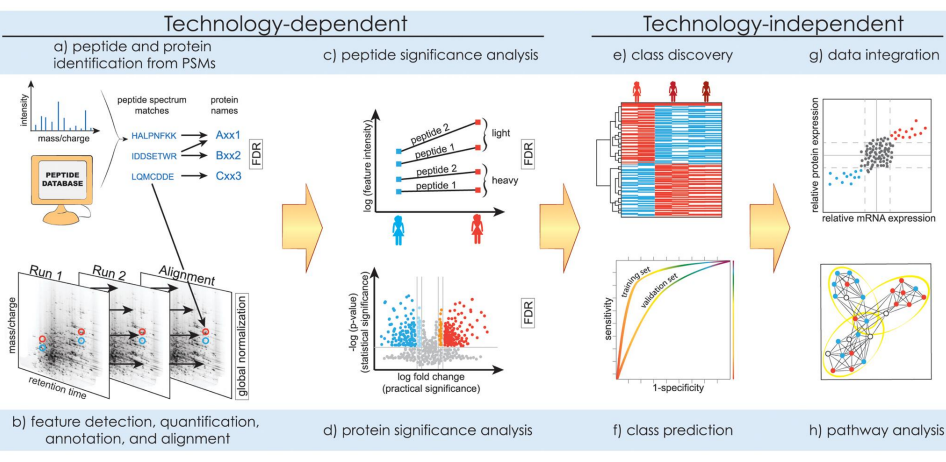
Enabling more sophisticated analysis of proteomic profiles

#### Limsoon Wong (Joint work with Wilson Wen Bin Goh)



## Proteomics is a system-wide characterization of all proteins





Kall and Vitek, PLoS Comput Biol, 7(12): e1002277, 2011

#### Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

#### Proteomics vs transcriptomics



- Proteomic profile
  - Which protein is found in the sample
  - How abundant it is
- Similar to gene expression profile. So typical gene expression profile analysis methods can be applied in theory...

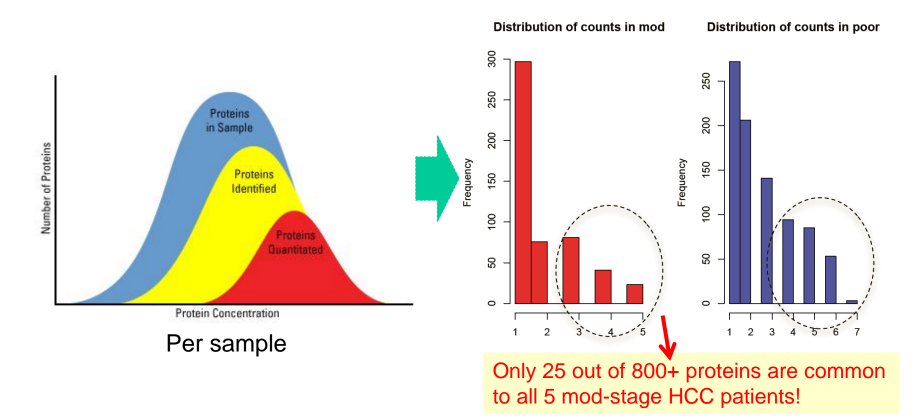
- Key differences
  - Profiling
    - Complexity: 20k genes vs 500k proteins
    - Dynamic range: > 10 orders of magnitude in plasma. Proteins cannot be amplified
  - Analysis
    - Much fewer features
    - Difficult to reproduce
    - Much fewer samples
    - Unstable quantitation

### Issues in proteomics: Coverage and consistency



4

#### **Technical incompleteness** How it affects real data



Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

## Using protein complexes to enhance proteomics: Basic ideas



### A postulate and some math

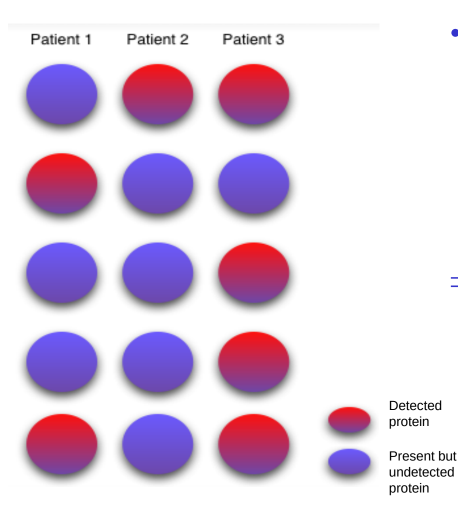


- Postulate: The chance of a protein complex being present in a sample is proportional to the fraction of its constituent proteins being correctly reported in the sample
- Suppose proteomics screen has 75% reliability; a complex comprises proteins A, B, C, D, E; and screen reports A, B, C, D only
- $\Rightarrow$  Complex has 60% (= 0.75 \* 4 / 5) chance to be present
- ⇒ The unreported protein E also has ≥ 60% chance to be present, as presence of the complex implies presence of all its constituents
  ⇒ improving coverage
- ⇒ Each of the reported proteins (A, B, C, and D) individually has 90% (= 100% \* 0.6 + 75% \* 0.4) chance of being true positive, whereas a reported protein that is isolated has a lower 75% chance of being true positive
  - $\Rightarrow$  removing noise

Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

## An intuition





- Suppose the failure to form a protein complex causes a disease
  - If any component protein is missing, the complex can't form
- ⇒ Diff patients suffering from the disease can have a diff protein component missing
  - Construct a profile based on complexes?

#### **Reference complexes**



8

• In this talk, human complexes (of size at least 5) from CORUM are used as reference complexes

 It is possible to use subnets generated from pathway and PPI databases. However these such subnets vary significantly depending on network databases and subnet-generation algo used

So I do not consider these...

