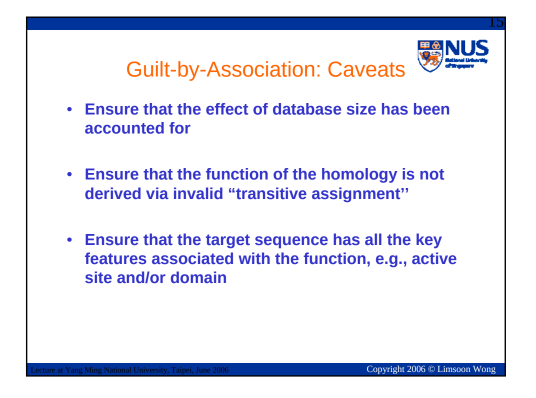
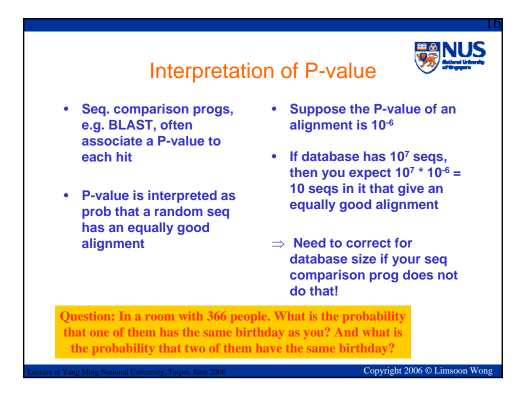
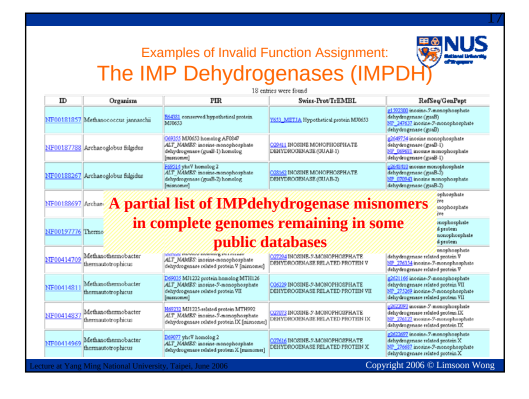
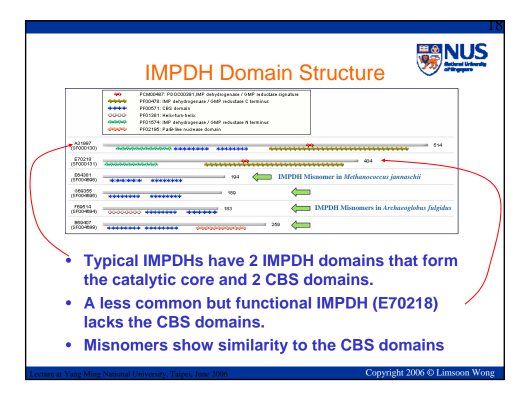


