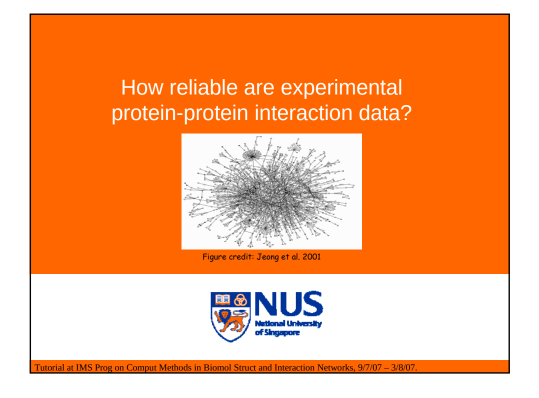
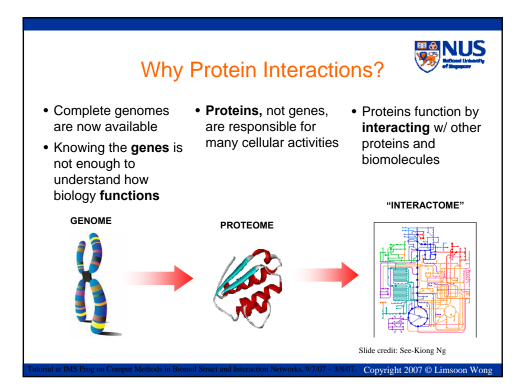
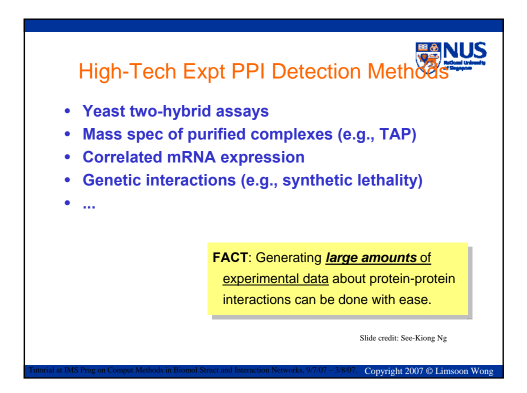
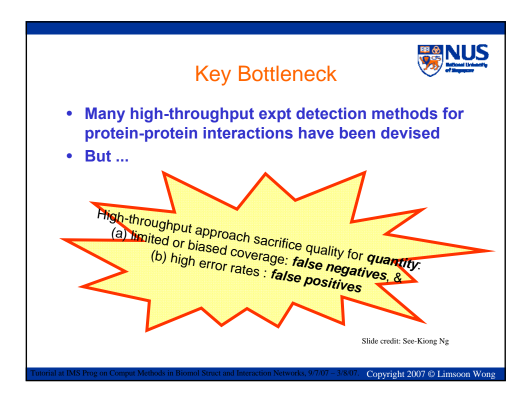


Outline
Reliability of experimental PPI data
Identification of false positives
 Interaction generality
 Interaction generality 2
 Interaction pathway reliability
 – FS Weight
 Meso-scale network motifs
 Identification of false negatives
Uses of (cleansed) PPI data
 Protein function prediction w/o homology info
 Protein complex prediction
Tutorial at IMS Prog on Comput Methods in Biomol Struct and Interaction Networks, 9/7/07 – 3/8/07. Copyright 2007 © Limsoon Wong



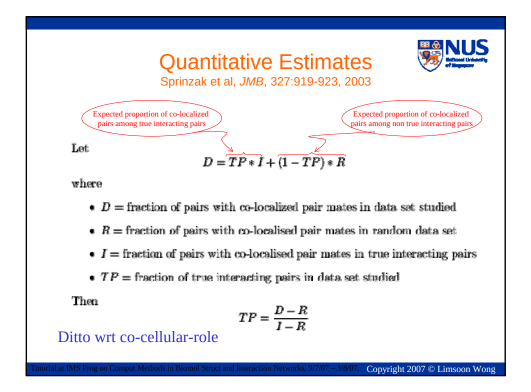


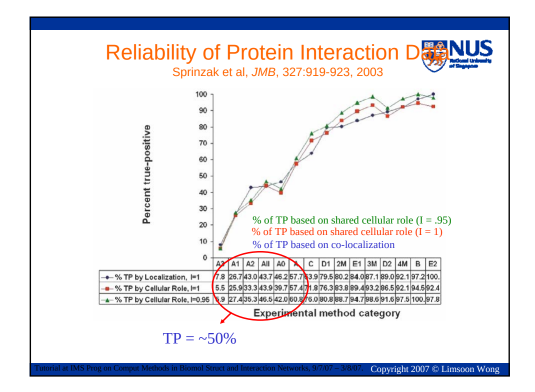




Experimental method category*	Number of interacting pair	rs Co-localization ^b (%)	Co-cellular-role ^b (%)
All: All methods	9347	64	49
A: Small scale Y2H	1861	73	62
A0: GY2H Uetz et al. (published results)	956	66	45
A1: GY2H Uetz et al. (unpublished results)	516	53	33
A2: GY2H Ito et al. (core)	798	64	40
A3: GY2H Ito et al. (all)	3655	41	15
B: Physical methods	71	98	95
C: Genetic methods	1052	77	75
D1: Biochemical, in vitro	614	87	79
D2: Biochemical, chromatography	648	93	88
E1: Immunological, direct	1025	90	90
E2: Immunological, indirect	34	100	93
2M: Two different methods	2360	87	85
3M: Three different methods	1212	92	94
4M: Four different methods	570	95	93

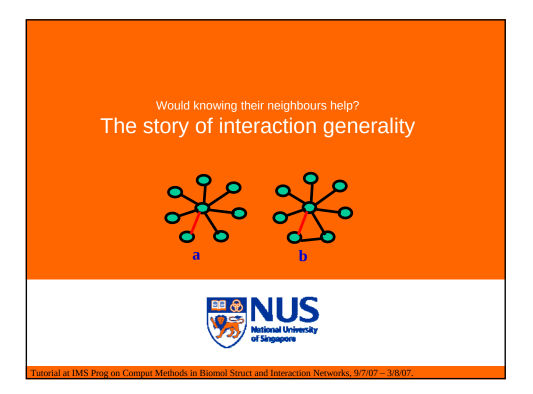
Copyright 2007 © Limsoon Won

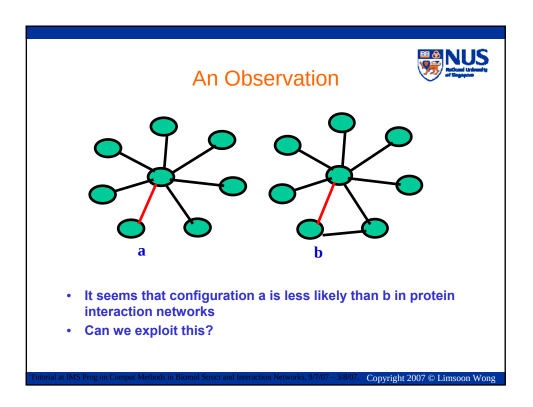


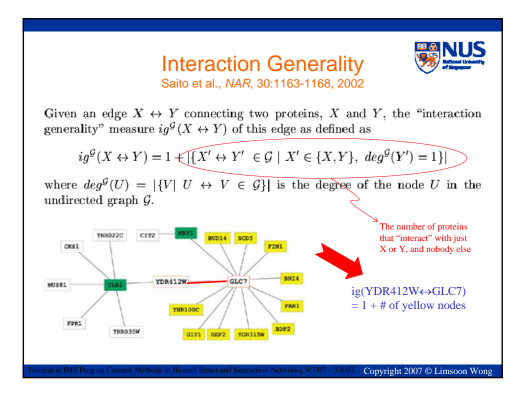


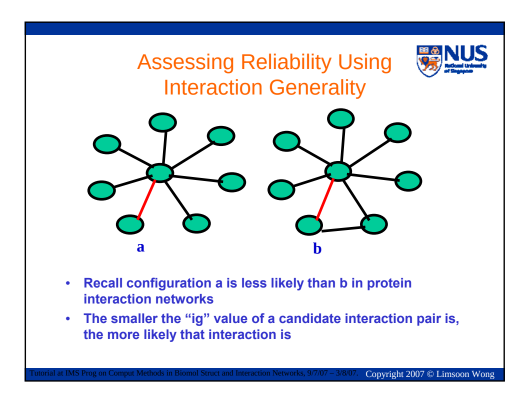
A	Are We There Yet?			
	Coverage	Data quality		
DNA genome sequence	99% of genome sequence	99.9% correct		
mRNA profiling	80-90% of transcripts represented	90% of spots are good data		
Protein interaction data	<u>10-30%</u> of interactions catalogued	50-70% of interactions are spurious		