Improving coverage in proteomic profiles



## Guo et al. Nature Medicine, 21, 407, 2015 Lots of missing values in real proteomics datasets

Normal

53

-

General

nm.3807-S4.xls [Read-Only] [Compatibility Mode] - Microsoft Excel

Bad

Good

Neutral

Pas	te 🛷 Form	at Painter	B <i>I</i> <u>U</u> ∗	💷 •   🆄	<u>• A</u> •	E <mark>=</mark> ∃	律 律	📲 Merge &	Center *	\$ ~ %	• .0 .00 .00 ≯.0	Condition Formattin	nal Format g ≠ as Table	Check 0	ell E	xplanator	y Inpu	ut	Linked	Cell 🛛	lote	-	Insert Delet	te Format	Clear *	Sort & Filter * 1			
	Clipboard	G.	Fo	nt	Fa		Alignme	nt	Tai	Numb	er 🕞						Styles						Cells	s		Editing			
	X30	•	f <sub>x</sub>	NA																									*
	Α	В	С	D	E	F	G	Н	I.	J	К	L	М	N	0	Р	Q	R	S	Т	U	V	W	Х	Y	Z	AA	AB	AC 🔺
		GeneSy		kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTis	kidneyTi 💻
	protein	mbol	kidneyTisue1	ue2	ue3	ue4	ue5	ue6	ue7	ue8	ue9	ue10	ue11	ue12	ue13	ue14	ue15	ue16	ue17	ue18	ue19	ue20	ue21	ue22	ue23	ue24	ue25	ue26	ue27
2		ACAA1			237958.5					92200.62				NA	222387.2	NA	177211	27857.94				77962.17		23827.06	302761.4		2064.747		122386.3
3	P05166	PCCB	246687.75	70504.27	253890.9	NA				321442.7				139288.5			245595.9	30488.41	221565			65477.99		NA		41974.24			175808.5
4	Q96RP9 Q15417	GFM1 CNN3	37872.59722 28364.89722	NA NA	40359.89 NA	NA NA	73975.35 NA		52272.02		34506.99 10577.49	35176.2		23060.3 33388.93	91995.3 27593.38	NA 40821.22	37735.48	33491.8 24964.95	48208.46 32403	47858.24 NA	39584.44 24907.94	NA 46053.92	67976.03 NA	23631.74 NA	46763.48 25129.86	NA 42948.4		53619.99 26438.35	23207.51
		S100A16	20504.05722 NA	35176.2	NA	66058.39	NA	30674.6	1804.538		10377.49 NA	32524.27 NA	11359.64	NA	18677.58	49821.32 41493.97	12617.18	24904.93	52405 NA	NA	24307.34 NA	36422.79	NA			42546.4 31161.06		20438.33	NA
7	P62820	RAB1A	NA	NA	NA	NA	NA		54417.16		NA	68503.39	NA	NA	NA	NA	NA	NA	NA	NA	32596.28	NA	NA	54839	NA		2064.747	NA	NA
8		PON1	NA			18128.35	NA	33573.36		NA	NA	NA	NA	59432.1	NA		36282.92	16953.34	NA	NA		45107.13	NA	19506.67			109838.9	NA	NA
9	Q9UL46	PSME2	33680.65278	99968.93	59047.33	145114.2	33256.26	141575.7	77962.17	75727.38	64365.04	121022.2	40286.83	114480.8	40567.01	104458.4	42876.78	83666.14	55954.92	62742.03	33768.27	111940.8	59915.42	151558.9	38443.16	113145.5	79024.33	73747.38	40140.37
10	P08237	PFKM	39644.09722	NA	54240.61	NA	136064	NA	1804.538	62845.97	141296.3	100616.3	137596.7	NA	140860.9	NA	96590.73	NA	92823.65	51085.24	155550.8	NA	47697.29	NA	136064	NA	2064.747	58618.05	143381.1
11		CAT	292456.0528	149632.6	239229.2	24964.95	258247.1	220764.4	540115.8	133921.9	284934.5	367784.7	293727.3	179981.9	259314.6	124294.3	204722.1	77070.33	109006.7	136875.9	290924.4	163095.2	237958.5	31389.75	271920.4	227900.3	499422.8	150524.5	294964.3
		CTNNBL1	NA	NA	NA	NA	NA	NA	1804.538	NA	NA	NA	NA	NA	NA	NA	NA		37621.73		NA	NA	NA	NA	NA	NA	2064.747	NA	NA
		C11orf54		77225.75			365975.5					119242.7		263299.1	474797	229655.9	427428	143697	124568		441856.5	74156.41							375463.4
	P31948	STIP1	76018.00556	83236.9			75613.89		98642.34		77709.53	282315.9		122386.3		129969.2		124568	108554.7			92656.4	85600.47				91127.04		122047.2
	O94901 Q99714	SUN1	57623.33889 175372.7444	NA 114480.8	NA	NA 75400.28	72273.86		1804.538 218888	NA 269679.7	NA 170177.4	NA 165385.0	58063.49 202618.2	NA 117389.5	NA 191537	NA 41135.21	NA 196208.5	NA 151044.7	NA 210269.6	NA 294964.3	60013.66 183893	NA	NA 170081.0		71252.19 233372.9	NA	2064.747 196996.8	NA 293727.3	NA 174540.8
	Q15833	STXBP2					16316.33	51400.47 NA	1804.538	203073.7 NA		17309.98		14224.85	12617.18	41155.21 NA	14224.85	9837.458	210205.0	294904.3 5634.228		28846.59			17380.49	NA			13166.66