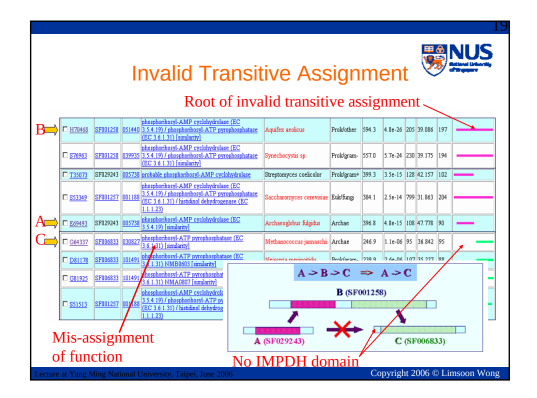
	14
E	xample Alignment with $PTP\alpha$
	632 bits (1629), Expect = e-180 s = 294/302 (97%), Positives = 294/302 (97%)
Query: 1	SPSTNRKYPPLPVDKLEEE INRRMADDNKLFREEFNALPACP IQATCEAASXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
Sbjct: 202	SPSTNRKYPPLPVDKLEEEINRRMADDNKLFREEFNALPACPIQATCEAAS R SPSTNRKYPPLPVDKLEEEINRRMADDNKLFREEFNALPACPIQATCEAASKEENKEKNR 261
Query: 61	YVN ILPYDHSRVHLTPVEGVPD SDY I NASF I NGYQEKNKF I AAQGPKEET VNDFWRMI WE 120 YVN I LPYDHSRVHLTPVEGVPD SDY I NASF I NGYOEKNKF I AAQGPKEET VNDFWRMI WE
Sbjct: 262	YVNILFYDHSRVHLIFYEGYFDSDYINGSFINGYQENNFIAAQGFREEIVNDFWRMIWE YVNILFYDHSRVHLIFYEGYPDSDYINGSFINGYQENNFIAAQGFREETVNDFWRMIWE 321
Query: 121	QNTATIVMVTNLKERKECKCAQYWPDQGCWTYGNVRVSVEDVTVLVDYTVRKFCIQQVGD 180 ONTATIVMVTNLKERKECKCAQYWPDOGCWTYGNVRVSVEDVTVLVDYTVRKFCIOOVGD
Sbjct: 322	QNTATIVMVTNLKERKECKCAQVWPDQGCWTYGNVRVSVEDVTVLVDYTVRKFCIQQVGD 381
Query: 181	VTNRKPQRLITQFHFTSWPDFGVPFTPIGMLKFLKKVKACNPQYAGAIVVHCSAGVGRTG 240 VTNRKPORLITOFHFTSWPDFGVPFTPIGMLKFLKKVKACNPQYAGAIVVHCSAGVGRTG
Sbjct: 382	VINRKPORLITOFHFTSWPDFGVFFTFIGMLKFLKKVKACNPQYAGAIVVHCSAGVGRTG 441
Query: 241	TFVVIDAMLDMMHSERKVDVYGFVSRIRAQRCQMVQTDMQYVFIYQALLEHYLYGDTELE 300 TFVVIDAMLDMMHSERKVDVYGFVSRIRAQRCOMVOTDMOYVFIYQALLEHYLYGDTELE
Sbjct: 442	TFVVTDAMLDMMHSERVVDVTGFVSRTRAQRCQMVQTDMQYFTTQALLEHYLYGDTELE TFVVTDAMLDMMHSERKVDVYGFVSRTRAQRCQMVQTDMQYVFTYQALLEHYLYGDTELE 501
Lecture at Yang Ming National V	University, Taipei, June 2006 © Limsoon Wong



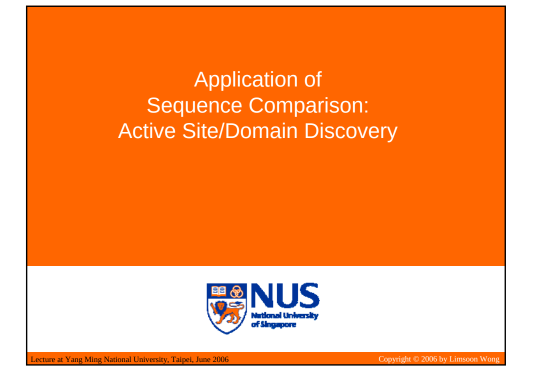


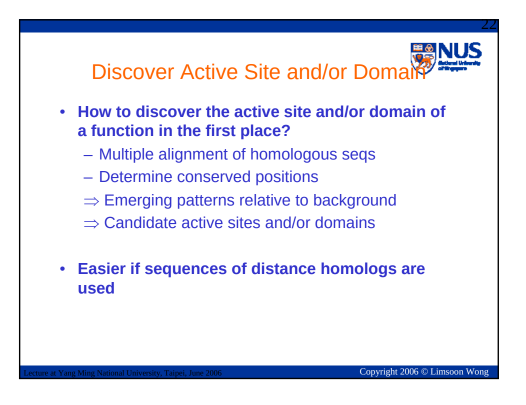






Emerging Pattern
Typical IMPDH Functional IMPDH w/o CBS
Image: PCM00497: PD D000391 J.MP dehydrogenase / GMP reductase signature Image: PF00477: ISB dehydrogenase / GMP reductase C terminus Image: Pr0077: ISB domain Image: PF0077: ISB domain Image: PF0077: ISB domain Image: PF0077: ISB domain Image: PF0178: ISB
A31997 (\$F000130) // /////////////////////////////////
Import Import<
(SF004966) • • • • • • • • • • • • • • • • • •
860407 (\$F004899) ***********************************
 Most IMPDHs have 2 IMPDH and 2 CBS domains Some IMPDH (E70218) lacks CBS domains ⇒ IMPDH domain is the emerging pattern
at Yang Ming National University, Taipei, June 2006 Copyright 2006 © Limsoon Wor