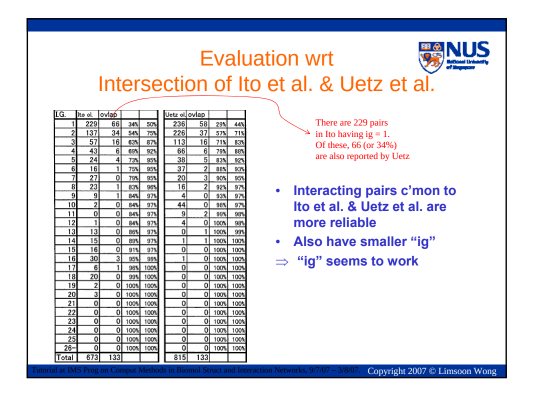


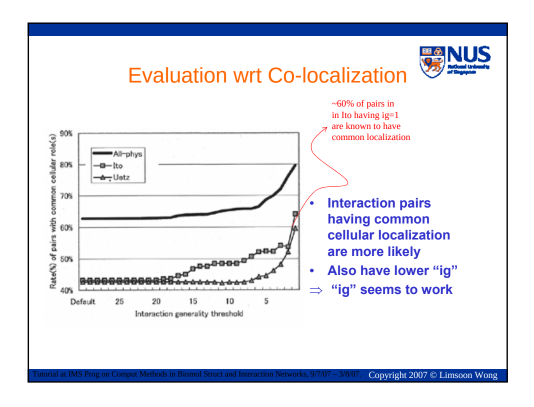


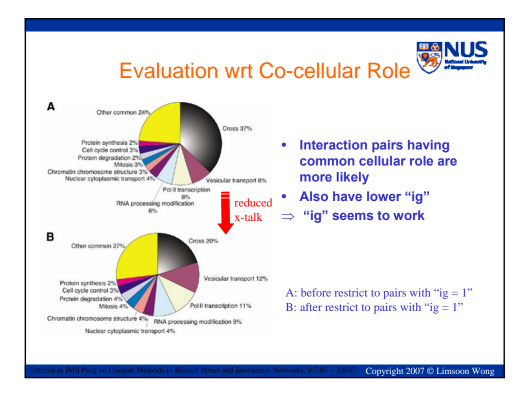


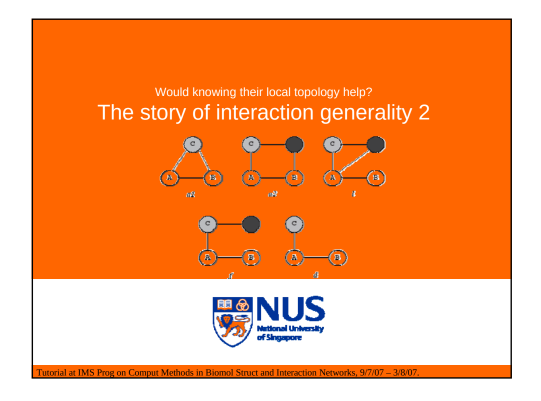


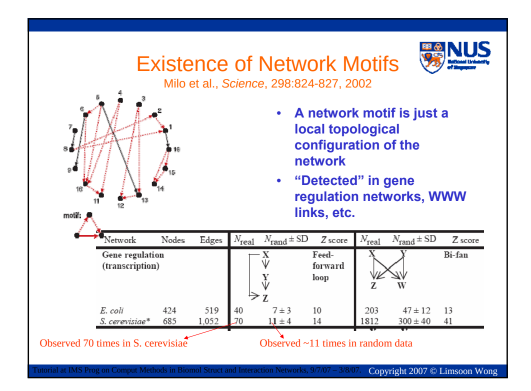


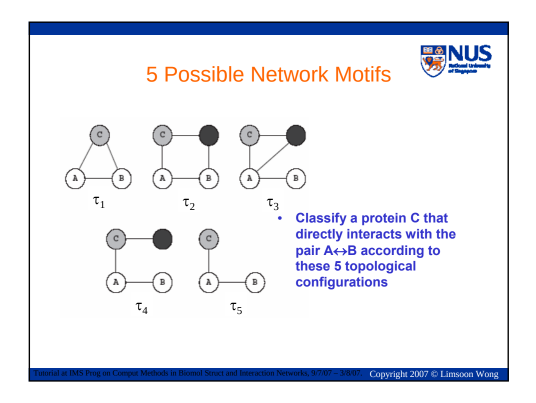


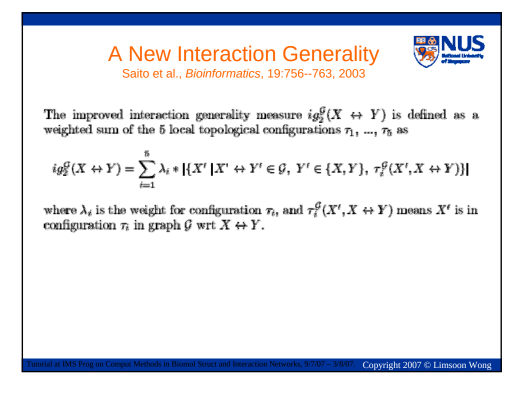


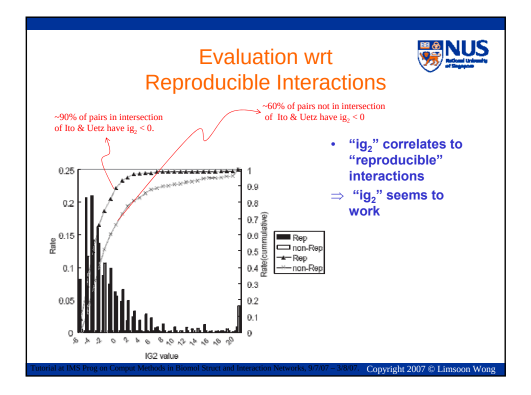


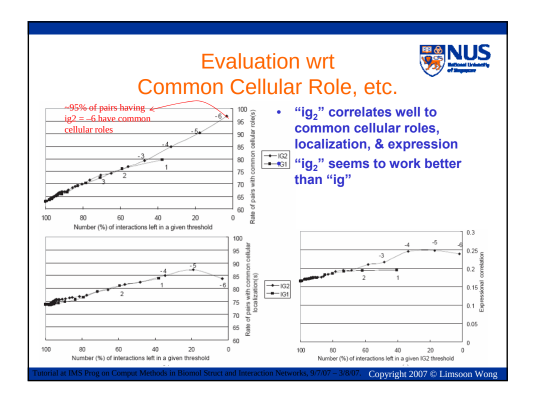


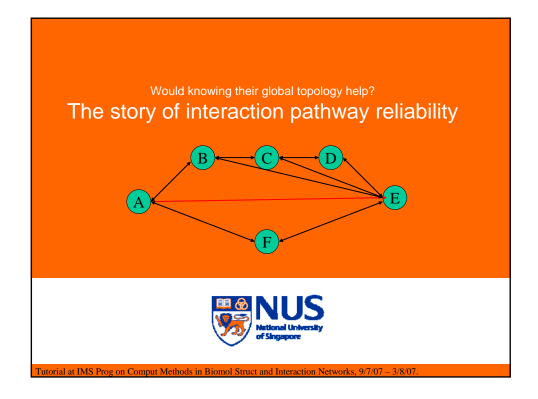


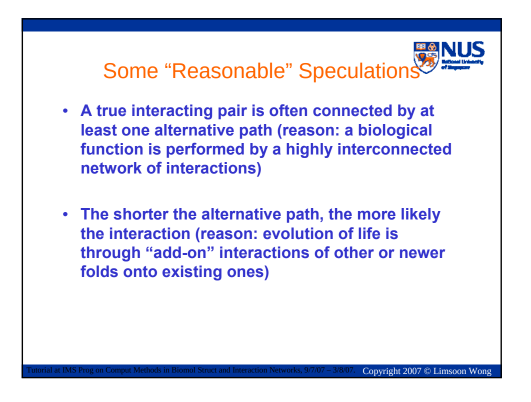


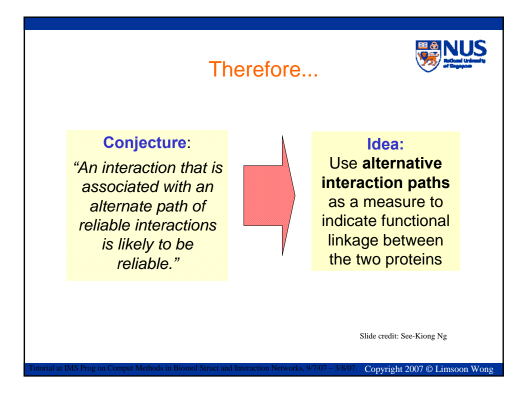


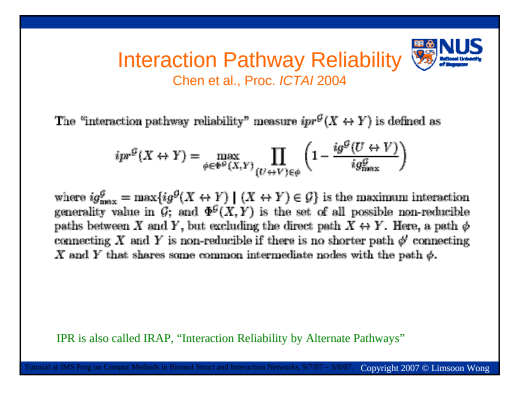


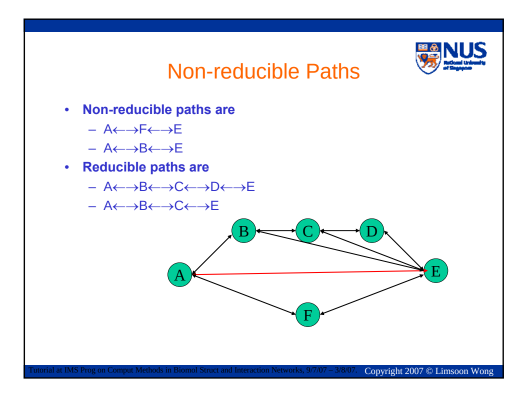


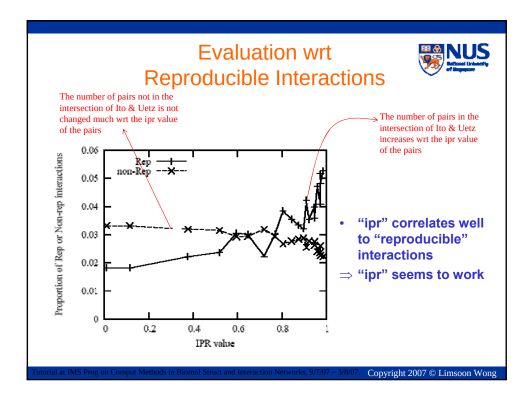


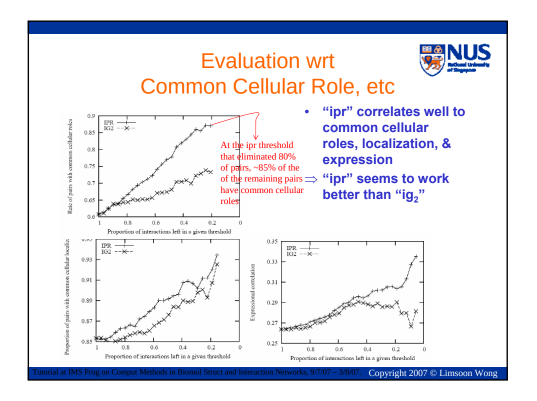


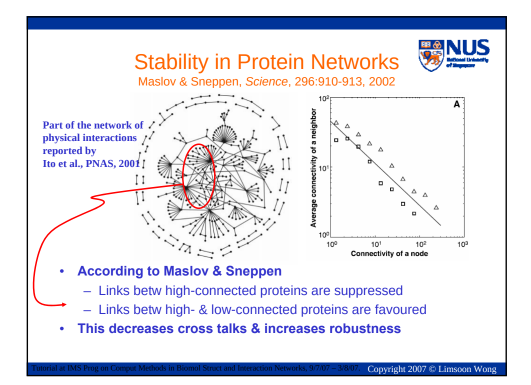


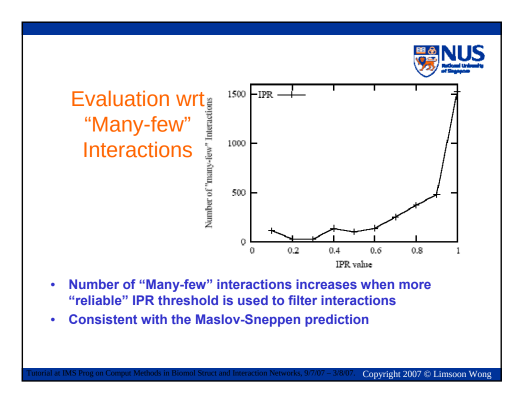


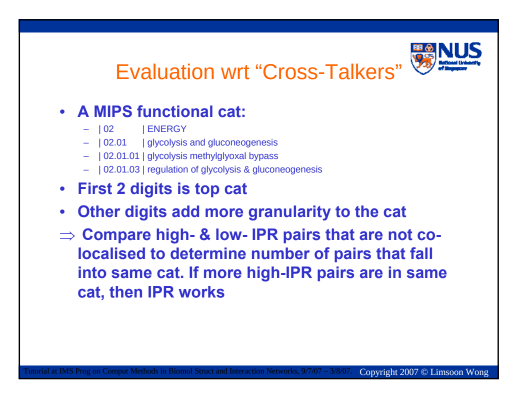


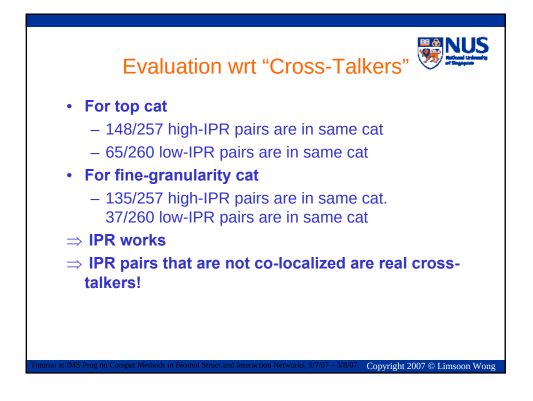




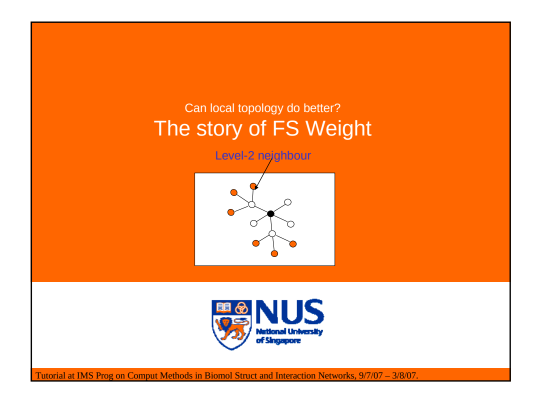


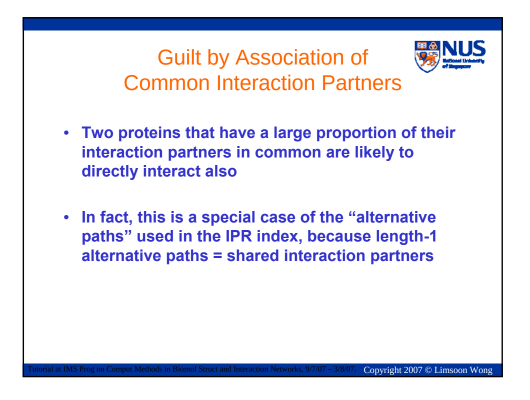


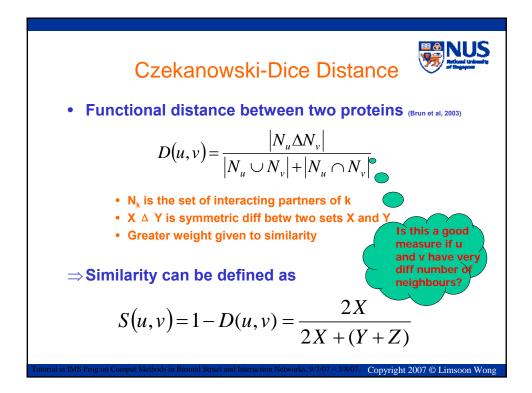