	P08195	SLC3A2	50797.625				85345.18	NA	1804.538	NA	77850.57	NA	100616.3	NA	76579.02	NA	44010.16		NA	NA	80199.58		72273.86			NA	2064.747		76292.57
	P26038	MSN					241992.3		164343.5					310472.5		393512.7		427428	390317.5			446678.9				423963.5			441856.5
20	P09104	ENO2	NA	144058.2	NA	184650.5	NA	137596.7	126146.3	21831.56	NA	NA	NA	119650.8	NA	404349.8	NA	48438.29	57080.76	NA	NA	151558.9	NA	181096.8	NA	123793.9	2064.747	NA	NA
21	P07148	FABP1	1219163.714	34579.48	861796.3	NA	940142	NA	1804.538	NA	1130692	NA	1057986	NA	789446.1	NA	221565	NA	NA	NA	1162786	32336.43	805128.4	NA	970053.3	NA	2064.747	NA	1300718
22	Q96Q11	TRNT1	NA	NA	NA	NA	NA	NA	1804.538	NA	NA	NA	NA	NA	NA	NA	NA	NA	37098.09	35565.03	NA	NA	NA	NA	NA	NA	2064.747	NA	NA
	015083	ERC2	NA	NA	NA	85740.42	NA		1804.538	NA	83390.33	NA	NA	NA	NA	NA	NA	142306.8	NA	NA	NA	NA	NA	72396.48	NA	NA	2064.747		70213.43
	Q15911	ZFHX3	NA	NA						NA	243050.1	NA	189860.5	NA	NA	NA	NA	457756.2	NA	NA	NA	NA	NA	NA	NA	NA	2064.747		252846.2
	Q9BUR5	APOO	35479.70278	NA			40140.37					54737.36		13642.38		NA	40140.37	NA	NA	10649.17	34436.2	NA	36956.08		47858.24	NA			20057.06
	Q9UJ83 Q8WUM4	HACL1	417999.9306 50008.50556	NA 34991.44	435248.4	NA 50108.55	336790.8 59047.33	227161.7 41611.18			276628.6	INA 134561 7	274264.6	NA 61553.77	317227.1	271920.4 65262.99	336790.8 68597.03	NA 59827.38	NA 73200.35	372485.6 75049.44	446678.9 64108.37	NA 40359.89	390317.5 70903.29	NA 49636.31					333342.7 37386.23
	P53597	SUCLG1		99433.59		94932.09	310472.5				275420.7			101732.7	245595.9	108554.7		89524.72	192915.6		357417.6	96737.9	205171.6						245595.9
	O00186	STXBP3	NA	28468.21	NA	NA	NA	19019.68		NA	NA	NA		21949.83	NA	NA	NA	NA	NA	NA		29005.53	NA	NA	NA		2064.747	NA	NA
	Q8N335	GPD1L	52415.71111	NA	59328.51	NA	54240.61	21949.83		91466.47	45427.61	109273.7	50443.03	NA	52700.48		45502.32	NA	57623.34		54737.36	NA	62380.69	NA			152627.3		49636.31
31	P08621	SNRNP70	48594.65	51791.05	47269.07	86082.28	44306.32	53026.19	1804.538	NA	59432.1	54839	49636.31	60605.33	52477.21	NA	NA	72977.35	74546.25	82242.07	33003.64	60605.33	49636.31	93224.91	NA	56917.54	2064.747	NA	50797.63
	Q969V6	MKL1	NA	91325.89	55954.92	NA	74269.09	80102.57	1804.538	NA	71906.43	NA	NA	152627.3	72497.5	72497.5	89662.88	51690.71	68707.95	41576.85	72021.55	92973.8	NA	NA	NA	88904.66	2064.747	NA	NA
	P08311	CTSG	NA	NA	46154.89	NA	NA	67879.78		NA	53026.19	NA	NA	68927.99	NA	NA	NA	NA		78414.15	NA	NA	46895.88	NA	NA	56514.53		NA	NA
	Q9UKU7	ACAD8			50179.16	NA	64601.65					28070.84	41974.24	NA	41840.21	NA	42678.39	NA			46053.92	NA	49467.07	NA	61900.08	NA			44605.86
	Q86X76	NIT1	75613.88611	NA		63988.55	80199.58				70389.43	NA		75506.47	78547.77	84980.21	76153.19	NA		40935.27		NA	59540.84	70713.02		73278.36			52415.71
	P05162 P23946	LGALS2 CMA1	33491.8 NA	NA NA	35565.03 NA	NA NA	52415.71	36825.06	1804.538 1804.538	23560.07	18592.77	NA NA		72761.18	35479.7	50008.51	24907.94 NA	NA NA			34916.06	NA NA	30730.15	NA NA			2064.747 53240.82	NA NA	25737.06 NA
	P23946 P01834	IGKC	462133.8694	885197.1	692332.5	484624	NA 296507.9	NA 462133.9		NA 319228.4	NA 659554.4		NA 312295.6	NA 52/1995 /	NA 566103.9	NA 692332.5					NA 499422.8	1130692	NA 706520.3		NA 322906.2			NA 310472.5	643593
	P01854 P14868	DARS	12567.36389	110112		136875.9	30209.1			114195.5				182241.6		201171.9		247871.5	161420			114678.3	54839				2064.747		53026.19
		DCTPP1	NA	NA	NA	NA	NA		1804.538			11589.48		27509.79	NA	NA	NA		87070.11		NA	NA	NA	NA	NA		2064.747		NA 🔻
	▶ ► sTa								1																				•
Rea																											100%	Θ	Ū€
																													4.25 DM



