: Medium ?
 Use Sequence Information (optional) ?

Chain 1:	<input type="radio"/> PDB: 4HHB:A ? OR <input checked="" type="radio"/> User File: C:\casp7\301\foldpro1.pdb Browse... Chain ID: <input type="text"/> ? <input type="checkbox"/> Use Fragment From: <input type="text"/> To: <input type="text"/> (optional) ? Sequence numbering <input type="text"/>
Chain 2:	<input type="radio"/> PDB: 4HHB:B ? OR <input checked="" type="radio"/> User File: C:\casp7\301\ROBETTA_TS1.pdl Browse... Chain ID: <input type="text"/> ? <input type="checkbox"/> Use Fragment From: <input type="text"/> To: <input type="text"/> (optional) ? Sequence numbering <input type="text"/>

USR1:_(size=395) vs USR2:_(size=395) Structure Alignment

Rmsd = 2.4Å Z-Score = 6.6
 Sequence identity = 42.8%
 Aligned/gap positions = 332/105

Sequence alignment based on structure alignment.

Sequence alignment based on structure alignment. Position numbers according to sequence (starting from 1) and according to PDB are given as SSSS/PPPP, SSSS - sequence, PPPP - PDB.

USR1: _ -

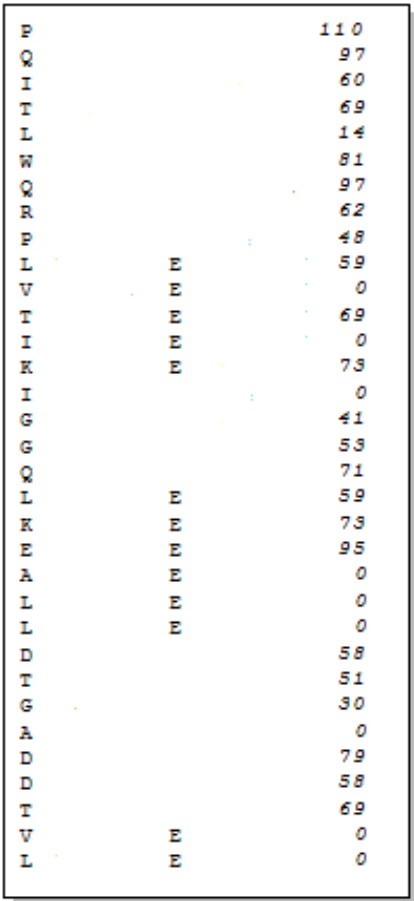
USR2: _ -

USR1: 4/5 PPQIRIPATYLRGGTSKGVFFRLEDLPE-----SCRVPGEARDRLFMRVIGSPDPYAA
 USR2: 6/7 QRIRIPATYLRGGTSKGVFFRL-----EDLPESCRVPGEARDRLFMRVIGSPDPYA---A

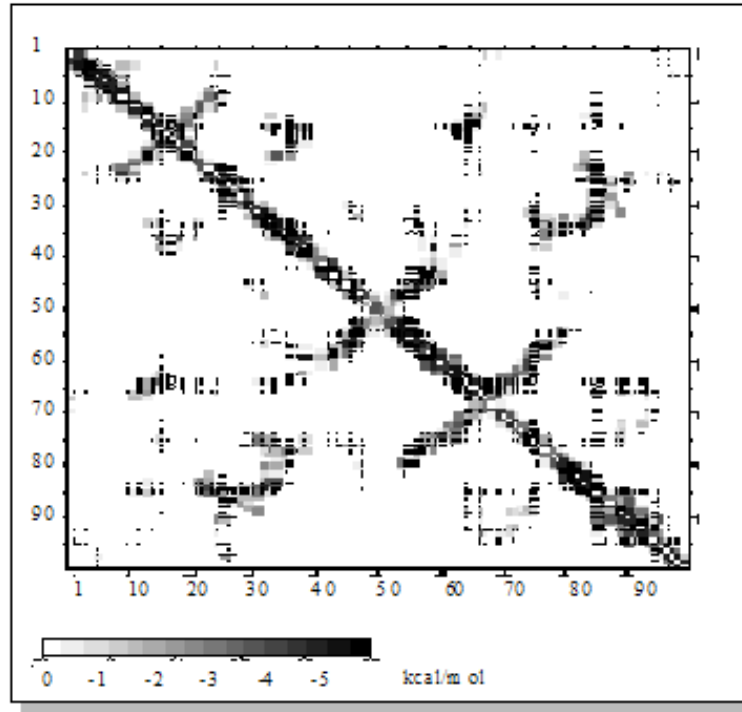
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 USR2: 57/58 HIDGMGGATSSTSKCVILSKSSQPGHDVDYLYGQVSIKPFVDWSGNCNLSTGAGAFAL

USR1: 117/118 HAGLVDPARIPEDGICEVRIWQANIGKTIIAHVFPVSGGQVQETGDFELDGVTFPAAEIVL
 USR2: 117/118 HAGLVDPARIPEDGICEVRIWQANIGKTIIAHVFPVSGGQVQETGDFELDGVTFPAAEIVL

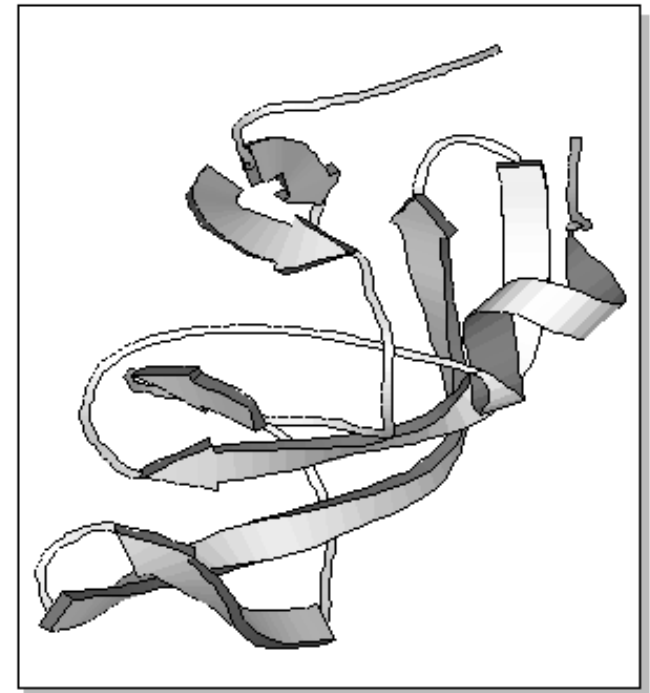
Notation: protein structure 1D, 2D, 3D



1D



2D



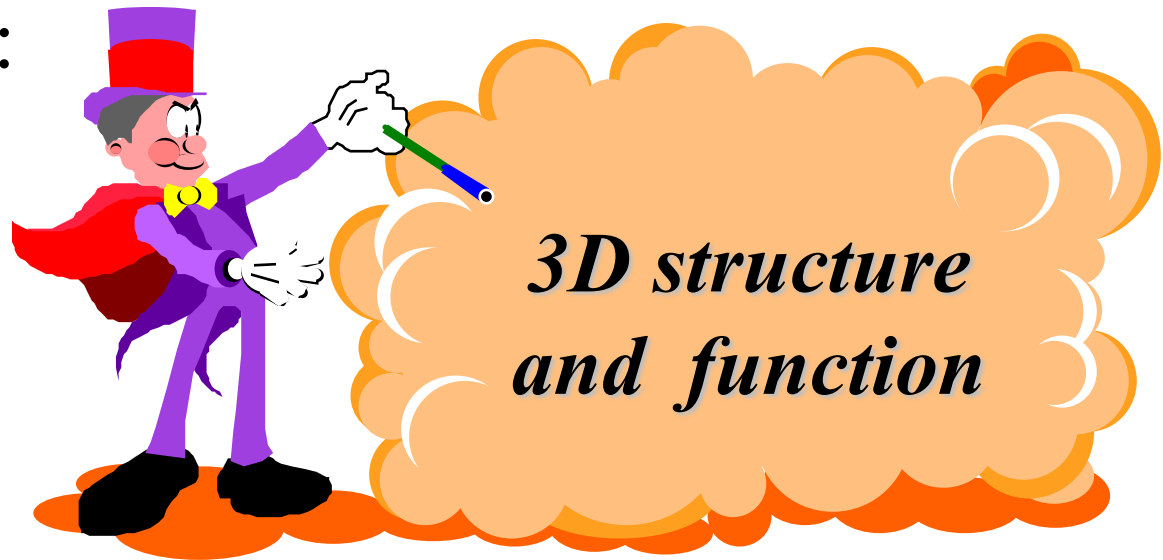
3D

Goal of structure prediction

- Epstein & Anfinsen, 1961:
sequence uniquely determines structure

- INPUT: sequence

- OUTPUT:



CASP – Olympics of Protein Structure Prediction

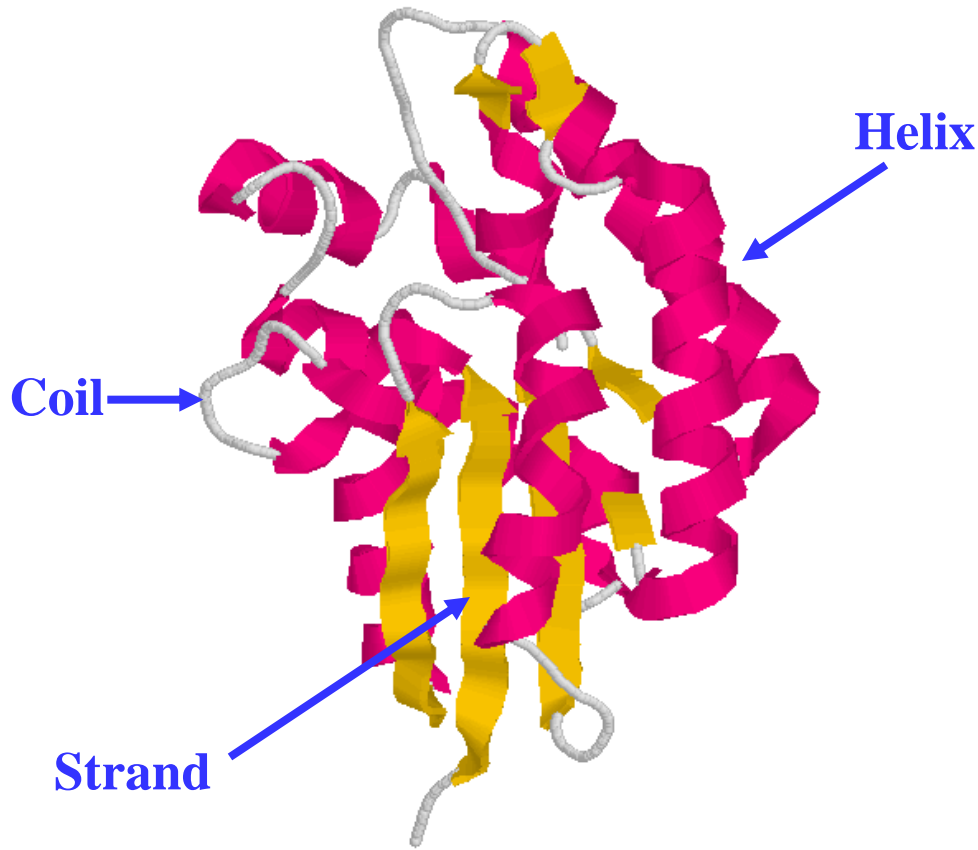
- Critical Assessment of Techniques of Protein Structure Prediction
- 1994,1996,1998,2000,2002,2004,2006, 2008, 2010
- Blind Test, Independent Evaluation
- CASP9: 116 targets



1D Structure Prediction

- Secondary structure
- Solvent accessibility
- Disordered regions
- Domain boundary

1D: Secondary Structure Prediction



MWLKKFGINLLIGQSV...



**Neural Networks
+ Alignments**



CCCC**HHHH**CCC**SSSS**...

Widely Used Tools (~78-80%)

SSpro 4.1: http://sysbio.rnet.missouri.edu/multicom_toolbox/

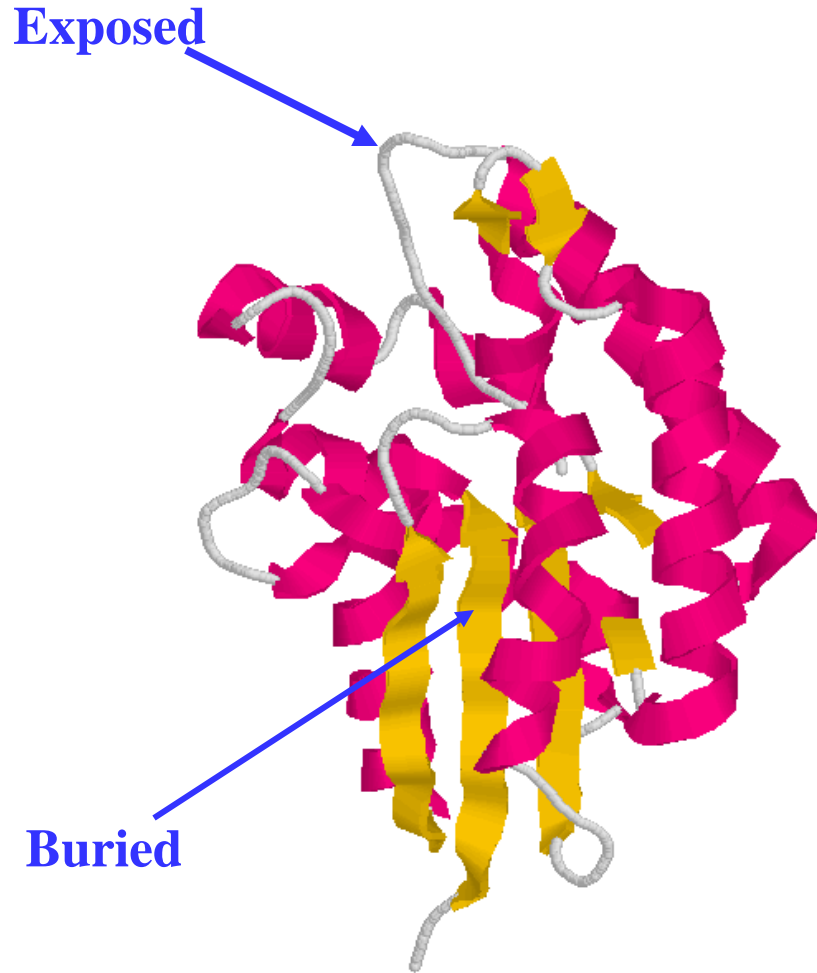
Distill: <http://distill.ucd.ie/porter/>

PSI-PRED: <http://bioinf.cs.ucl.ac.uk/psipred/psiform.html>
software is also available

SAM: http://compbio.soe.ucsc.edu/SAM_T08/T08-query.html

PHD: <http://www.predictprotein.org/>

1D: Solvent Accessibility Prediction



MWLKKFGINLLIGQSV...

Neural Networks
+ Alignments

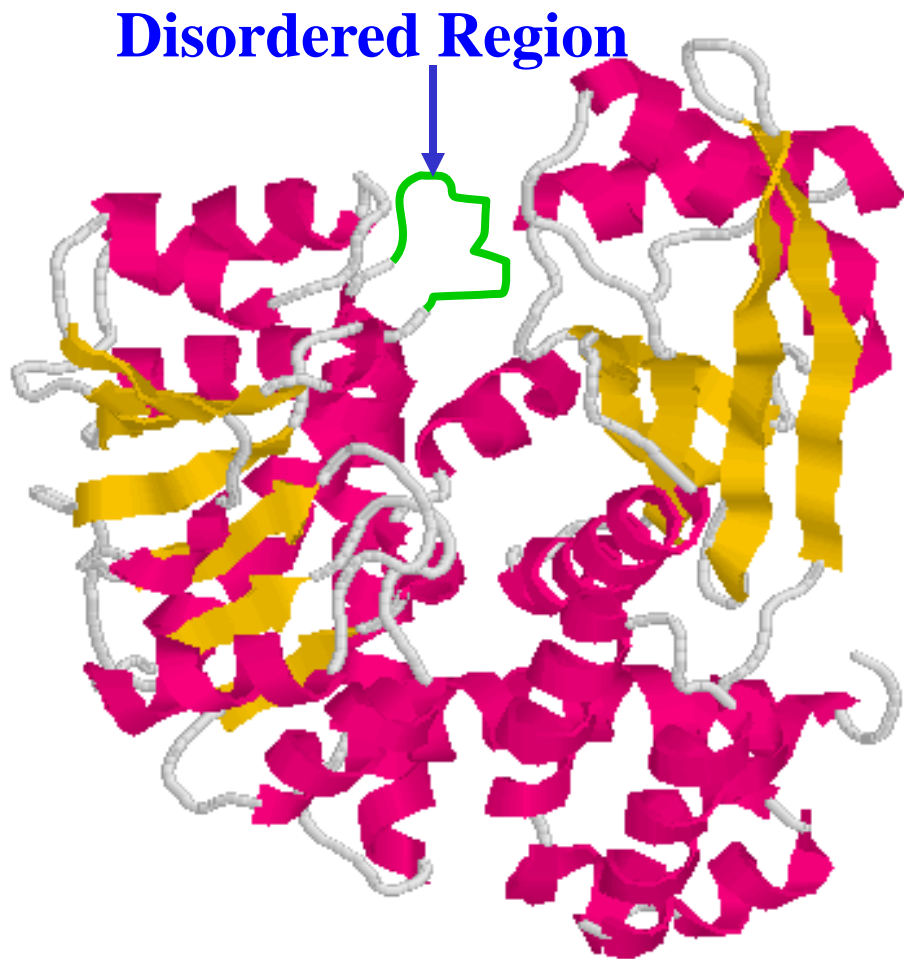
eeeeee**bbbbbb**eeeebb...