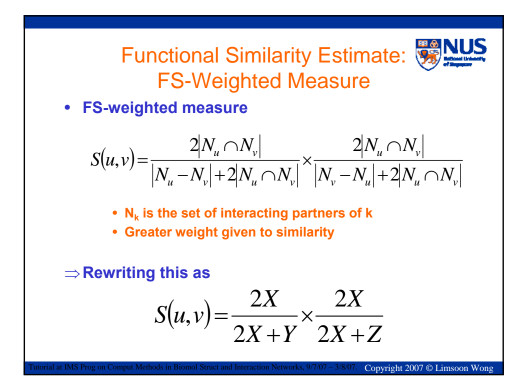


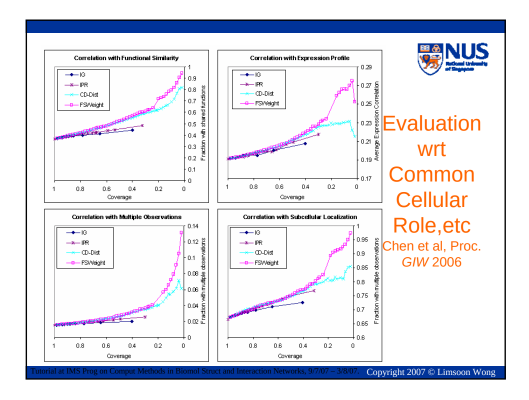
ProteinA	Cellular Localization	ProteinB	Cellular Localization	Functional Pathway
YDR299w	nucleolus-protein transport	YLR208w	cytoplasm-release of transport vesicles from ER	Vesicular transport (Golgi network)
YOL018c	endosome, ER- syntaxin SNARE	YMR117c	spindle pole body- spindle pole component	Cellular import
YDL154w	nucleus-recombination	YBR133c	cytoplasm- neg. regulator of kinase	Meiosis and budding
YGL192w	nucleus-put. Adenosine methyltransferase for sporulation	YBR057c	cytoplasm-meiosis potentially in premeiosis DNA synth	Development of asco-basido -zygo spore
YDR299w	nucleolous- protein transport	YPL085w	cytoplasm,ER-veiscle coat protein interacts cytoplasm, with sec23p	both in vesicular transport
YEL013w	vacuole-phosphorylated protein which interacts with Atg13p for cyto to vacuole targeting vacuole targeting	YFL039c	cytoskeleton-actin	Protein targeting and budding