27

R

Σ AutoSum

🐺 Fill 🔻

-

Calculation

a 🕜 🗖 🗗 🔀

Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

P

S

🕅 🛃 🧐 🕶 🖓 🖛

👗 Cut

Copy · Paste

File

e

Home

Insert

Calibri

Page Layout

Formulas

× 11

Data

Review

View

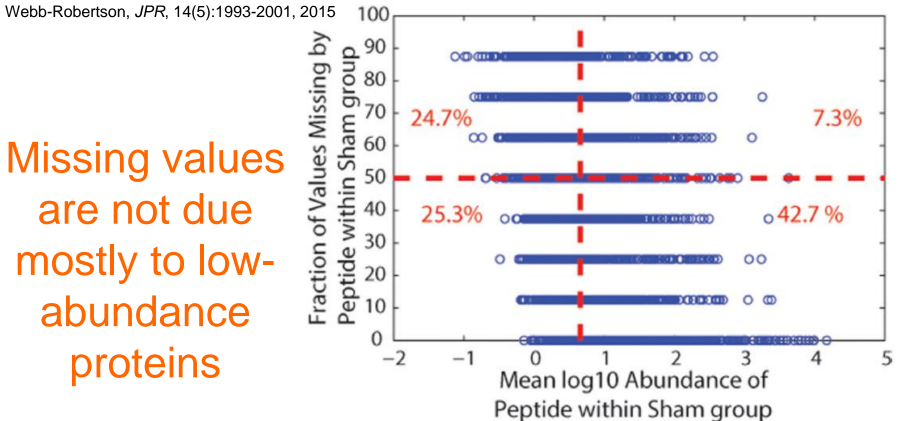
Dr.

Acrobat

🖥 Wrap Text

#### Copyright 2016 © Limsoon Wong

📖 EN 🔺 🍪 🔁 🏴 🕪

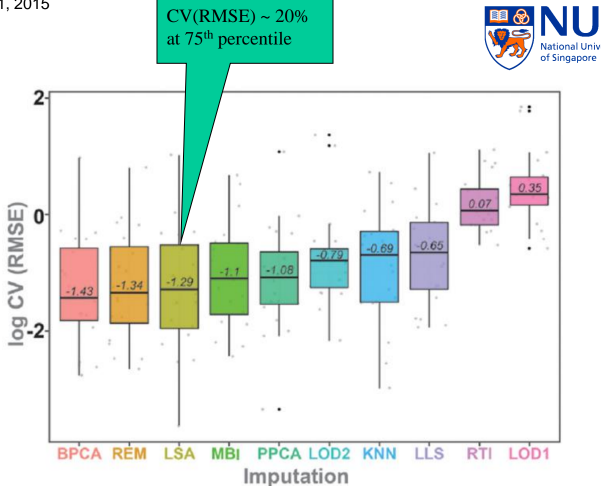


#### Figure 1.

Average log<sub>10</sub> intensity as measured by peptide peak area in the control group versus fraction of missing values and peptide counts associated with bins corresponding to the fraction of missing data comparing phenotypes and exposures for datasets from (A) human plasma and (B) mouse lung. The control group for the human plasma is the normal glucose tolerant (NGT) samples, and the sham group for the mouse lung is the regular weight mice with no lipopolysaccharide (LPS) exposure. The vertical red line represents median average intensity, and the horizontal red line represents the point that 50% of the values are missing.

Webb-Robertson, *JPR*, 14(5):1993-2001, 2015

Current imputation methods don't work very well



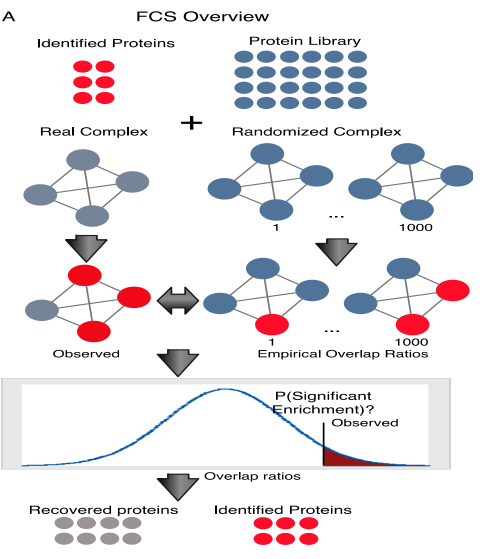
#### Figure 2.

Boxplot of the average  $\log_{10} \text{CV}(\text{RMSE})$  for the imputed dilution series datasets (Table 1) at the (A) peptide and (B) protein levels. The lower line represents the 25th percentile, the upper line of the box represents the 75th percentile, and the inner line corresponds to the median  $\log_{10} \text{CV}(\text{RMSE})$ .





- Rescue undetected proteins from high-scoring protein complexes
- Goh et al. Comparative network-based recovery analysis and proteomic profiling of neurological changes in valporic acid-treated mice. *JPR*, 12(5):2116-2127, 2013



## Other methods for rescuing missing proteins



14

#### • CEA

- Generate cliques from PPIN
- Rescue missing proteins from cliques containing lots of high-confidence proteins
- Li et al. Network-assisted protein identification and data interpretation in shotgun proteomics. *Mol. Syst. Biol.*, 5:303, 2009

#### MaxLink

- Map high-confidence proteins ("seeds") to PPIN
- Rescue proteins that interact many seeds but few non-seeds
- Goh et al. Int J Bioinformatics Research and Applications, 8(3/4):155-170, 2012

• PEP

- Map high-confidence proteins to PPIN
- Extract neighbourhood & predict protein complexes using CFinder
- Rescue undetected proteins from high-ranking predicted complexes
- Goh et al. A Network-based pipeline for analyzing MS data---An application towards liver cancer. *J. Proteome Research*, 10(5):2261-2272, 2011

Goh et al. Comparative network-based recovery analysis and proteomic profiling of neurological changes in valporic acid-treated mice. JPR, 12(5):2116-2127, 2013