Accuracy: 79% at 25% threshold

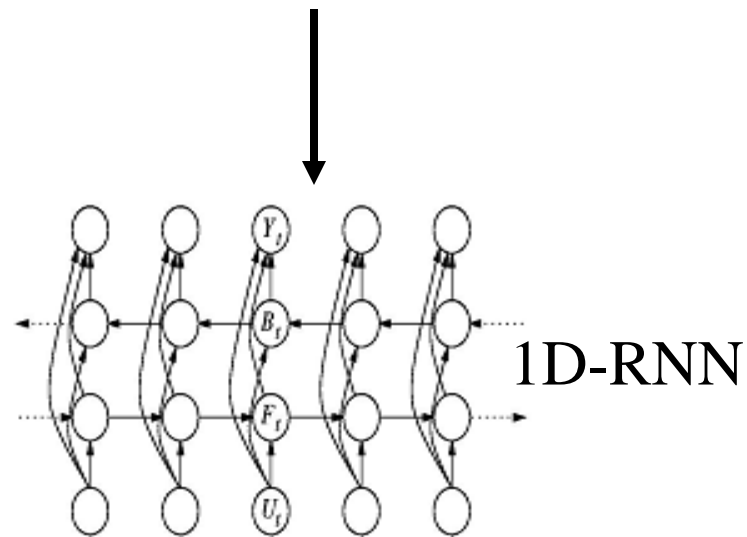
Widely Used Tools (78%)

- ACCpro 4.1: software:
http://sysbio.rnet.missouri.edu/multicom_toolbox/
- SCRATCH: <http://scratch.proteomics.ics.uci.edu/>
- PHD: <http://www.predictprotein.org/>
- Distill: <http://distill.ucd.ie/porter/>

1D: Disordered Region Prediction Using Neural Networks



MWLKKFGINLLIGQSV...



OOOOODDDDOOOOO...

93% TP at 5% FP

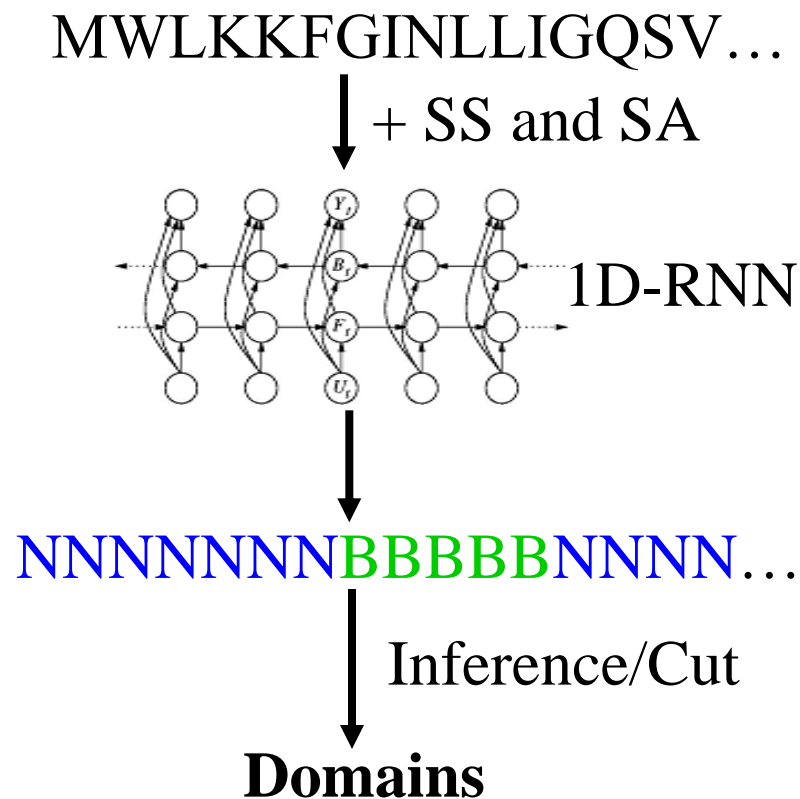
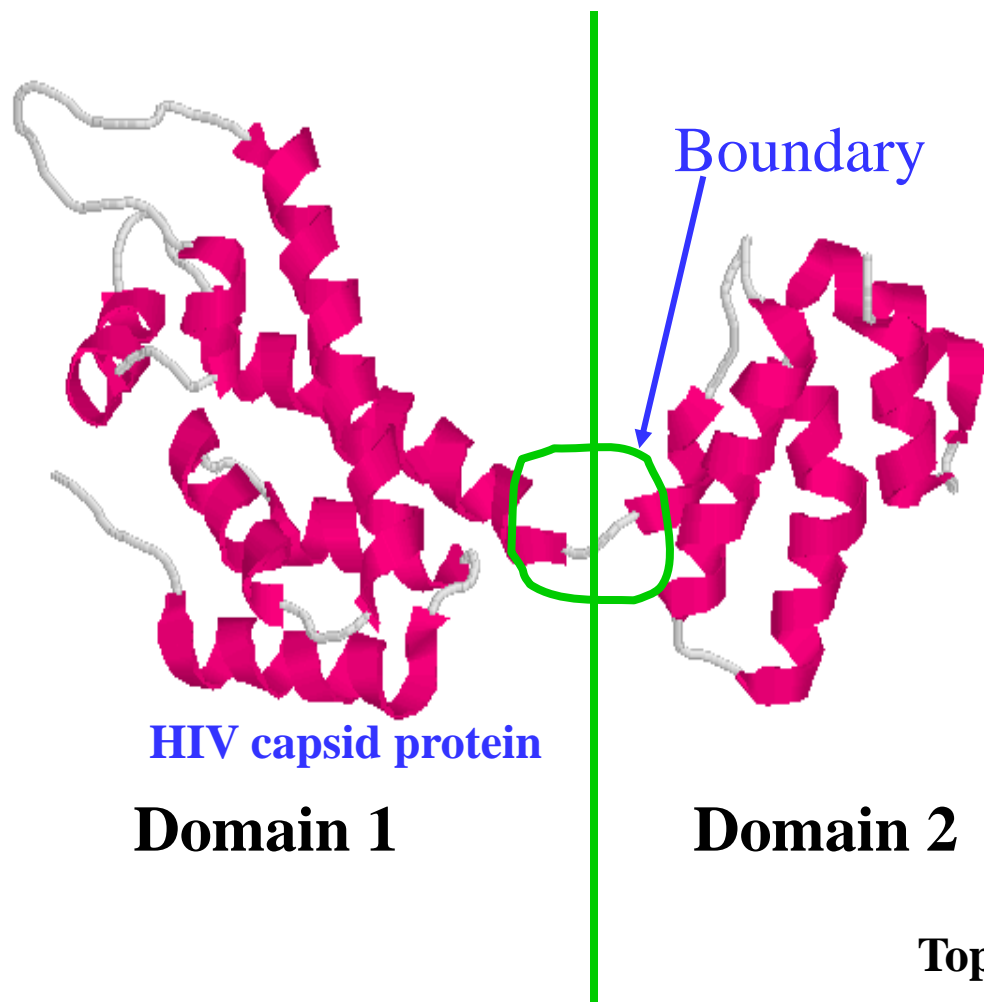
Tools

PreDisorder: http://sysbio.rnet.missouri.edu/multicom_toolbox/

A collection of disorder predictors:

<http://www.disprot.org/predictors.php>

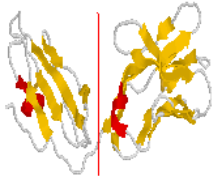
1D: Protein Domain Prediction Using Neural Networks



Top *ab-initio* domain predictor in CAFASP4

Tools

DOMAC: <http://casp.rnet.missouri.net/domac.html>



DOMAC: An Accurate, Hybrid Protein Domain Prediction Server
[\(Help\)](#)

Email address(where the prediction will be sent):

Target Name(required):

Protein sequence(one plain sequence, no headers):

Reference:

J. Cheng. DOMAC: An Accurate, Hybrid Protein Domain Prediction Server. Nucleic Acids Research, vol. 35, pp. w354-w356, 2007.