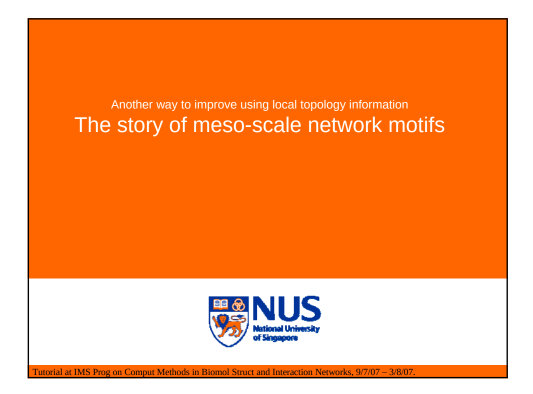


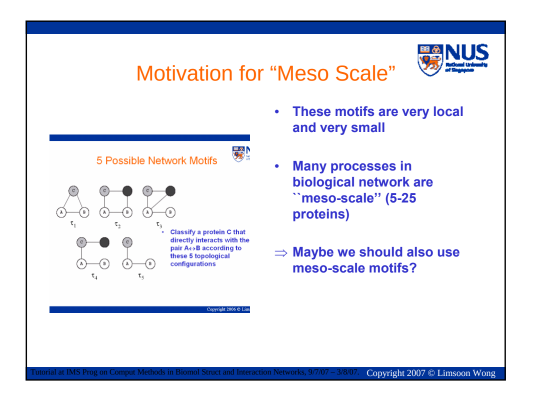


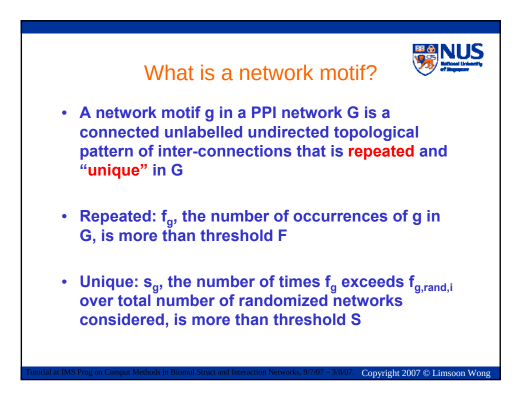


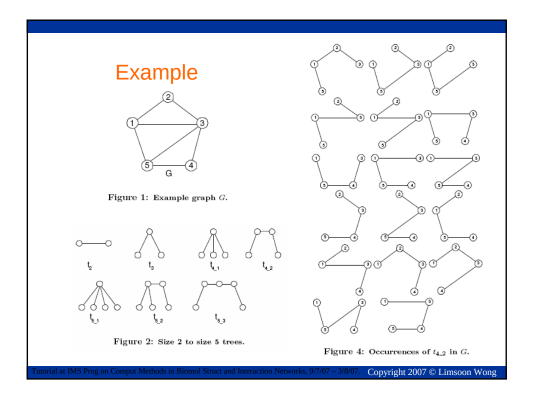


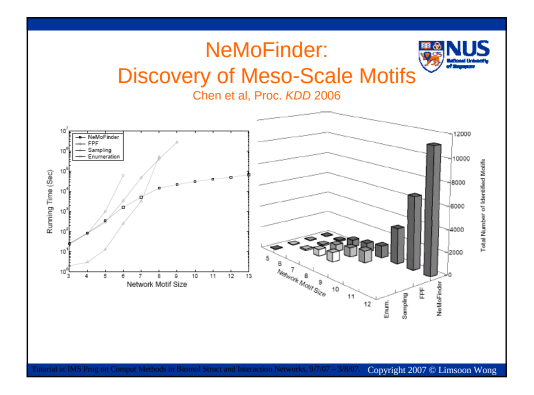


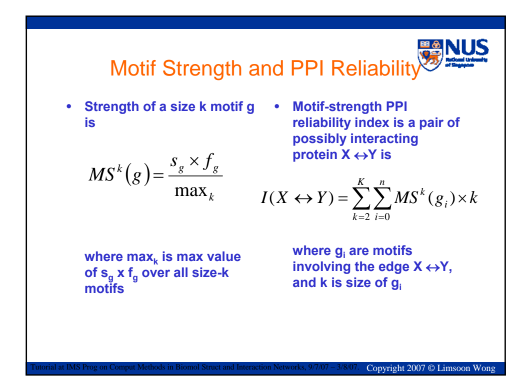


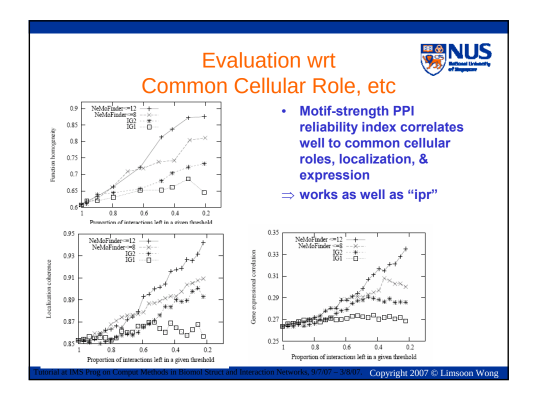


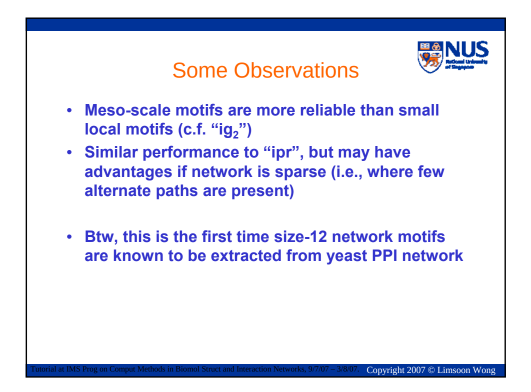


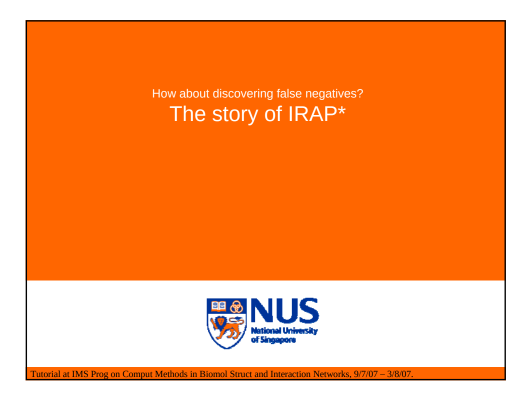


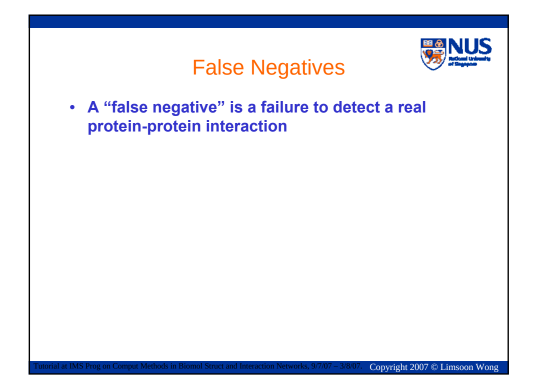


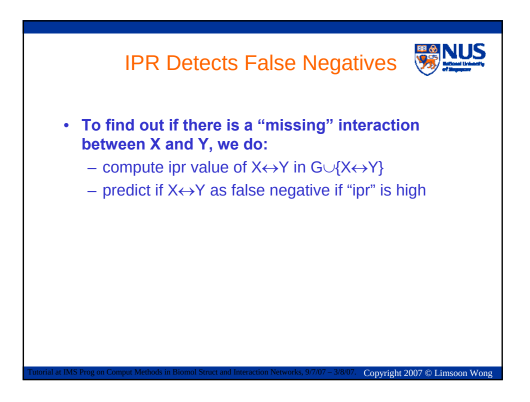


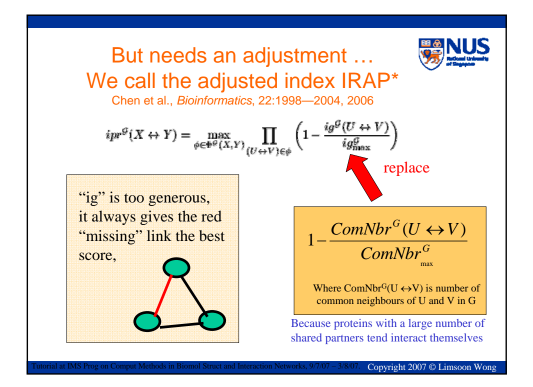


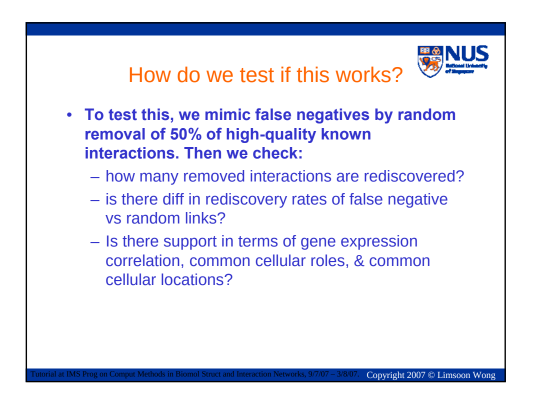


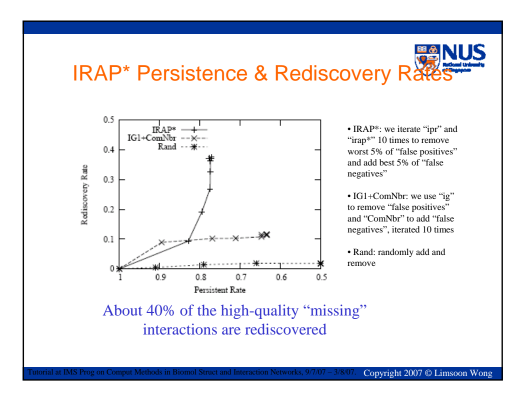


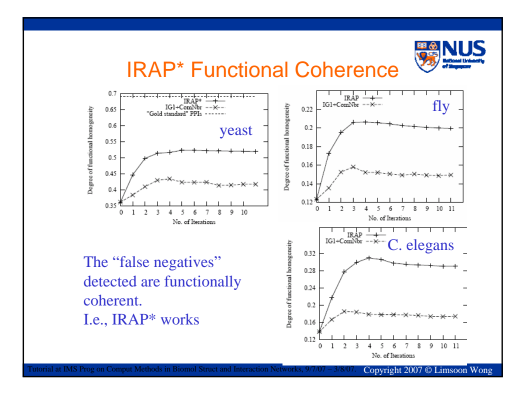










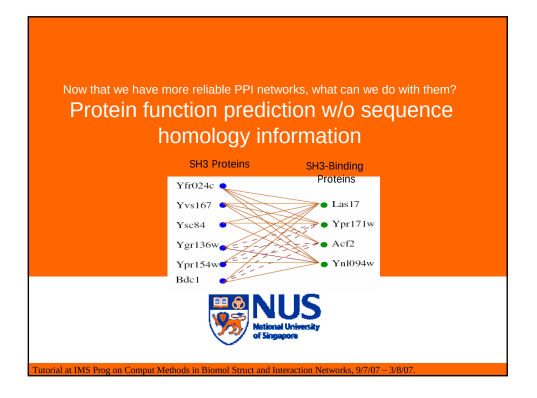


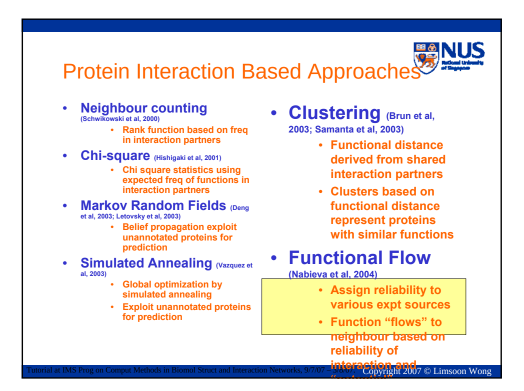