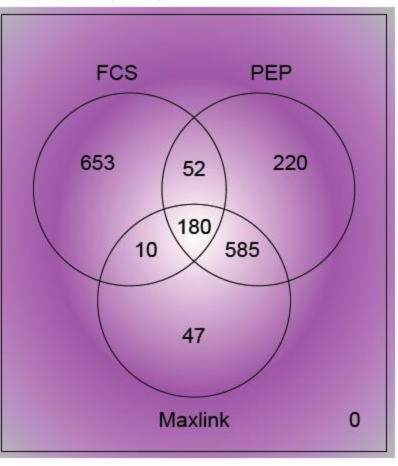
## iTRAQ experiment



- Valporic acid (VPA)-treated mice vs control
  - VPA or vehicle injected every 12 hours into postnatal day-56 adult mice for 2 days
  - Role of VPA in epigenetic remodeling
- MS was scanned against IPI rat db in round #1 - 291 proteins identified
- MS was scanned against UniProtkb in round #2 498 additional proteins identified
- All recovery methods ran on round #1 data and the recovered proteins checked against round #2



Moderate level of agreement of reported proteins between various recovery methods FCS (Real Complexes)



#### **Performance comparison**



17

Method	Novel Suggested Proteins	Recovered proteins	Recall	Precision
PEP	1037	158	0.317	0.152
Maxlink	822	226	0.454	0.275
FCS (predicted)	638	224	0.450	0.351
FCS (complexes)	895	477	0.958	0.533

• Looks like running FCS on real complexes is able to recover more proteins and more accurately

### SWATH experiment



- If there are technical replicates, they should have reported the same proteins. So we can run FCS on one replica, and see whether the predicted missing proteins show up in other replicas
- If there are multiple biological replicates (i.e. patients of the same phenotype), we can run FCS on one of them, and check on the others
- Proteomics data used: Renal cancer
  - Guo et al. Nature Medicine, 21(4):407-413, 2015
  - 6 pairs of normal vs cancer ccRCC tissues
  - SWATH in duplicates

# ~20% of predicted missing proteins are supported by $\geq$ 1 reported peptide in the screen



A Strategy 1 (complex to proteins in the peptide list back to self)

	· · ·		· · · · · · · · · · · · · · · · · · ·	·		
Sample	N T1-> N T1	N T2 -> N T2	C T1-> C T1	C T2 -> C T2		
1	0.203 0	0.220 0	0.186 0.001	0.191 0		
	985 200	937 206	823 153	911 174		
2	0.204 0	0.222 0	0.194 0.004	0.215 0		
	936 191	889 197	904 175	918 197		
3	0.197 0	0.212 0	0.241 0	0.225 0		
	972 191	950 201	849 205	840 189		
4	0.223 0	0.232 0	0.215 0.001	0.211 0		
	943 210	948 220	925 199	930 196		
5	0.225 0	0.201 0	0.209 0	0.185 0		
	912 205	964 194	877 183	904 167		
6	0.249 0	0.215 0	0.233 0	0.241 0		
	883 220	977 210	886 206	927 223		

Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

## ~20% of predicted missing proteins are supported by $\geq$ 1 reported peptide in the replicate



20

**B** Strategy 2 (complex to proteins in the peptide list in the other replicate)

Sample	N T1-> N T2	N T2 -> N T1	C T1-> C T2	C T2 -> C T1
1	0.212 0	0.210 0	0.198 0	0.182 0
	985 209	937 197	823 163	911 166
2	0.213 0	0.216 0	0.205 0	0.202 0.001
	936 199	889 192	904 185	918 185
3	0.212 0	0.196 0	0.218 0	0.249 0
	972 206	950 186	849 185	840 209
4	0.224 0	0.233 0	0.197 0.002	0.222 0
	943 211	948 221	925 182	930 206
5	0.188 0.002	0.235 0	0.185 0	0.209 0
	912 171	964 227	877 162	904 189
6	0.224 0	0.246 0	0.227 0	0.249 0
	883 198	977 240	886 201	927 231

Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

## But ~25% of predicted missing proteins are supported by peptides in the screen or replicate



**C** Strategy 3 (complex to proteins in the peptide list union of self and other replicate)

Sample	N T1-> N T12	N T2 -> N T12	C T1-> C T12	C T2 -> C T12	
1	0.248 0	0.258 0	0.238 0	0.229 0.001	
	985 244	937 242	823 196	911 209	
2	0.248 0	0.260 0	0.225 0	0.234 0.001	
	936 232	889 231	904 203	918 215	
3	0.243 0	0.241 0	0.274 0	0.281 0	
	972 236	950 229	849 233	840 236	
4	0.268 0	0.280 0	0.251 0	0.263 0	
	943 253	948 265	925 232	930 245	
5	0.254 0	0.267 0	0.241 0	0.238 0	
	912 232	964 257	877 211	904 215	
6	0.280 0	0.275 0	0.269 0	0.283 0	
	883 247	977 269	886 238	927 262	

Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

#### ~25% FCS-predicted missing protein are supported by peptides in screen/replicate. Can we do better?

**Recall this postulate:** 

The chance of a protein complex being present is proportional to the fraction of its protein members being correctly reported in the screen

Presence of complex implies presence of all member proteins

**PROTREC:** Rank predicted missing proteins by

Prob(Protein p is present but unreported) = Max<sub>complex C contains p</sub> Prob(C is present)

Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

Copyright 2016 © Limsoon Wong



N1\_T12 N1\_T12 Α 1400 1.0 1200 0.8 1000 0.6 800 Proportion FCS Count 600 0.4 400 0.2 200 0.0 0 0.9 0 01 02 03 04 05 06 07 08 0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1 Prob(Protein p is present but unreported) Prob(Protein p is present but unreported) N1\_T12 В N1\_T12 1:0 1000 0.8 800 PROTREC 0.6 600 Count Proportion 0.4 400 200 0.2 0 0.0 0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1 0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1 Prob(Protein p is present but unreported) Prob(Protein p is present but unreported) **Original Screen** Validated Unvalidated

