

## Bioinformatics 2

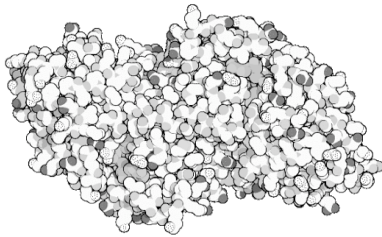
### Protein (Interaction) Networks

Armstrong, 2009

- Biological Networks in general
- Metabolic networks
- Briefly review proteomics methods
- Protein-Protein interactions
- Protein Networks
- Protein-Protein interaction databases
- An example

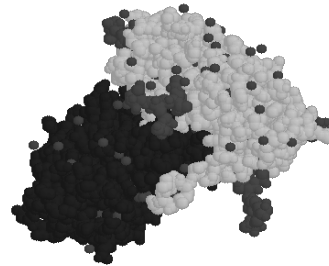
Armstrong, 2009

### alcohol dehydrogenase



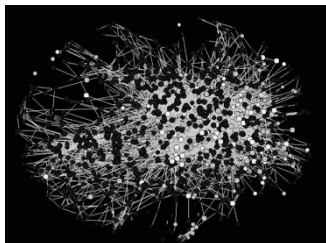
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### ricin (A and B)



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### synaptic proteome



Armstrong, 2009

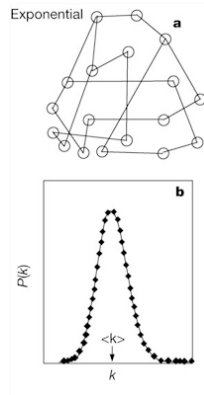
## Biological Networks

- Genes - act in cascades
- Proteins - form functional complexes
- Metabolism - formed from enzymes and substrates
- The CNS - neurons act in functional networks
- Epidemiology - mechanics of disease spread
- Social networks - interactions between individuals in a population
- Food Chains

Armstrong, 2009

## Large scale organisation

- First networks in biology generally modeled using classic random network theory.
- Each pair of nodes is connected with probability  $p$
- Results in model where most nodes have the same number of links  $\langle k \rangle$
- The probability of any number of links per node is  $P(k) \approx e^{-k}$



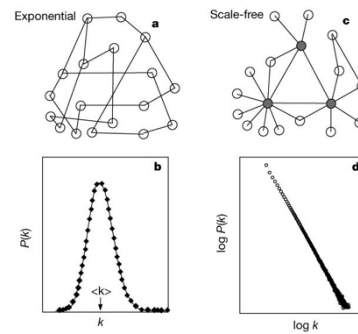
Armstrong, 2009



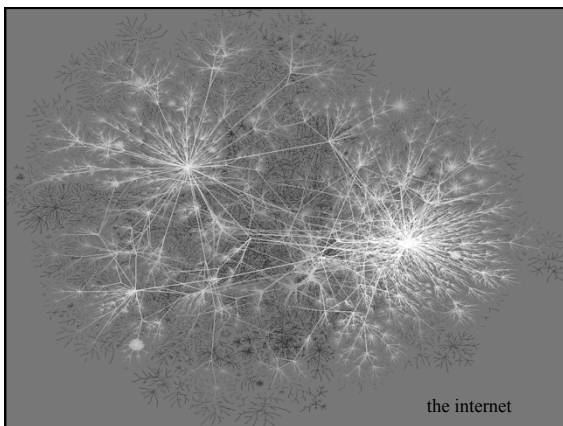
## Non-biological networks

- Research into WWW, internet and human social networks observed different network properties
  - 'Scale-free' networks
  - $P(k)$  follows a power law:  $P(k) \approx k^{-\gamma}$
  - Network is dominated by a small number of highly connected nodes - hubs
  - These connect the other more sparsely connected nodes

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Armstrong, 2009



## Small worlds

- General feature of scale-free networks
  - any two nodes can be connected by a relatively short path
  - average between any two people is around 6
    - What about SARS???
  - 19 clicks takes you from any page to any other on the internet.