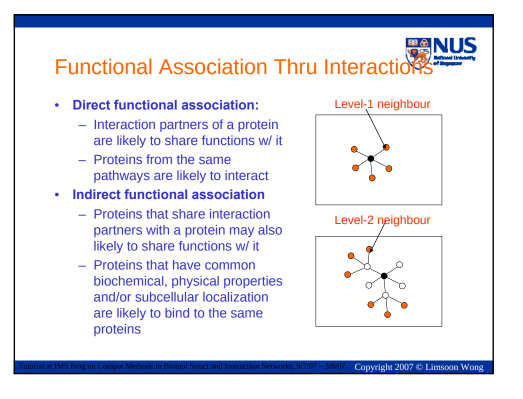
Conclusions

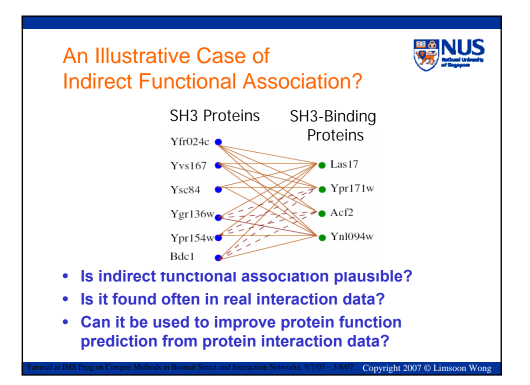
- There are latent local & global network "motifs" that indicate likelihood of protein interactions
- These network "motifs" can be exploited in computational elimination of false positives & false negatives from highthroughput Y2H expt & possibly other highly erroneous interaction data
- IPR & meso-scale motifs are the most effective topologically-based computational measure for assessing the reliability (false positives) of proteinprotein interactions detected by highthroughput methods
- IPR/IRAP* can discover new interactions (false negatives) not detected in the expt PPI network

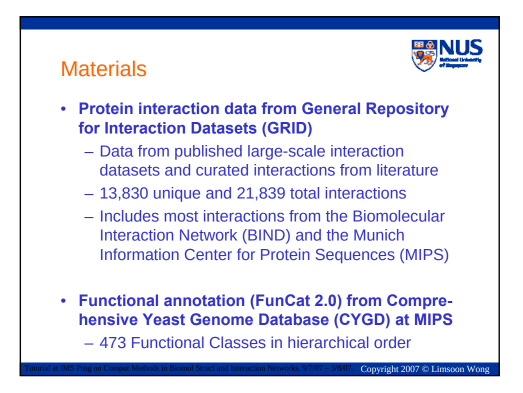