Ranking by PROTREC significantly improves precision of FCS predictions

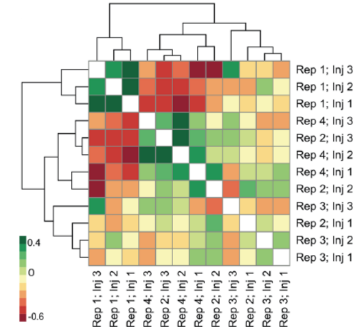
Improving consistency in proteomic profile analysis



## Proteomic profiles generally not NUS consistent, even for technical replicates

#### A human kidney tissue

- Guo et al. Nature Medicine, 21(4):407-413, 2015
- Digested in quadruplicates
- Analyzed in triplicates
- Clustering by proteins
  - Correlation betw replicates is not good (~0.4)
  - Technical replicates of the same biological replicate are not tightly clustered

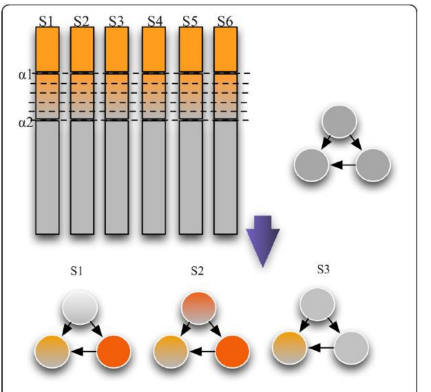


Goh et al. Quantitative proteomics signature profiling based on network contextualization. *Biology Direct*, 10:71. 2015

qPSP



27



**Fig. 1** Schematic demonstrating qPSP's fuzzification procedure. First, alpha1 at top 10 % was defined. An alpha2 was defined from top 10-20 %. To place less confidence in the lower-scoring alpha2, proteins that fall within this range were grouped into 5 bins with descending weights. The modulated hit-rates for each sample could then be used for generating each sample's proteomic signature profile

- Features are complexes
- Feature values are fuzzy weighted proportion of proteins in a complex

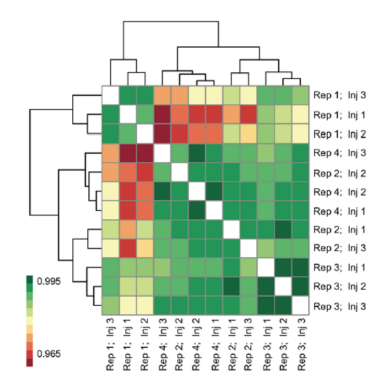
- score(C,S<sub>i</sub>) =  $\Sigma_{p \in C} fs(p,S_i) / |C|$ 

• Complex C is significant if  $\{score(C,S_i) \mid S_i \in A\}$  is very different by t-test from  $\{score(C,S_i) \mid S_i \in B\}$ 

## Consistency of qPSP



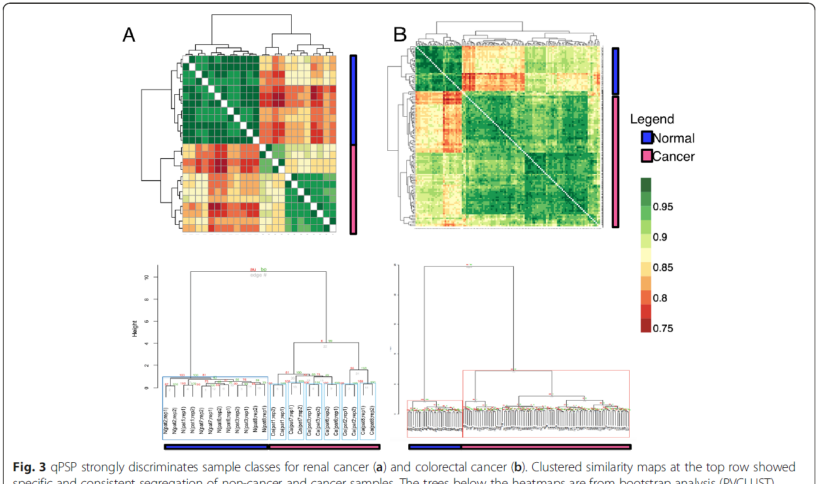
- Clustering of benchmarking control data based on protein complexes (i.e. qPSP)
  - Correlation betw replicates is >0.95
    - Cf. 0.4 based on proteins
  - Technical replicates are better clustered



## Application to renal & colorectal cancers



29



specific and consistent segregation of non-cancer and cancer samples. The trees below the heatmaps are from bootstrap analysis (PVCLUST), which demonstrates that the discrimination between sample classes based on qPSP hit-rates is highly stable

#### Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

# Further improving consistency, as well as catching significant low-abundance complexes



## ESSNet, adapted for proteomics



31

- Let g<sub>i</sub> be a protein in a given protein complex
- Let p<sub>j</sub> be a patient
- Let q<sub>k</sub> be a normal
- Let  $\Delta_{i,j,k} = \text{Expr}(g_i,p_j) \text{Expr}(g_i,q_k)$
- Test whether  $\Delta_{i,j,k}$  is a distribution with mean 0

- Null hypothesis is "Complex C is irrelevant to the difference between patients and normals, and the proteins in C behave similarly in patients and normals"
- No need to restrict to most abundant proteins
- ⇒ Potential to reliably detect low-abundance but differential proteins

Lim et al. A quantum leap in the reproducibility, precision, and sensitivity of gene expression profile analysis even when sample size is extremely small. *JBCB*, 13(4):1550018, 2015