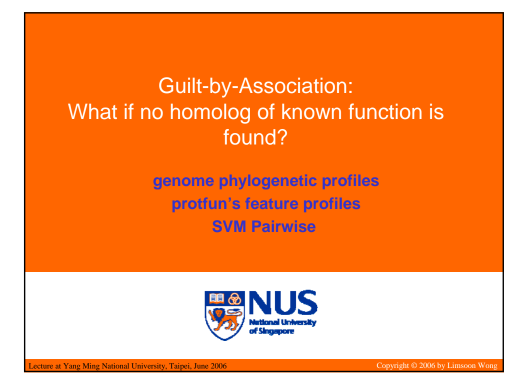
Multiple Alignment of PTPs

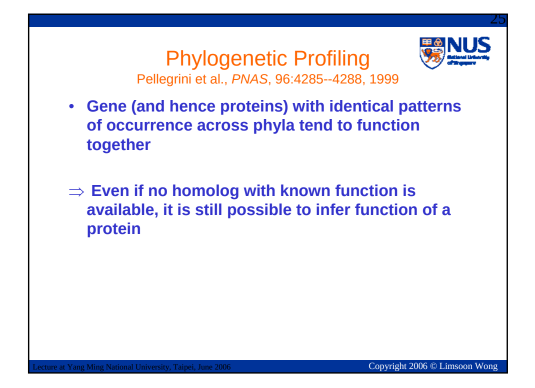


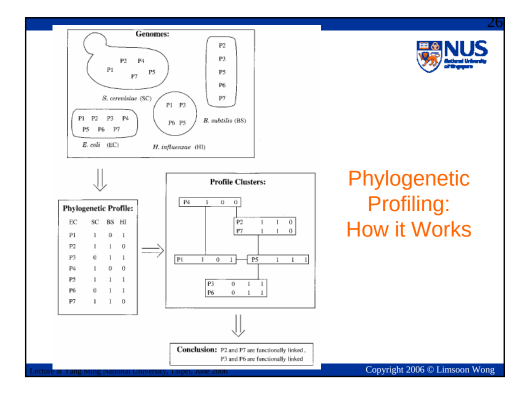
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FHFTSWPDFGVPFTPIGMLKFLKKVKACNP--QYAGAIVVHCSAGVGRTGTFVVIDAMLD gi|126467| gi|2499753 FHFTGWPDHGVPYHATGLLSFIRRVKLSNP--PSAGPIVVHCSAGAGRTGCYIVIDIMLD YHYTQWPDMGVPEYALPVLTFVRRSSAARM--PETGPVLVHCSAGVGRTGTYIVIDSMLQ gi|462550| gi|2499751 FHFTSWPDHGVPDTTDLLINFRYLVRDYMKQSPPESPILVHCSAGVGRTGTFIAIDRLIY gi|1709906 FOF TAUPDHGVPEHPTPFLAFLRRVKTCNP--PDAGPMVVHCSAGVGRTGCF IVIDAMLE LHFTSWPDFGVPFTPIGMLKFLKKVKTLNP--VHAGPIVVHCSAGVGRTGTFIVIDAMMA gi|126471| gi|548626| ${\tt FHFTGWPDHGVPYHATGLLSFIRRVKLSNP--PSAGPIVVHCSAGAGRTGCYIVIDIMLD}$ gi|131570| ${\tt FHFT} GWPD {\tt H} GVPY {\tt H} {\tt ATGLL} GFV {\tt R} QVKS {\tt KSP--PNAGPLVV} {\tt H} CSAGAGRTGCFIVIDIMLD$ gi|2144715 ${\tt fhftswpdhgvpdttdllinfrylvrdymkqsppespilvhcsagvgrtqtfiaidrliy}$..* *** *** . * ...****** ****.... ** ...

- Notice the PTPs agree with each other on some positions more than other positions
- These positions are more impt wrt PTPs
- Else they wouldn't be conserved by evolution
- \Rightarrow They are candidate active sites







Phylogenetic Profiling: P-value

The probability of observing by chance z occurrences of genes X and Y in a set of N lineages, given that X occurs in x lineages and Y in y lineages is