Copyright 2007 © Limsoon Wong

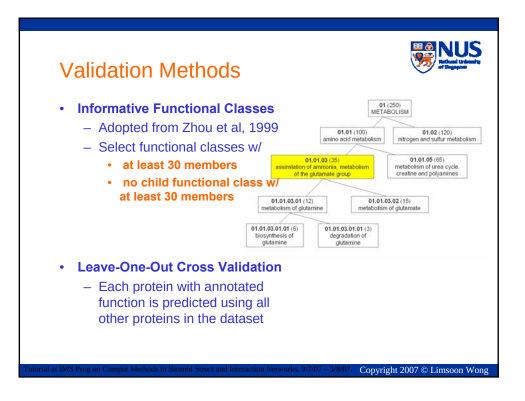


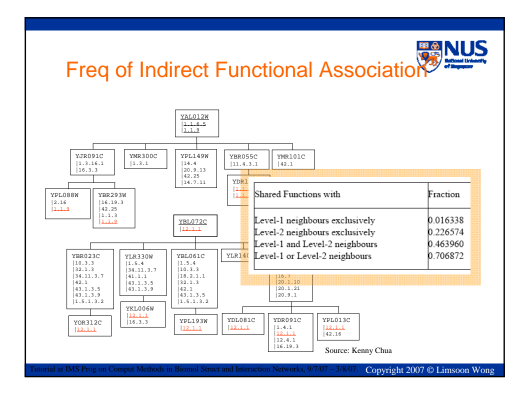


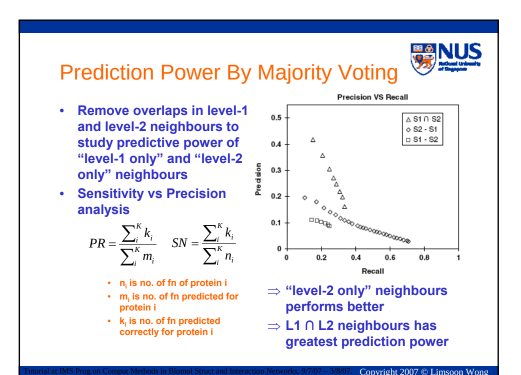


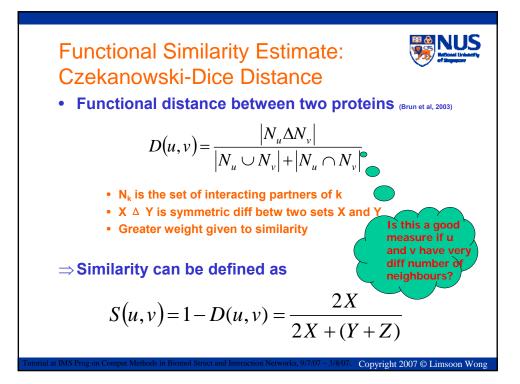


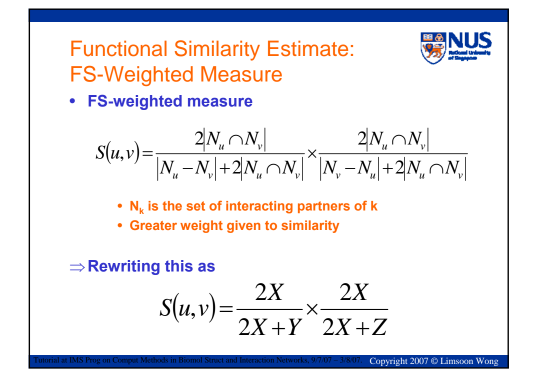


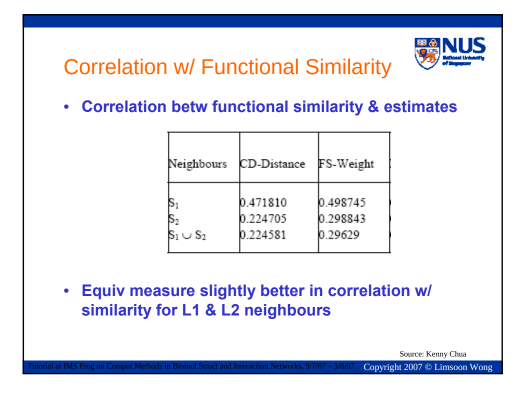














Reliability of Expt Sources

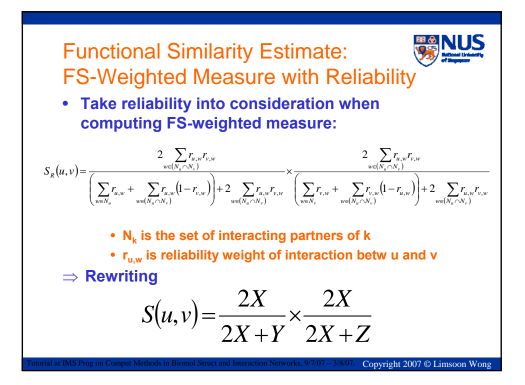
- Diff Expt Sources have diff reliabilities
 - Assign reliability to an interaction based on its expt sources (Nabieva et al, 2004)
- Reliability betw u and v computed by:

$$r_{u,v} = 1 - \prod_{i \in E_{u,v}} (1 - r_i)$$

- r_i is reliability of expt source i,
- E_{u,v} is the set of expt sources in which interaction betw u and v is observed