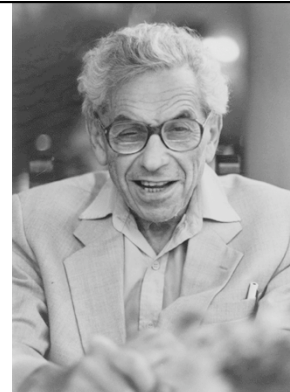
Armstrong, 2009

## 6 degrees of separation..?

- Stanley Milgram's work in late 1960's
- Sent letters to people in Nebraska
- Target unknown person in Massachusetts
- Average 6 'jumps' to reach target

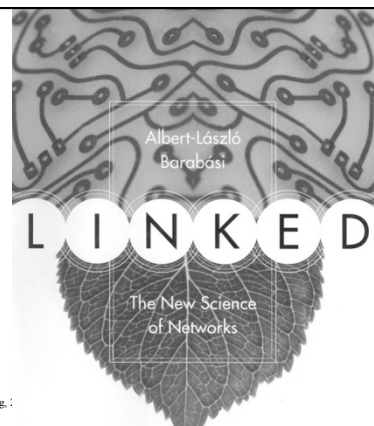
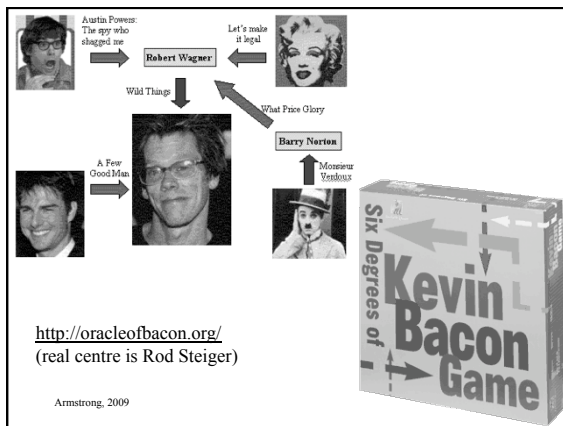
(only 5% got there)

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Armstrong, 2009

Paul Erdős, the most prolific mathematician who ever lived, has no home and no job, but he has reached the world for over fifty years, inspiring other mathematicians. From the *Biography* N to a Number: A Portrait of Paul Erdős by George Gonyea



## Biological organisation

Jeong et al., 2000 *The large-scale organisation of metabolic networks. Nature* 407, 651-654

- Pioneering work by Oltvai and Barabási
- Systematically examined the metabolic pathways in 43 organisms
- Used the WIT database
  - 'what is there' database
  - <http://wit.mcs.anl.gov/WIT2/>
  - Genomics of metabolic pathways



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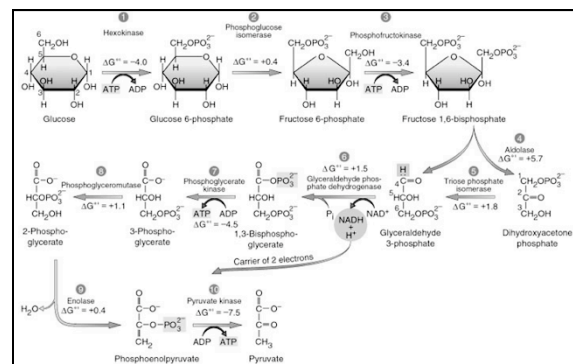
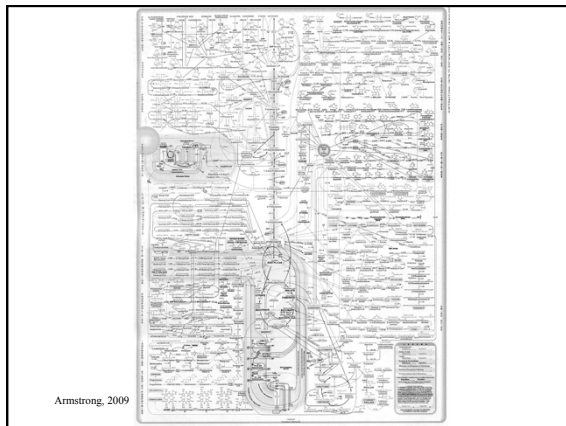
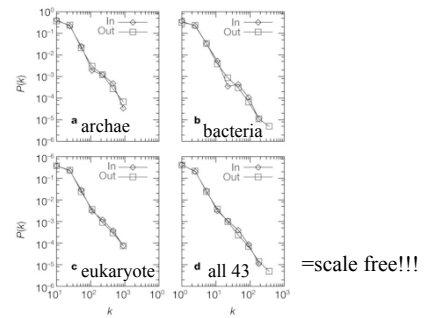