[Dr. Jianlin Cheng's Bioinformatics and Systems Biology Laboratory](#)
[Department of Computer Science](#)
[University of Missouri](#)

Cheng, Nucleic Acids Research, 2007

DoBo

Protein domain boundary prediction by integrating evolutionary signals and machine learning


Have a question? Maybe it's answered in the [FAQ](#)

Job Details


Job title (optional)

Sequence

Plain sequence. Spaces, newlines and any FASTA header will be ignored.
Minimum sequence length is 90 residues.

Confidence level 

Set a minimum threshold for the confidence of domain boundary predictions.

Single/multi-domain classification 

Run an additional check to classify query as a single or multi-domain protein.

Web: http://sysbio.rnet.missouri.edu/multicom_toolbox/index.html

Reference:

J. Eickholt, X. Deng, and J. Cheng. **DoBo: Protein Domain Boundary Prediction by Integrating Evolutionary Signals and Machine Learning.** *BMC Bioinformatics*. 12:43, 2011.

1. Input query

LNKGQRHIKIREIIMS...

2. Identify homologous sequences w/ PSI-BLAST



3. Extract pairwise alignments

```

Query 1 LNKGQRHIKIREIIMSNDIETQDELVDRLREAGFNVTQATVSRDIKEMQLVKVPMANGRY 60
Sbjct 1 MNKGQRHIKIREIIANKEIETQDELVDILRNEGFNVTQATVSRDIKELHLVKVPLHDGRY 60
...
Query 6 RHIKIREIIMSNDIETQDELVDRLREAGFNVTQATVSRDIKEMQLVKVPMANGRYKYSLP 65
Sbjct 5 RHSKILEILNKYEVEVTQEDLTEYLREAGINVTQATVSRDIRQMMLVKVMTKSGKYKYAAY 64
...
Query 1 LNKGQRHIKIREIIMSNDIETQDELVDRLREAGFNVTQATVSRDIKEMQLVKVPMANGRY 60
Sbjct 1 MNKGQRHIKIREIIANKEIETQDELVDILRNEGFNVTQATVSRDIKELHLVKVPLHDGRY 60

```

4. Form multiple sequence alignment



5. Identify domain boundary signals

Gap 45 residues or longer



Remaining sequence longer than 45 residues



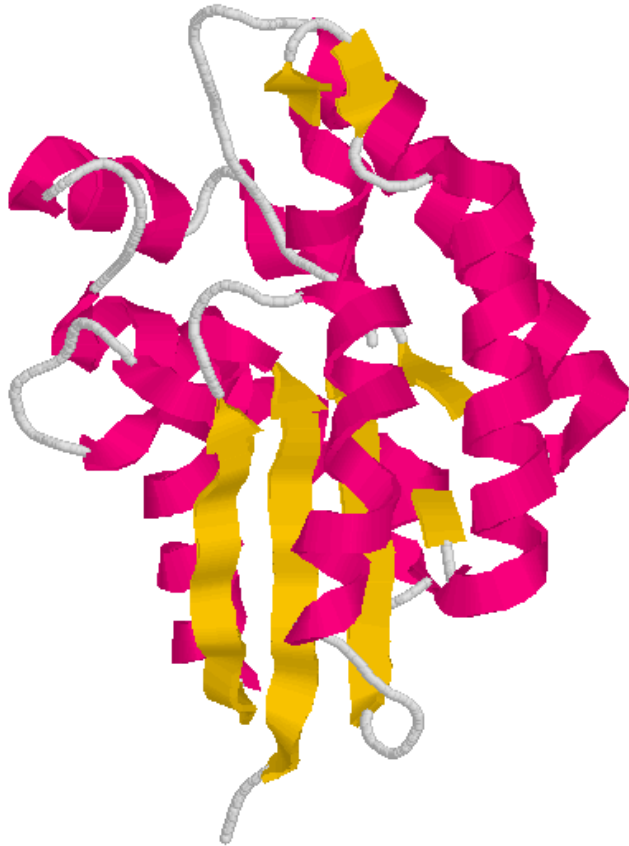
Domain boundary signal (indicated by large arrows)

Project 1

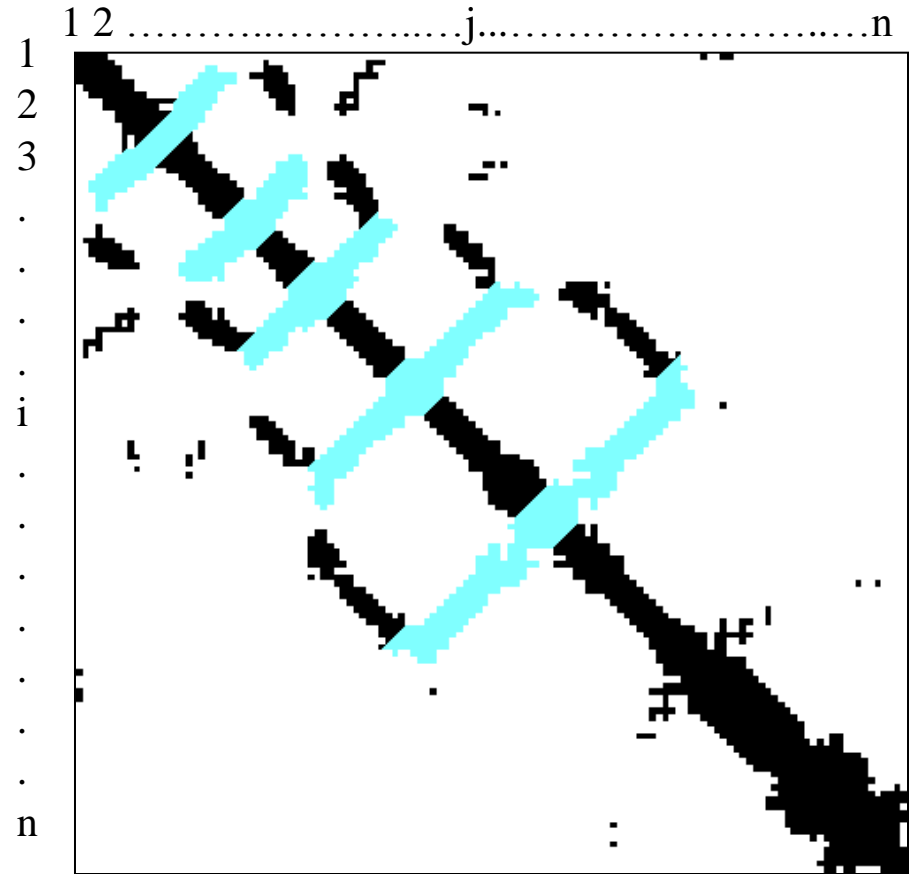
- Predict Secondary Structure, Solvent Accessibility, Disorder Regions of soybean transcription factors
- Data:
http://casp.rnet.missouri.edu/marc/muii_7005/SEQ_TFP_90_2500.txt
- Select **10** proteins to make predictions

2D: Contact Map Prediction

3D Structure



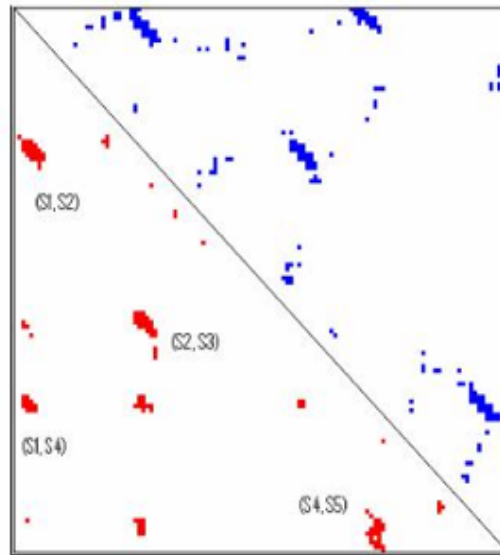
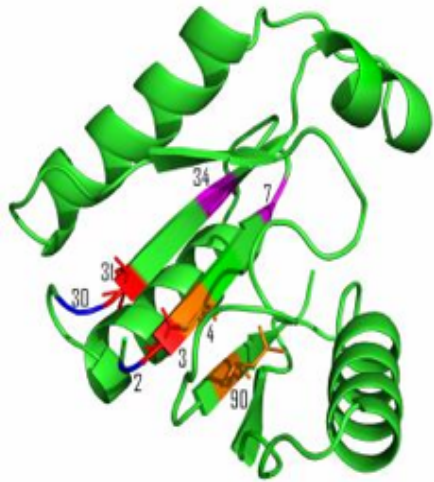
2D Contact Map