Source	Reliability
Affinity Chromatography	0.823077
Affinity Precipitation	0.455904
Biochemical Assay	0.666667
Dosage Lethality	0.5
Purified Complex	0.891473
Reconstituted Complex	0.5
Synthetic Lethality	0.37386
Synthetic Rescue	1
Two Hybrid	0.265407

Copyright 2007 © Limsoon We



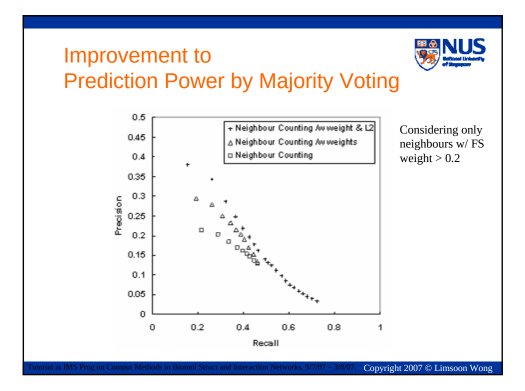


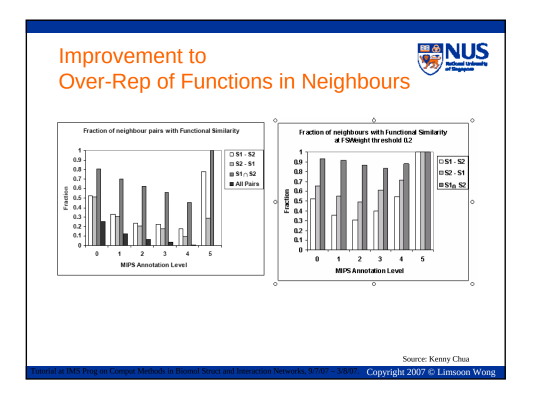
Copyright 2007 © Limsoon Wo

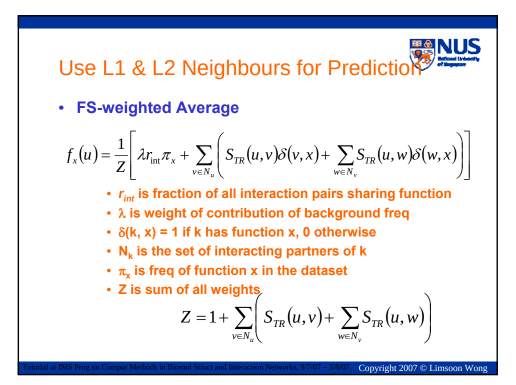
Integrating Reliability

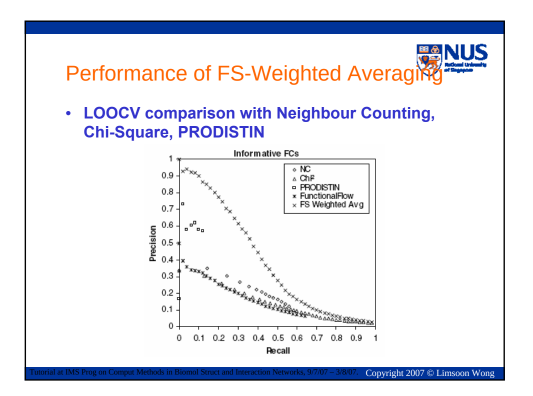
• Equiv measure shows improved correlation w/ functional similarity when reliability of interactions is considered:

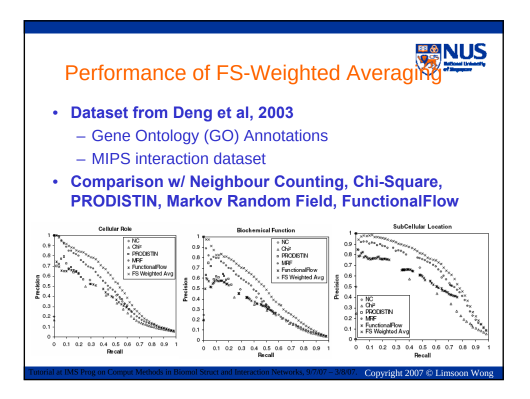
Neighbours	CD-Distance	FS-Weight	FS-Weight R
S 1	0.471810	0.498745	0.532596
S_2	0.224705	0.298843	0.375317
$S_1 \cup S_2$	0.224581	0.29629	0.363025









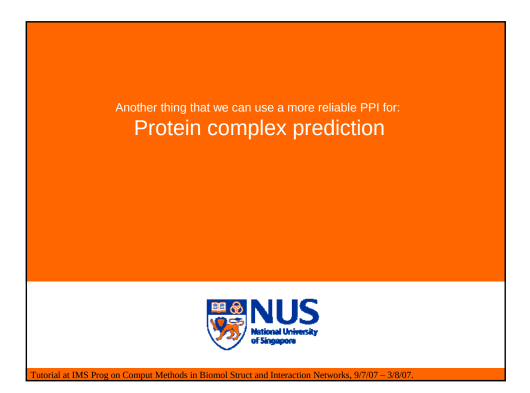




Copyright 2007 © Limsoon Won

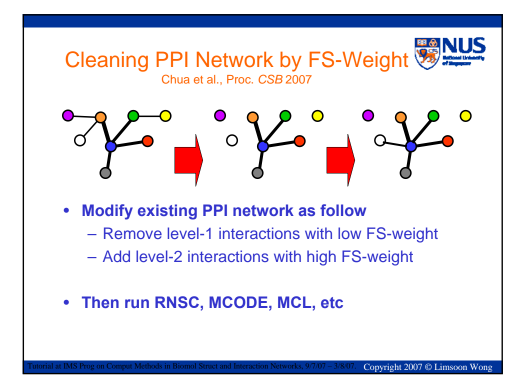
Conclusions

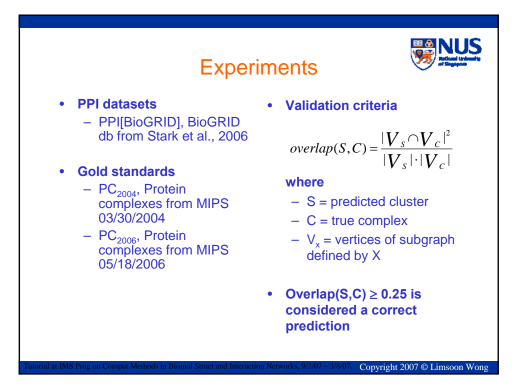
- Indirect functional association is plausible
- It is found often in real interaction data
- It can be used to improve protein function prediction from protein interaction data
- It should be possible to incorporate interaction networks extracted by literature in the inference process within our framework for good benefit