Image taken from <http://fig.cox.miami.edu/~cmallery/255/255atp/255makeatp.htm>

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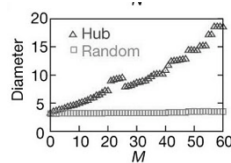


## Using metabolic substrates as nodes



## Random mutations in metabolic networks

- Simulate the effect of random mutations or mutations targeted towards hub nodes.
  - Measure network diameter
  - Sensitive to hub attack
  - Robust to random



## Consequences for scale free networks

- Removal of highly connected hubs leads to rapid increase in network diameter
  - Rapid degeneration into isolated clusters
  - Isolate clusters = loss of functionality
- Random mutations usually hit non hub nodes
  - therefore robust
- Redundant connectivity (many more paths between nodes)

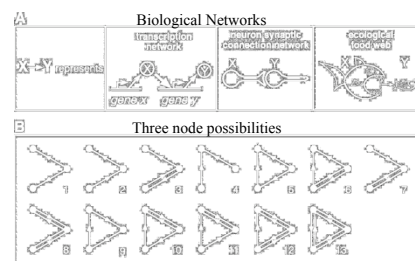
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## Network Motifs

- Do all types of connections exist in networks?
- Milo et al studied the transcriptional regulatory networks in yeast and E.Coli.
- Calculated all the three and four gene combinations possible and looked at their frequency

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Milo et al. 2002 Network Motifs: Simple Building Blocks of Complex Networks. Science 298: 824-827



Armstrong, 2009

## Gene sub networks

| Network                                | Nodes | Edges | $N_{real}$ | $N_{rand} \pm SD$ | Z score | $N_{real}$ | $N_{rand} \pm SD$ | Z score |
|--|-------|-------|------------|-------------------|---------|------------|-------------------|---------|
| <b>Gene regulation (transcription)</b> |       |       |            |                   |         |            |                   |         |
|  |       |       |            |                   |         |            |                   |         |
| <i>E. coli</i>                         | 424   | 519   | 40         | 7 ± 3             | 10      | 203        | 47 ± 12           | 13      |
| <i>S. cerevisiae</i> *                 | 685   | 1,052 | 70         | 11 ± 4            | 14      | 1812       | 300 ± 40          | 41      |