Distance Threshold = 8\AA

Contact Prediction

- SVMcon:
<http://casp.rnet.missouri.edu/svmcon.html>
- NNcon:
<http://casp.rnet.missouri.edu/nncon.html>
- SCRATCH:
<http://scratch.proteomics.ics.uci.edu/>
- SAM: <http://compbio.soe.ucsc.edu/HMM-apps/HMM-applications.html>



NNcon: Protein Contact Map Prediction Using Artificial Neural Networks ([Help](#))

Email address(where the prediction will be sent):

Target Name(required):

Protein sequence(one plain sequence, no headers, and length < 1000 amino acids; an example sequence is [here](#)):

Predict

Two Methodologies for 3D Structure Prediction

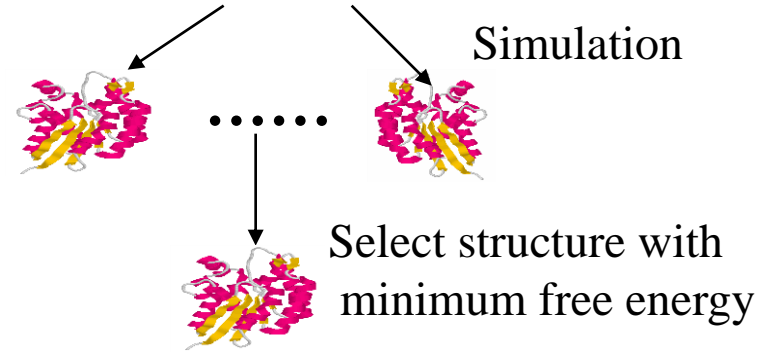
- AB Initio Method (physical-chemical principles / molecular dynamics, knowledge-based approaches)
- Template-Based Method (knowledge-based approaches)

Two Approaches

• Ab Initio Structure Prediction

Physical force field – protein folding
Contact map - reconstruction

MWLKKFGINLLIGQSV...



• Template-Based Structure Prediction

Query protein

MWLKKFGINKH...



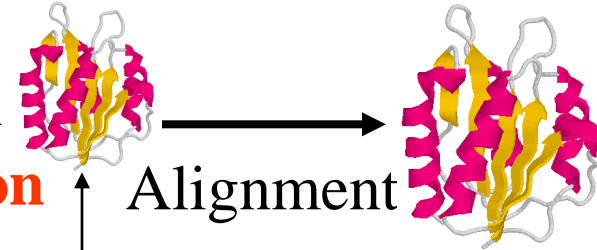
Protein Data Bank

Fold

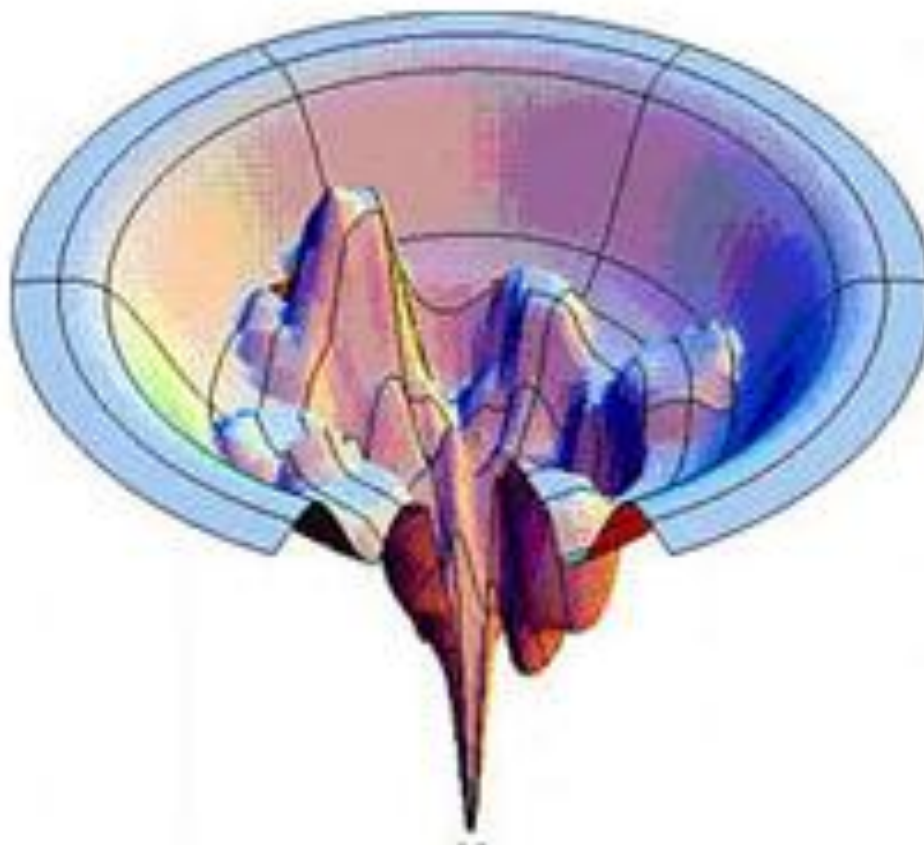
Recognition

Alignment

Template

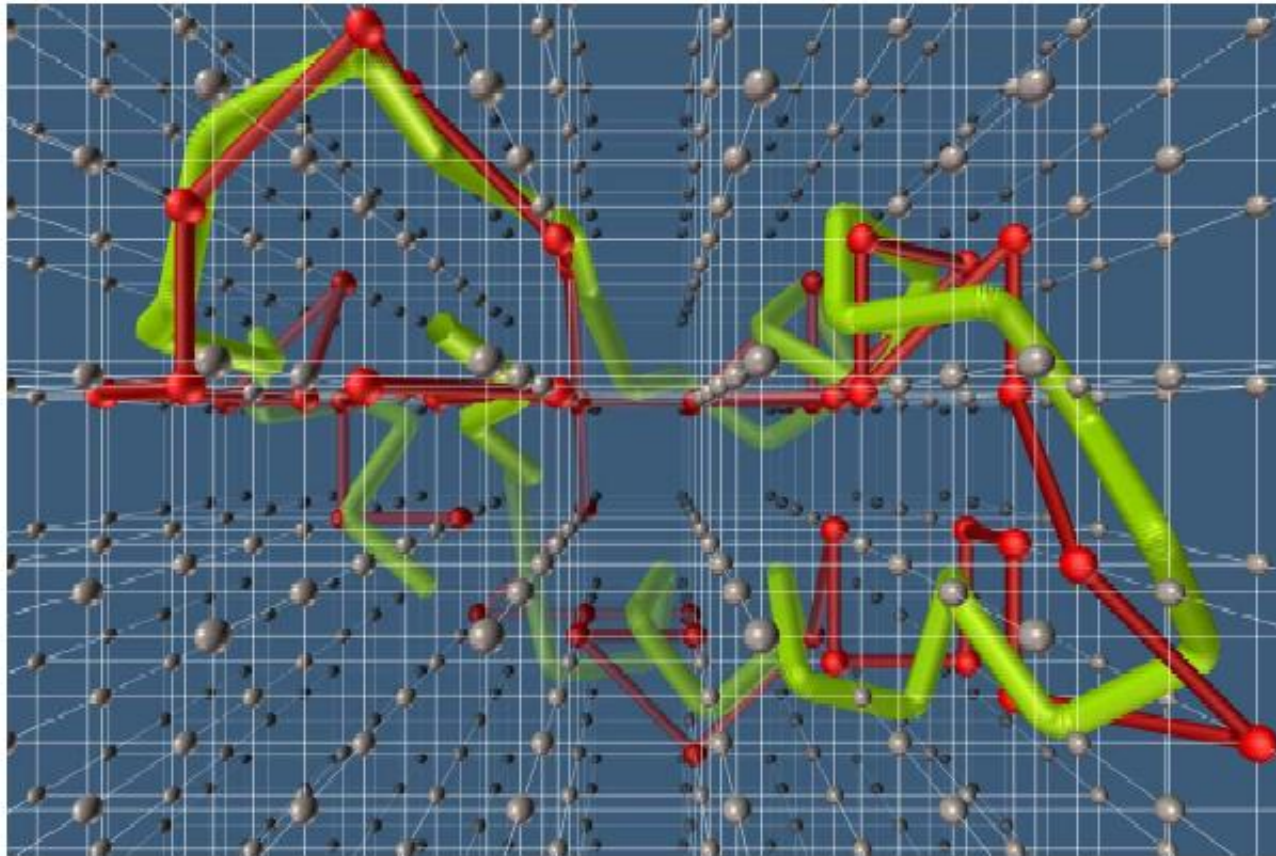


Protein Energy Landscape



C. Park, 2005

Markov Chain Monte Carlo Simulation



Template-Based Structure Prediction

1. Template identification
2. Query-template alignment
3. Model generation
4. Model evaluation
5. Model refinement

Notes: if template is easy to identify, it is often called **comparative Modeling or homology** modeling. If template is hard to identify, it is often called **fold recognition**.