### Five methods to compare with



- Network-based methods
  - Hypergeometric enrichment (HE)
  - Direct group analysis (DG), similar to GSEA
  - qPSP, Goh et al., *Biology Direct*, 10:71, 2015
  - PFSNET, Goh & Wong, JBCB, 14(5):16500293, 2016
- Standard t-test on individual proteins (SP)

#### Simulated data

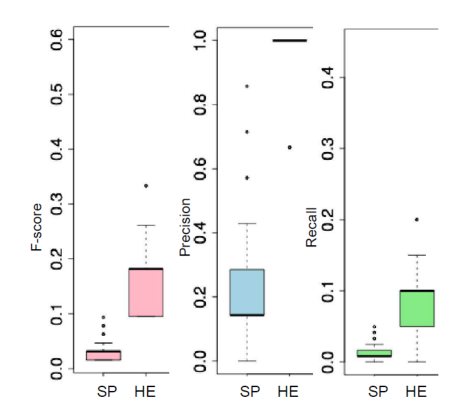


- Simulated datasets from Langley and Mayr
  - D.1.2 is from study of proteomic changes resulting from addition of exogenous matrix metallopeptidase (3 control, 3 test)
  - D2.2 is from a study of hibernating arctic squirrels (4 control, 4 test)
- Both D1.2 and D2.2 have 100 simulated datasets, each with 20% significant features
  - Effect sizes of these differential features are sampled from one out of five possibilities (20%, 50%, 80%, 100% and 200%), increased in one class and not in the other
- Significant artificial complexes are constructed with various level of purity (i.e. proportion of significant proteins in the complex)
  - Equal # of non-significant complexes are constructed as well



## SP shows poor performance on simulated data.

Can networkbased methods do better?



Supplementary Figure 1 Single protein (SP) precision-recall performance on D1.2. The f-score

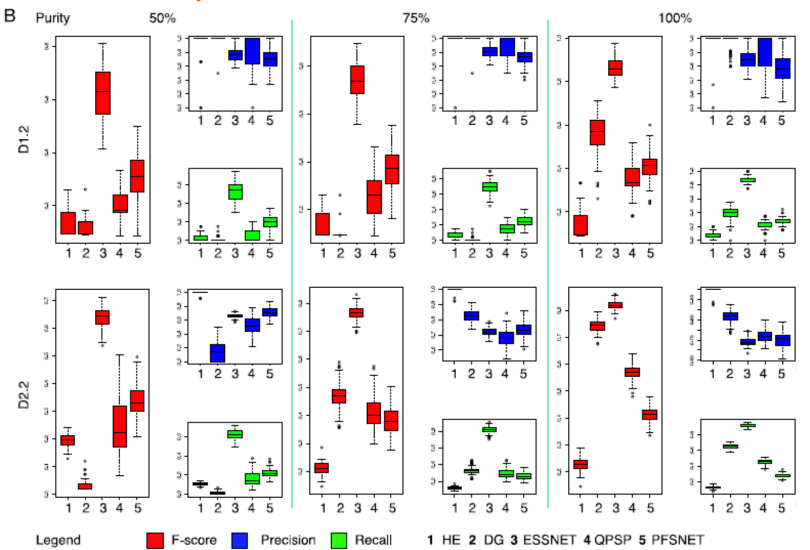
(pink), precision (blue) and recall (green) shows that SP performs abysmally on simulated data. HE is

shown next to SP as a reference.

## ESSNET shows excellent recall/precision on simulated data



35

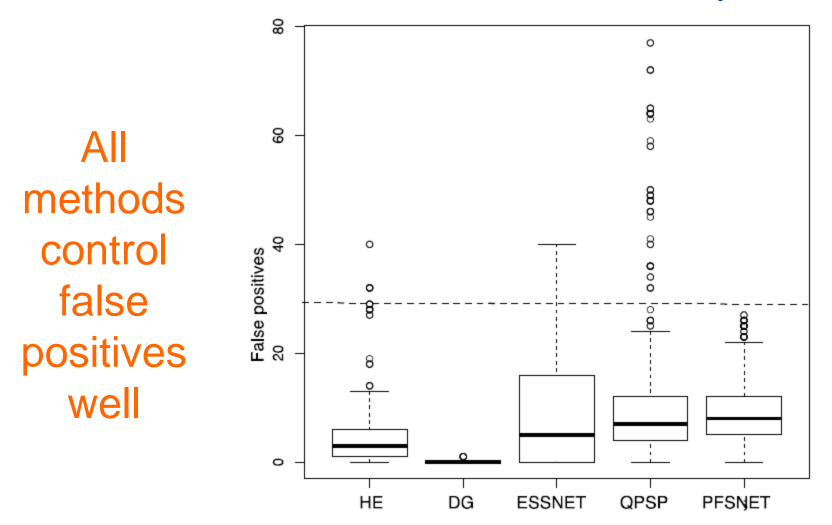


Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016

Renal cancer control data (RCC)

- 12 runs originating from a human kidney tissue digested in quadruplicates and analyzed in triplicates
- Excellent for evaluating false-positive rates of feature-selection methods
  - Randomly split the 12 runs into two groups.
    Report of any significant features between the groups must be false positives





Dash line corresponds to expected # of false positives at alpha 0.05 (~30 complexes)