Heavy bias in both yeast and E.coli towards these two sub network architectures

Armstrong, 2009

| Network  | Nodes   | Edges  | $N_{real}$ | $N_{rand} \pm SD$ | Z score | $N_{real}$ | $N_{rand} \pm SD$ | Z score |
|--|---------|--------|------------|-------------------|---------|------------|-------------------|---------|
| <b>Gene regulation (transcription)</b>                     |         |        |            |                   |         |            |                   |         |
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| <b>Feed-back</b>   |         |        |            |                   |         |            |                   |         |
| <i>E. coli</i>   | 424     | 519    | 125        | 80 ± 10           | 3.7     | 137        | 55 ± 11           | 5.3     |
| <i>S. cerevisiae</i> *                                     | 685     | 1,052  | 227        | 110 ± 10          | 2.0     | 335        | 130 ± 10          | 2.0     |
| <b>Electronic circuits (forward logic chips)</b>           |         |        |            |                   |         |            |                   |         |
| Little Rock  | 92      | 984    | 3218       | 3120 ± 50         | 2.1     | 7295       | 2220 ± 210        | 25      |
| Vision   | 83      | 391    | 1182       | 1020 ± 20         | 7.2     | 1357       | 230 ± 50          | 23      |
| St. Martin   | 42      | 205    | 469        | 450 ± 10          | N5      | 382        | 150 ± 20          | 12      |
| Chameleon  | 31      | 87     | 80         | 52 ± 4            | N5      | 26         | 5 ± 2             | 8       |
| Cochella   | 29      | 243    | 279        | 255 ± 12          | 3.6     | 181        | 80 ± 20           | 5       |
| Clayton  | 25      | 49     | 264        | 250 ± 5           | 5.5     | 197        | 80 ± 20           | 13      |
| W. Brook   | 21      | 159    | 381        | 330 ± 7           | 5.3     | 267        | 50 ± 7            | 32      |
| <b>Electronic circuits (digital fractional amplifiers)</b> |         |        |            |                   |         |            |                   |         |
| u1580  | 10,383  | 14,240 | 434        | 2 ± 2             | 285     | 1040       | 1 ± 1             | 3200    |
| u4856  | 20,717  | 34,234 | 413        | 10 ± 3            | 120     | 1739       | 6 ± 2             | 800     |
| u9847  | 23,843  | 33,463 | 612        | 3 ± 2             | 400     | 2044       | 1 ± 1             | 2500    |
| u124   | 5,344   | 8,167  | 211        | 2 ± 1             | 340     | 754        | 1 ± 1             | 1000    |
| u1322  | 8,671   | 13,813 | 493        | 2 ± 1             | 225     | 1616       | 1 ± 1             | 4000    |
| <b>World Wide Web</b>                                      |         |        |            |                   |         |            |                   |         |
| u208   | 122     | 180    | 10         | 1 ± 1             | 9       | 4          | 1 ± 1             | 3.8     |
| u420   | 252     | 399    | 20         | 1 ± 1             | 18      | 10         | 1 ± 1             | 30      |
| u834   | 512     | 829    | 40         | 1 ± 1             | 35      | 23         | 1 ± 1             | 25      |
| <b>Armstrong</b>   |         |        |            |                   |         |            |                   |         |
| u1448  | 328,729 | 1,448  | 3,164      | 30 ± 2            | 100     | 6,569      | 340 ± 10          | 15,000  |
| u1449  | 328,729 | 1,448  | 3,164      | 30 ± 2            | 100     | 6,569      | 340 ± 10          | 15,000  |

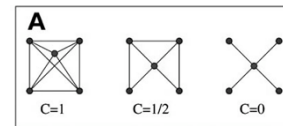
## What about known complexes?

- OK, scale free networks are neat but how do all the different functional complexes fit into a scale free proteome arrangement?
  - e.g. ion channels, ribosome complexes etc?
- Is there substructure within scale free networks?
  - Examine the clustering co-efficient for each node.

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## Clustering co-efficients and networks.

- $C_i = 2n / (k_i(k_i - 1))$
- $n$  is the number of direct links connecting the  $k_i$  nearest neighbours of node  $i$
- A node at the centre of a fully connected cluster has a  $C$  of 1

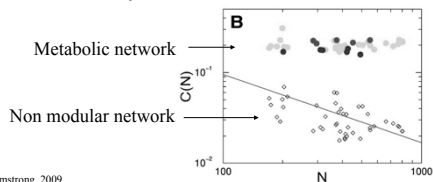


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## Clustering co-efficients and networks.

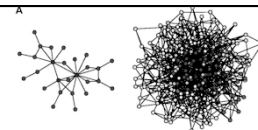
Ravasz et al., (2002) Hierarchical Organisation of Modularity in Metabolic Networks. Science 297, 1551-1555

- The modularity (ave C) of the metabolic networks is an order of magnitude higher than for truly scale free networks.