TARGET

TEMPLATE

ASILPKRLFGNCEQTSDEGLK
IERTPLVPHISAQNVCLKIDD
VPERLIPERASFQWMNDK



ASILPKRLFGNCEQTSDEGLKIERTPLVPHISAQNVCLKIDDVPERLIPE
MSVIPKRLYGNCEQTSEEAIRIEDSPIV---TADLVCLKIDEIPERLVGE



Modeller

- Need an alignment file between query and template sequence in the PIR format
- Need the structure (atom coordinates) file of template protein
- You need to write a simple script (Python for version 8.2) to tell how to generate the model and where to find the alignment file and template structure file.
- Run Modeller on the script. Modeller will automatically copy coordinates and make necessary adjustments to generate a model.
- See project step 5-8 for more details.

An PIR Alignment Example

Diagram illustrating a PIR alignment example with annotations:

- Template id
- Template structure file id
- Structure determination method
- Start index
- End index
- Query sequence id

```
>P1;1SDMA
structureX:1SDMA: 1: : 344: : : :
KIRVYCRLRPLCEKEIIAKERNAIRSVDEFTVEHLWKDDKAKQHMYDRVFDGNATQDDVFEDTKYL
VQSAVDGYNVCIFAYGQTGSGKTFTIYGADSNPGLTPRAMSELFKIMKKDSNKFSFSLKAYMVELY
QDTLVDLLLPKQAKRLKLDIKKDSKGMVSVENVTVVVISSTYEELKTTIQRGSEQRHTTGTLMNEQS
SRSHLIVSVI IESTNLQTQAIARGKLSFVDLAGSERVKKEAQSINKSLSALGDVISALSSGNQHIP
YRNHKL TMLMSDSLGGNAK TLMFVNISP AENLDETHNSLT YASRVRSIVNDPSKNVSSKEVARLK
KLVS YWELEEI QDE*
>P1;bioinfo
: : : : : : : :
NIRVIARVRPVTKEDGEGPEATNAVTFDADDDSI I HLLHKGKPVSFELDKVFS PQASQQDVFQEVQ
ALVTSCIDGFNVCIFAYGQTGAGKTYTMEGTAENPGINQRALQLLFSEVQEKASDWEYTTITVSAAE
IYNEVLRDLLGKEPQEKLEIRLCPDGSQQLYVPGLTEFQVQSVDDINKVFEFGHTNRTTEFTNLNE
HSSRSHALLIVTVRGVDCSTGLRTTGKLNLDLAGSERVKGSGAEGSRLREAQHINKSLSALGDVI
AALRSRQGHV PFRNSKLT YLLQDSL S GDSK TLMV-----QVSPVEKNTSETLYSLKFAER---
-----VR*
```

Structure File Example (1SDMA.atm)

ATOM	1	N	LYS	1	-3.978	26.298	113.043	1.00	31.75	N
ATOM	2	CA	LYS	1	-4.532	25.067	113.678	1.00	31.58	C
ATOM	3	C	LYS	1	-5.805	25.389	114.448	1.00	30.38	C
ATOM	4	O	LYS	1	-6.887	24.945	114.072	1.00	32.68	O
ATOM	5	CB	LYS	1	-3.507	24.446	114.631	1.00	34.97	C
ATOM	6	CG	LYS	1	-3.743	22.970	114.942	1.00	36.49	C
ATOM	7	CD	LYS	1	-3.886	22.172	113.644	1.00	39.52	C
ATOM	8	CE	LYS	1	-3.318	20.766	113.761	1.00	41.58	C
ATOM	9	NZ	LYS	1	-1.817	20.761	113.756	1.00	43.48	N
ATOM	10	N	ILE	2	-5.687	26.161	115.522	1.00	26.16	N
ATOM	11	CA	ILE	2	-6.867	26.500	116.302	1.00	22.75	C
ATOM	12	C	ILE	2	-7.887	27.226	115.439	1.00	21.35	C
ATOM	13	O	ILE	2	-7.565	28.200	114.770	1.00	20.95	O
ATOM	14	CB	ILE	2	-6.513	27.377	117.523	1.00	21.68	C
ATOM	15	CG1	ILE	2	-5.701	26.563	118.526	1.00	21.13	C
ATOM	16	CG2	ILE	2	-7.782	27.875	118.200	1.00	18.96	C
ATOM	17	CD1	ILE	2	-5.368	27.325	119.787	1.00	21.39	C
ATOM	18	N	ARG	3	-9.120	26.737	115.461	1.00	22.04	N
ATOM	19	CA	ARG	3	-10.214	27.327	114.693	1.00	23.95	C
ATOM	20	C	ARG	3	-10.783	28.563	115.400	1.00	22.82	C
ATOM	21	O	ARG	3	-10.771	28.645	116.629	1.00	22.62	O
ATOM	22	CB	ARG	3	-11.327	26.290	114.510	1.00	26.34	C
ATOM	23	CG	ARG	3	-11.351	25.586	113.161	1.00	30.68	C
ATOM	24	CD	ARG	3	-10.004	25.034	112.771	1.00	35.43	C
ATOM	25	NE	ARG	3	-10.104	24.072	111.672	1.00	43.37	N
ATOM	26	CZ	ARG	3	-10.575	24.350	110.458	1.00	46.04	C
ATOM	27	NH1	ARG	3	-10.997	25.572	110.168	1.00	48.68	N
ATOM	28	NH2	ARG	3	-10.627	23.400	109.532	1.00	48.37	N
ATOM	29	N	VAL	4	-11.278	29.524	114.630	1.00	20.49	N
ATOM	30	CA	VAL	4	-11.853	30.724	115.225	1.00	17.59	C
ATOM	31	C	VAL	4	-13.082	31.211	114.471	1.00	18.31	C
ATOM	32	O	VAL	4	-13.030	31.446	113.264	1.00	16.37	O
ATOM	33	CB	VAL	4	-10.834	31.872	115.272	1.00	19.94	C
ATOM	34	CG1	VAL	4	-11.512	33.168	115.759	1.00	15.64	C
ATOM	35	CG2	VAL	4	-9.668	31.489	116.168	1.00	15.45	C

Modeller Python Script (bioinfo.py)

```
# Homology modelling by the automodel class
```

```
from modeller.automodel import * # Load the automodel class
```

```
log.verbose() # request verbose output
```

```
env = environ() # create a new MODELLER environment to build this model in
```

```
# directories for input atom files
```

```
env.io.atom_files_directory = './../atom_files'
```

```
a = automodel(env,
```

```
   alnfile = 'bioinfo.pir', # alignment filename
```

```
    knowns = '1SDMA', # codes of the templates
```

```
    sequence = 'bioinfo') # code of the target
```

```
a.starting_model= 1 # index of the first model
```

```
a.ending_model = 1 # index of the last model
```

```
                # (determines how many models to calculate)
```

```
a.make() # do the actual homology modelling
```