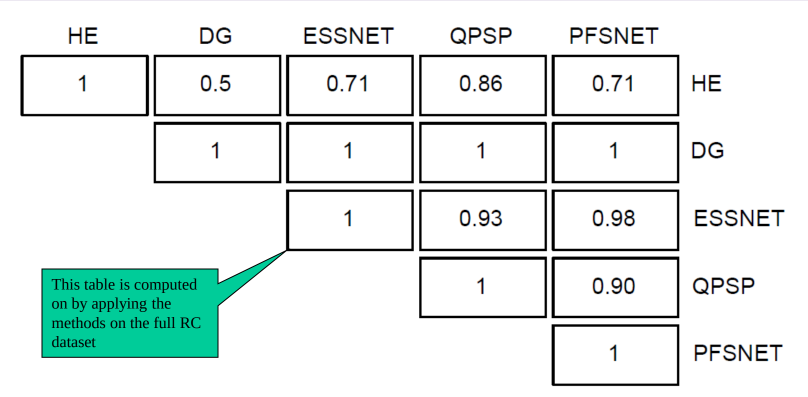


- Renal cancer data (RC)
- 12 samples are run twice so that we have technical replicates over 6 normal and 6 cancer tissues
- Excellent opportunity for testing reproducibility of feature-selection methods
  - A good method should report similar feature sets between replicates
- Can also test feature-selection stability
  - Apply feature-selection method on subsamples and see whether the same features get selected

## ESSNET & PFSNET show excellent reproducibility



Number of terms	HE	DG	ESSNET	QPSP	PFSNET
Replicate 1	4	1	35	86	45
Replicate 2	6	2	29	75	46
Overlaps	0.25	0.5	0.83	0.66	0.94





Number of features Sampling size Δ ΗE DG ESSNET QPSP PFSNET ŏ Feature-selection stability 0.8 \_\_\_\_ 0.4 Ο 

റ്റ

0.0

Sampling size

о

Δ

Δ

**ESSNET &** PFSNET show excellent stability

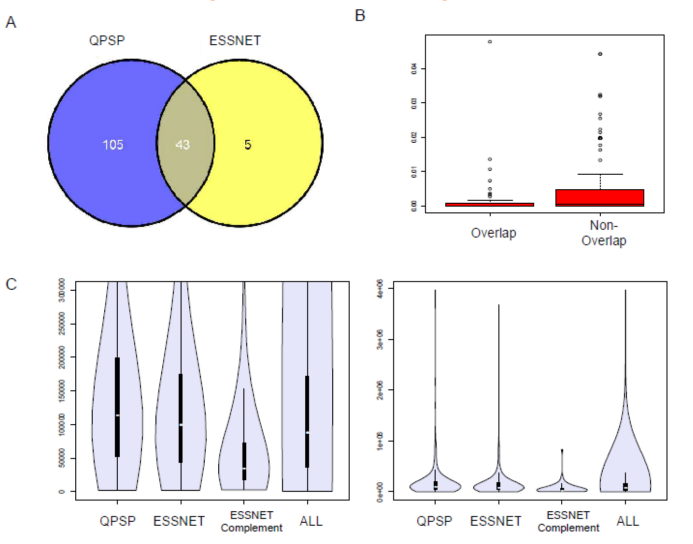
Talk given at IAS, CityU, Hong Kong, 11-15 Aug 2016



## ESSNET & PFSNET show excellent stability

	4	6	8	Mean
HE	0.022	0.016	0.047	0.030
DG	0.001	0.001	0.002	0.001
ESSNET	0.714	0.941	1.000	0.885
QPSP	0.149	0.282	0.991	0.470
PFSNET	1.000	1.000	1.000	1.000

## ESSNET can assay low-abundan



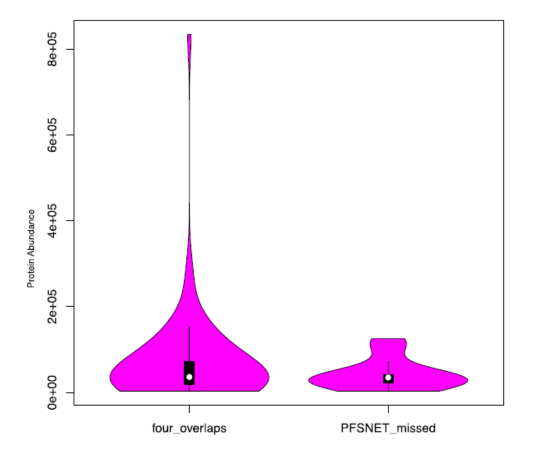
A: QPSP-ESSNET significantcomplex overlaps

42

B: P-value distribution for overlapping and non-overlapping QPSP complexes.

C: Sampling abundance distribution. The left panel is a zoom-in of the right. The y-axis is the protein abundance while the four categories are the distribution of abundances of complexes found in QPSP, ESSNET, **ESSNET** unique (complement), and all proteins in RC.

## ESSNET can assay low-abundan



Of the 5 ESSNETunique complexes, PFSNET can detect 4; the missed complex consists entirely of lowabundance proteins. 43

If p-value threshold is adjusted by Benjamini-Hochberg 5% FDR, PFSNET can detect only 3 of the 5 ESSNETunique complexes while ESSNET continues to detect them all.

#### **Concluding Remarks**





#### In conclusion...

## Contextualization (into complexes) can deal with coverage and consistency issues in proteomics

#### References



- Goh & Wong. Integrating networks and proteomics: Moving forward. Trends in Biotechnology, in press
- [FCS] Goh et al. Comparative network-based recovery analysis and proteomic profiling of neurological changes in valporic acid-treated mice. Journal of Proteome Research, 12(5):2116-2127, 2013
- [qPSP] Goh et al. Quantitative proteomics signature profiling based on network contextualization. *Biology Direct*, 10:71, 2015
- [PFSNET] Goh & Wong. Evaluating feature-selection stability in nextgeneration proteomics. Journal of Bioinformatics and Computational Biology,14(5):16500293, 2016
- [ESSNET] Goh & Wong. Advancing clinical proteomics via analysis based on biological complexes: A tale of five paradigms. Journal of Proteome Research, in press
- [PROTREC] Goh & Wong. Recoving missing proteins based on biological complexes. In preparation