$$P(z|N, x, y) = \frac{w_z * \overline{w_z}}{W}$$

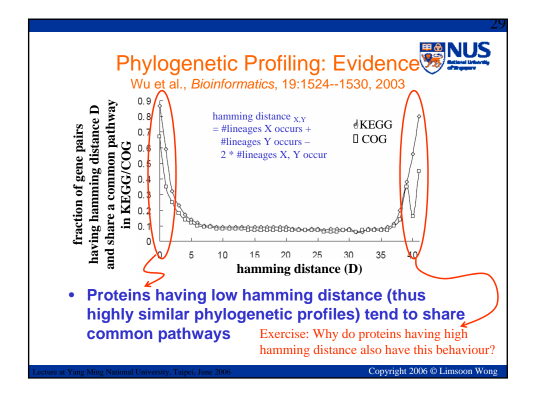
where

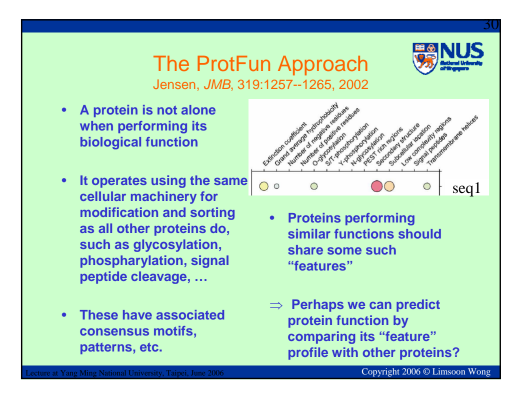
$$w_{z} = \binom{N}{z}$$
No. of ways to distribute z
co-occurrences over N
lineage's
$$\overline{w_{z}} = \binom{N-z}{x-z} * \binom{N-z}{y-z}$$
No. of ways to distribute
the remaining x - z and y - z
occurrences over the remaining
N - z lineage's
$$W_{z} = \binom{N}{x} * \binom{N}{y} + \underbrace{N_{z}}_{over N \text{ lineage's}}$$
No. of ways of
distributing X and Y
over N lineage's
without restriction

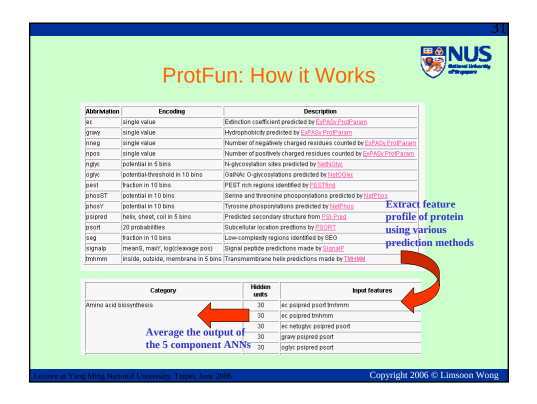
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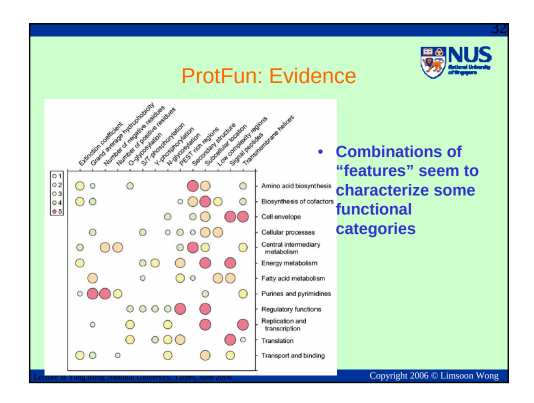
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Pellegrini et al., PNAS,		Evider -4288, 199	
Keyword	No. of non- homologous proteins in group	No. neighbors in keyword group	
Ribosome	60	197	27
Transcription	36	17	10
tRNA synthese and ligase	26	11	5
Membrane proteins*	25	89	5
Flagellar	21	89	3
Iron, ferric, and ferritin	19	31	2 2
Galactose metabolism	18	31	2
Molybdoterin and Molybdenum,			
and molybdoterin	12	6	1
Hypothetical [†]	1.084	108,226	8,440









	Prion	A4	TTHY	
Amino acid biosynthesis	0.011	0.011	0.011	• At the seq level,
Biosynthesis of cofactors	0.041	0.161	0.034	· · · · · · · · · · · · · · · · · · ·
Cell envelope	0.146	0.804	0.698	Prion, A4, & TTHY
Cellular processes	0.027	0.027	0.051	are dissimilar
Central intermediary metabolism	0.047	0.139	0.059	
Energy metabolism	0.029	0.023	0.046	
Fatty acid metabolism	0.017	0.017	0.023	DrotEun prodicto
Purines and pyrimidines	0.528	0.417	0.153	 ProtFun predicts
Regulatory functions	0.013 0.020	0.014 0.029	$0.014 \\ 0.040$	them to be cell
Replication and transcription Translation	0.020	0.029	0.040	envelope-related,
Fransport and binding	0.035	0.827	0.032	•
mansport and binding	0.031	0.827	0.812	tranport & binding
Enzyme	0.233	0.367	0.227	
Non-enzyme <	0.767	0.633	0.773	>
Oxidoreductase (EC 1)	0.070	0.024	0.055	This is in agreement
Transferase (EC 2)	0.031	0.208	0.037	w/ known
Hydrolase (EC 3)	0.101	0.090	0.208	
Isomerase (EC 4.–.–.–)	0.020	0.020	0.020	functionality of
Ligase (EC 5)	0.010	0.010	0.010	these proteins
Ligase (EC 5.–.–.–) Lyase (EC 6.–.–.–)	$0.010 \\ 0.017$	$0.010 \\ 0.078$	$0.010 \\ 0.017$	these proteins

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