Where to find structure file

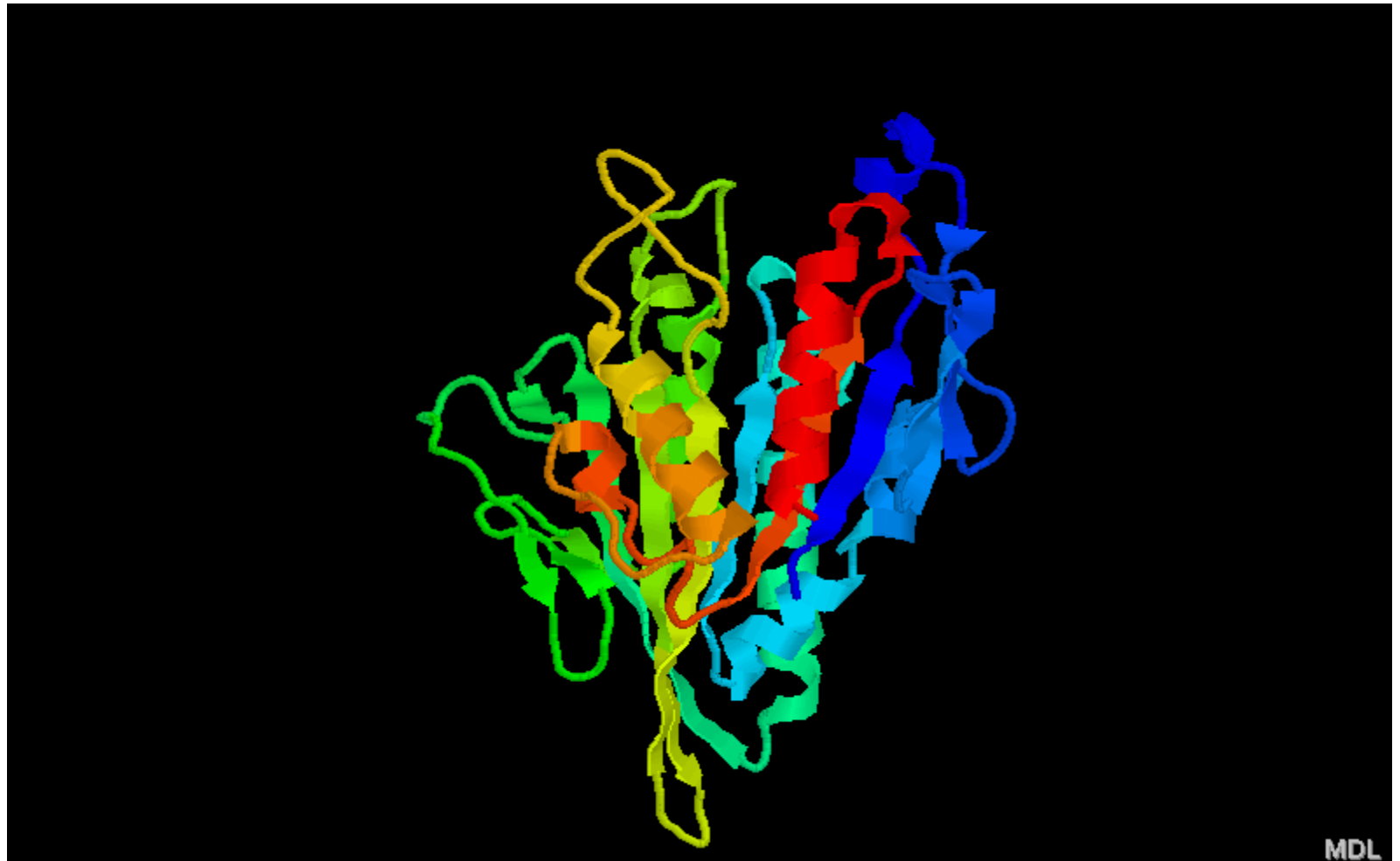
PIR alignment file name

Template structure file id

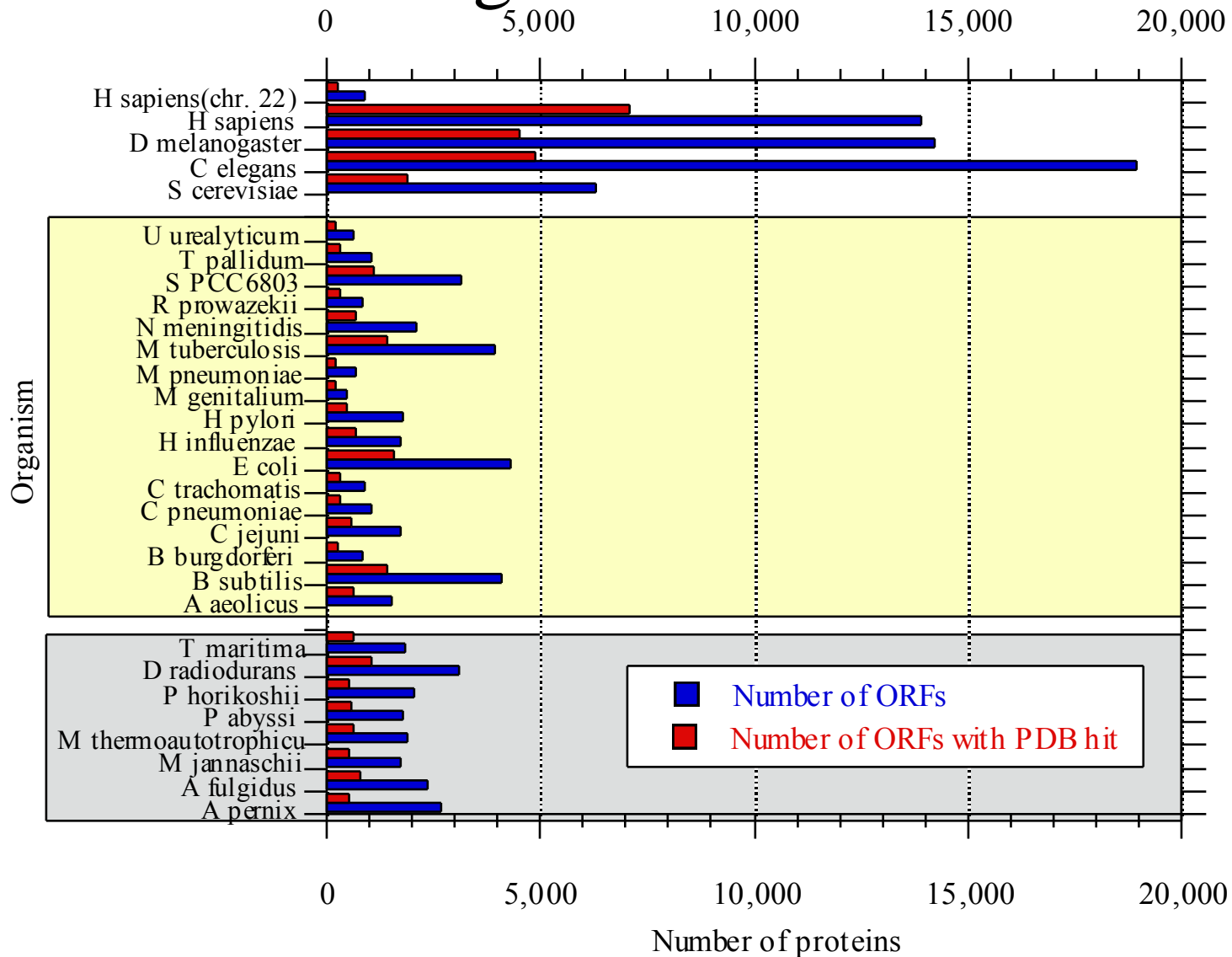
Query sequence id

Output Example

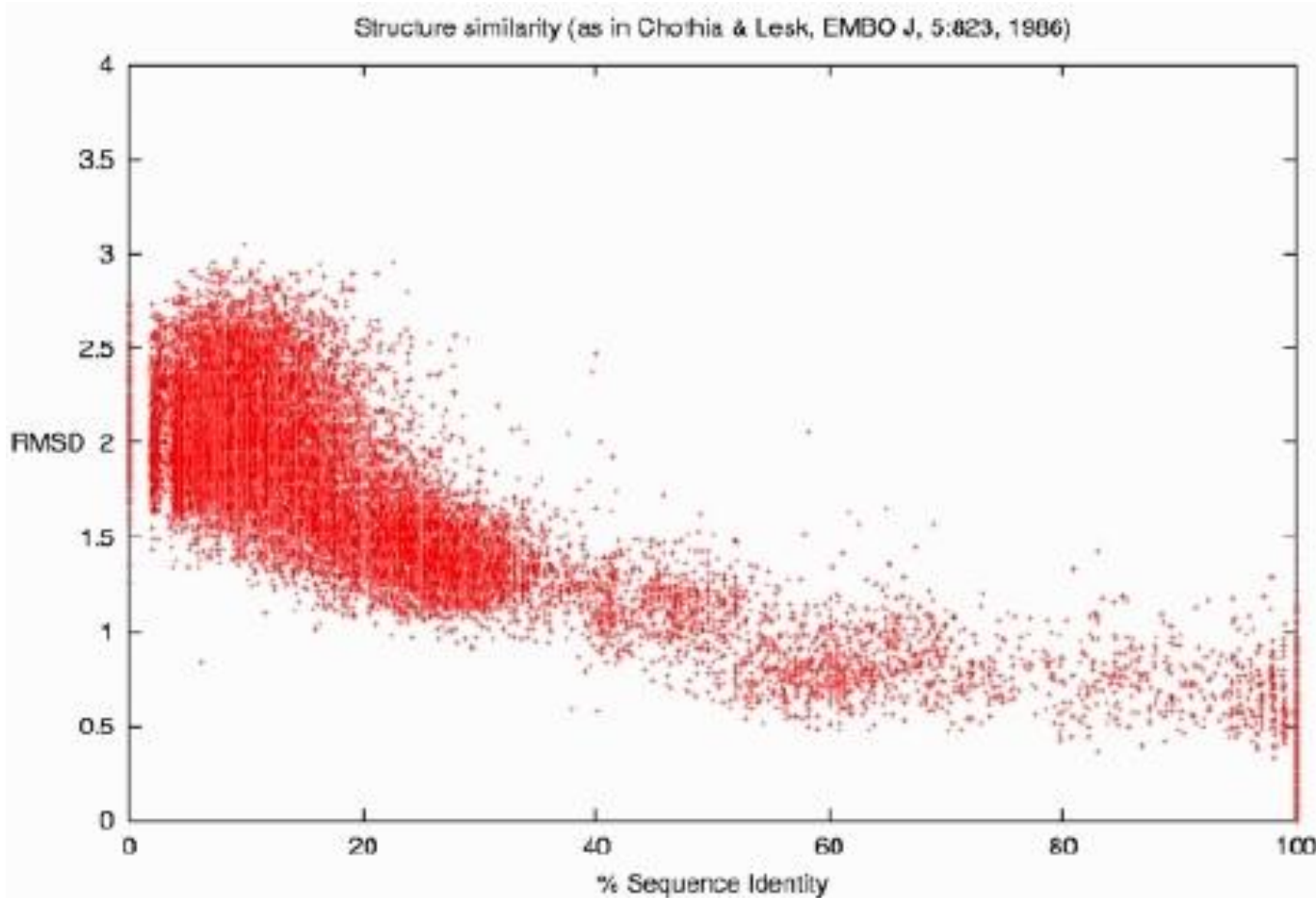
Command: mod8v2 bioinfo.py



Homology modelling for entire genomes



Sequence Identity and Alignment Quality in Structure Prediction



Superimpose
-> RMSD

%Sequence Identity: percent of identical residues in alignment
RMSD: square root of average distance between predicted structure and native structure.

3D Structure Prediction Tools

- MULTICOM (http://sysbio.rnet.missouri.edu/multicom_toolbox/index.html)
- I-TASSER (<http://zhang.bioinformatics.ku.edu/I-TASSER/>)
- HHpred (<http://protevo.eb.tuebingen.mpg.de/toolkit/index.php?view=hhpred>)
- Robetta (<http://robetta.bakerlab.org/>)
- 3D-Jury (<http://bioinfo.pl/Meta/>)
- FFAS (<http://ffas.ljcrf.edu/ffas-cgi/cgi/ffas.pl>)
- Sparks (<http://phyyz4.med.buffalo.edu/hzhou/anonymous-fold-sp3.html>)
- FUGUE (<http://www-cryst.bioc.cam.ac.uk/%7Efugue/prfsearch.html>)
- FOLDpro (<http://mine5.ics.uci.edu:1026/foldpro.html>)
- SAM (<http://www.cse.ucsc.edu/research/compbio/sam.html>)
- Phyre (<http://www.sbg.bio.ic.ac.uk/~phyre/>)
- 3D-PSSM (<http://www.sbg.bio.ic.ac.uk/3dpssm/>)
- mGenThreader (<http://bioinf.cs.ucl.ac.uk/psipred/psiform.html>)

Protein Model Quality Assessment



APOLLO: assessing protein single or multiple model(s) ([help](#))

Evaluating the absolute and/or relative qualities of multiple models or a single model