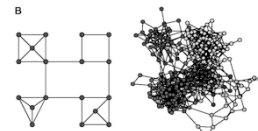


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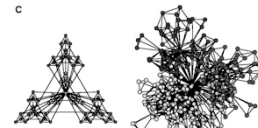
No modularity  
Scale-free



Highly modular  
Not scale free



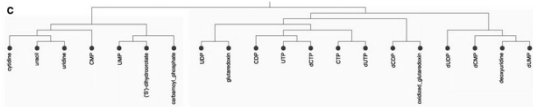
Hierarchical network  
Scale-free



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## Clustering on C

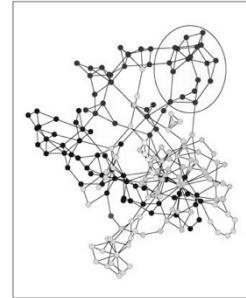
- Clustering on the basis of C allows us to rebuild the sub-domains of the network



- Producing a tree can predict functional clustered arrangements.

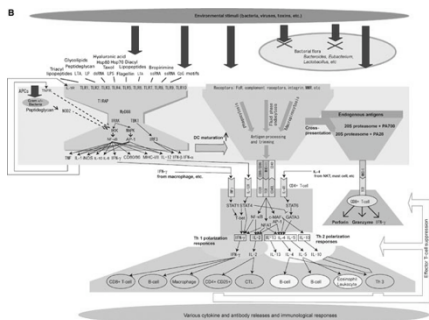
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## Cluster analysis on the network



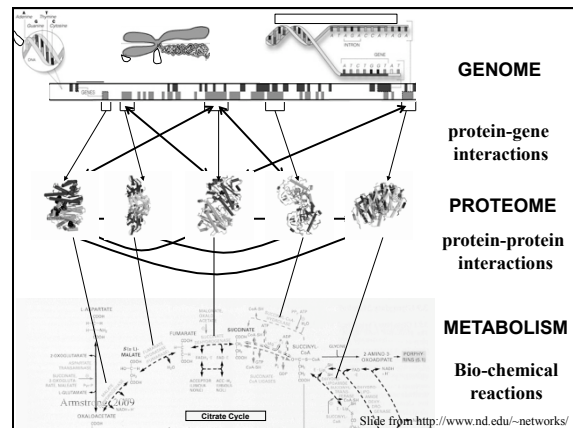
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## Bow-tie and nested bow-tie architectures



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[http://www.nature.com/msb/journal/v2/n1/fig\\_tab/msb4100039\\_F2.html](http://www.nature.com/msb/journal/v2/n1/fig_tab/msb4100039_F2.html)



Slide from <http://www.nd.edu/~networks/>

## Biological Profiling

- Microarrays
  - cDNA arrays
  - oligonucleotide arrays
  - whole genome arrays
- Proteomics
  - yeast two hybrid
  - PAGE techniques
  - Mass Spectrometry (Lecture 2)

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## Protein Interactions

- Individual Proteins form functional complexes
- These complexes are semi-redundant
- The individual proteins are sparsely connected
- The networks can be represented and analysed as an undirected graph

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## How to build a protein network

- Biological sample – how to you isolate your complex?
- What is in your complex?
- How is it connected?
  - Databases and Literature Mining
  - Yeast two hybrid screening & other cellular interaction assays
  - Mass-spec analysis
- Building and analysing the network
- An example

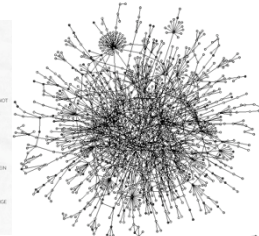
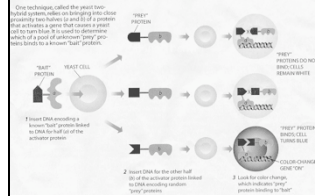
Armstrong, 2009

## Yeast protein network

**Nodes:** proteins

**Links:** physical interactions (binding)

### Finding Proteins That Interact



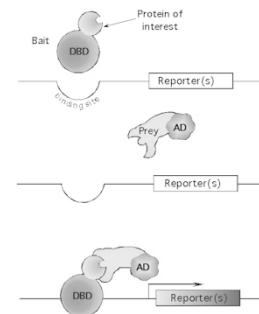
P. Uetz et al. *Nature* 403, 623-7 (2000).