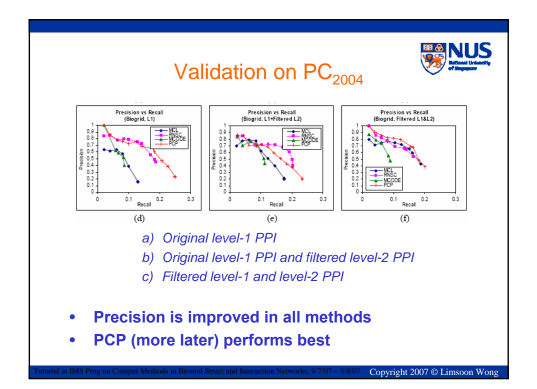


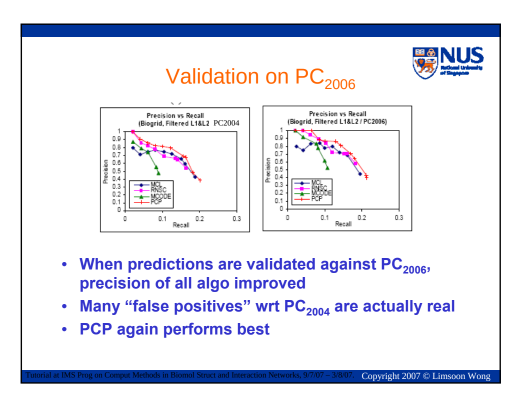
	RNSC	MCODE	MCL
Туре	Clustering, local search cost based	Local neighborhood density search	Flow simulation
Multiple assignment of protein	No	Yes	No
Weighted edge	No	No	Yes

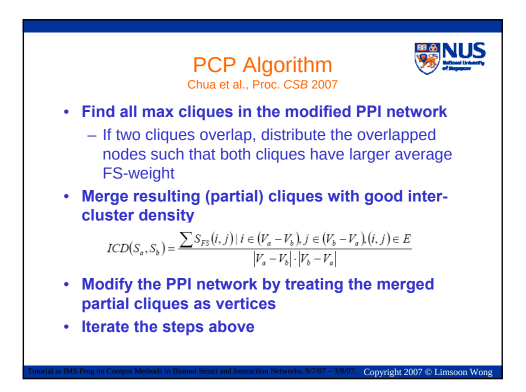
Copyright 2007 © Limsoon Wo

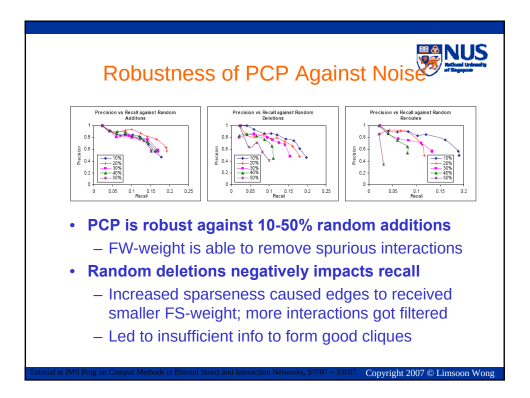


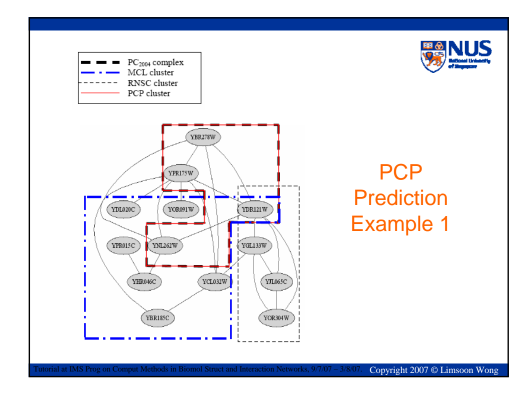


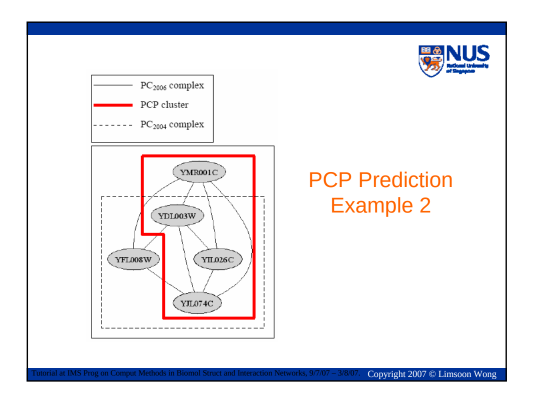


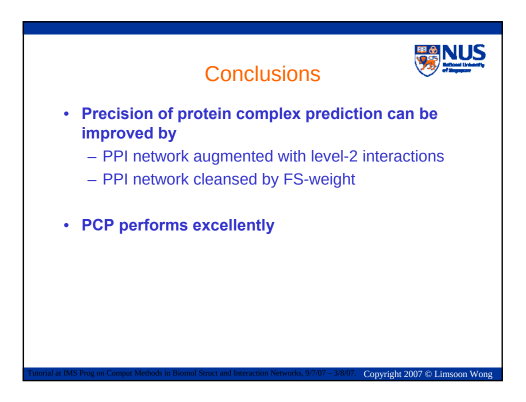










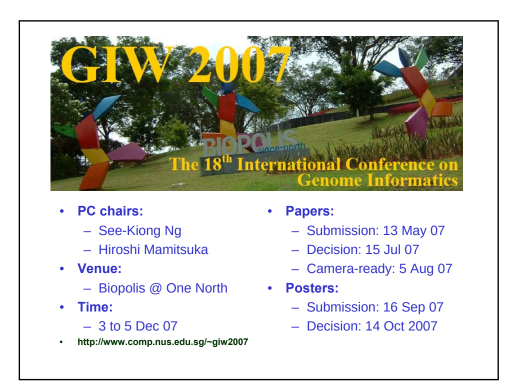




Copyright 2007 © Limsoon Wo

References

- J. Chen et al, "Increasing confidence of protein-protein interactomes", Proc. GIW 2006, pages 284—297
- J. Chen et al, "Systematic assessment of high-throughput protein interaction data using alternative path approach", Proc. ICTAI 2004, pages 368—372
- J. Chen et al, "Towards discovering reliable protein interactions from high-throughput experimental data using network topology", *Artificial Intelligence in Medicine*, 35:37—47, 2005
- J. Chen et al, "Increasing confidence of protein interactomes using network topological metrics", *Bioinformatics*, 22:1998–2004, 2006
- J. Chen et al, "NeMoFinder: Dissecting genome-wide protein-protein interactions with meso-scale network motifs", Proc. KDD 2006
- H.N. Chua et al. "Exploiting indirect neighbours and topological weight to predict protein function from protein-protein interactions", *Bioinformatics*, 22:1623–1630, 2006
- H.N. Chua et al. "Using indirect protein-protein interactions for protein complex prediction",
 Proc. CSB 2007, August 2007, to appear
- R. Saito et al, "Interaction generality, a measurement to assess the reliability of a proteinprotein interaction", NAR 30:1163--1168, 2002
- R. Saito et al, "Construction of reliable protein-protein interaction networks with a new interaction generality measure", *Bioinformatics* 19:756--763, 2003



RECOMB2008: 12th International Conference on Research in Computational Molecular Biology

- Conference Chair:
 Limsoon Wong
- PC chair:
 - Martin Vingron
- Venue:
 - UCC @ NUS
- Time:
 - 30 Mar to 2 Apr 08
- http://www.comp.nus.edu.sg/~recomb08

- Papers:
 - Submission: 24 Sept 07
 - Decision: 10 Dec 07
 - Camera-ready: 18 Jan 08
- Posters:
 - Submission: 14 Jan 08

Copyright 2007 © Limsoon Won

- Decision: 4 Feb 08