Upload a compressed file (i.e. zip or tar.gz) containing multiple models OR a single model text file in PDB format: (two multiple models examples: [example.zip](#), [example.tar.gz](#); a single model file example: [example](#))

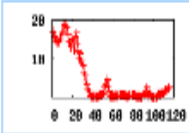
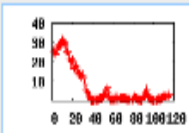
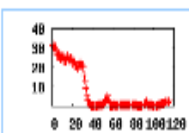
OR paste a single model in PDB format: ([example](#))

(Optional) Email address: (where the evaluation results will be sent to)

(To protect reviewer's anonymity, email account:
bioinformatics.test@gmail.com; password: bioinformatics;)

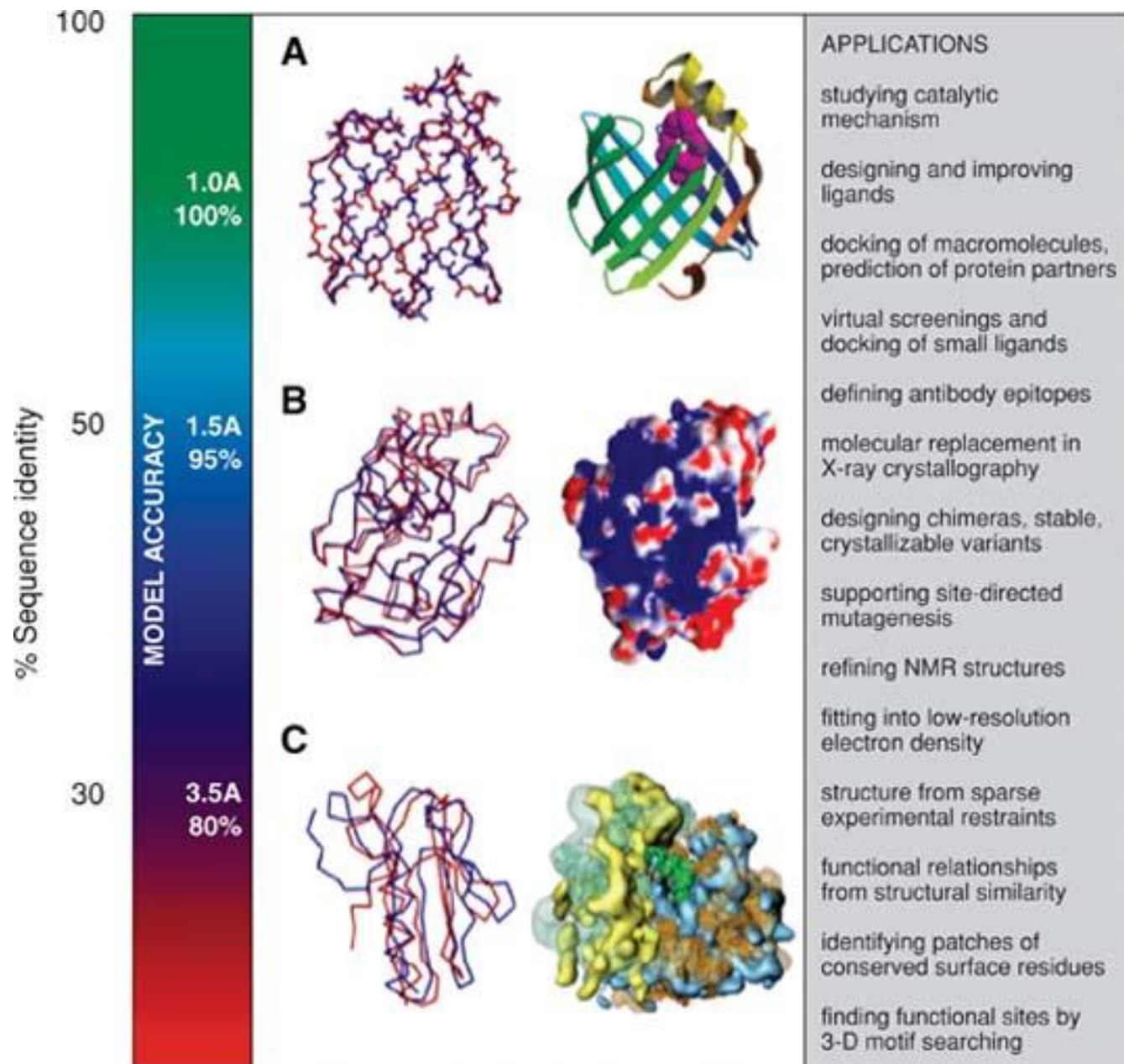
<http://sysbio.rnet.missouri.edu/apollo/>

APOLLO Output

Model Name	Absolute Score	Average Pairwise GDT-TS Score	Refined Average Pairwise Q Score	Local Quality (click to enlarge)
QUARK_TS1	0.713	0.619	0.654	
BAKER-ROSETTASERVER_TS1	0.668	0.503	0.516	
MULTICOM-NOVEL_TS1	0.649	0.638	0.811	

Application of Structure Prediction

- Structure prediction is improving
- Template-based structure become more and more practical. Particularly, comparative / homology modeling is pretty accurate in many cases.
- Comparative modeling has been widely used in drug design.
- Protein structure prediction (both secondary and tertiary) has become an indispensable tool of investigating function of proteins and mechanisms of biological processes.



Baker and Sali (2000)

J. Pevsner, 2005

Project 2

- Select 5 soybean proteins
- Predict 3D structures
- Visualize the structures

(data:http://casp.rnet.missouri.edu/marc/muii_7005/SEQ_TFP_90_2500.txt)