Slide from <http://www.nd.edu/~networks/>

## Yeast two hybrid

- Use two mating strains of yeast
- In one strain fuse one set of genes to a transcription factor DNA binding site
- In the other strain fuse the other set of genes to a transcriptional activating domain
- Where the two proteins bind, you get a functional transcription factor.

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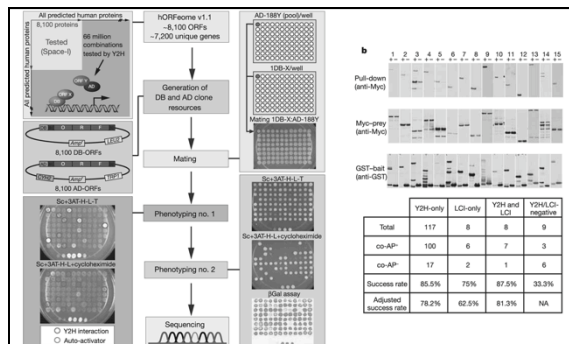


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## Data obtained

- Depending on sample, you get a profile of potential protein-protein interactions that can be used to predict functional protein complexes.
- False positives are frequent.
- Can be confirmed by affinity purification etc.

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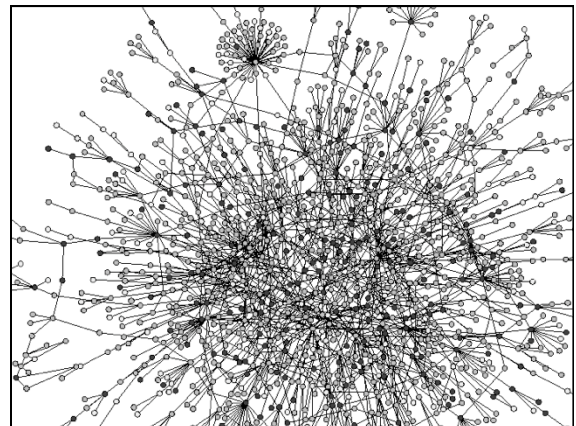
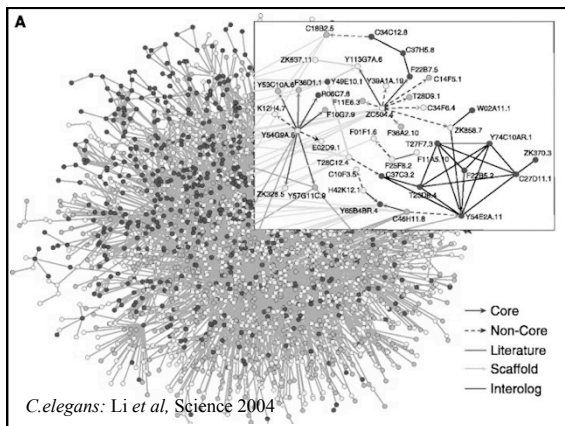
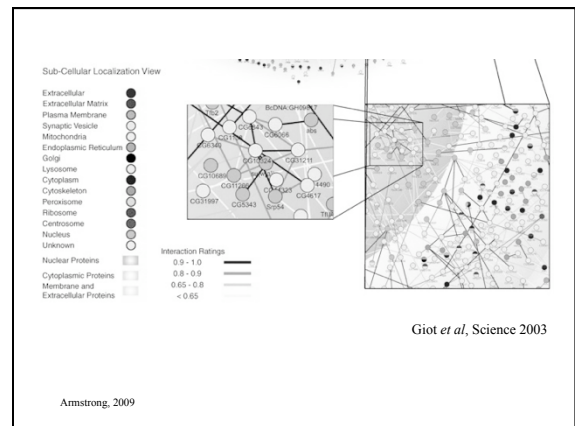
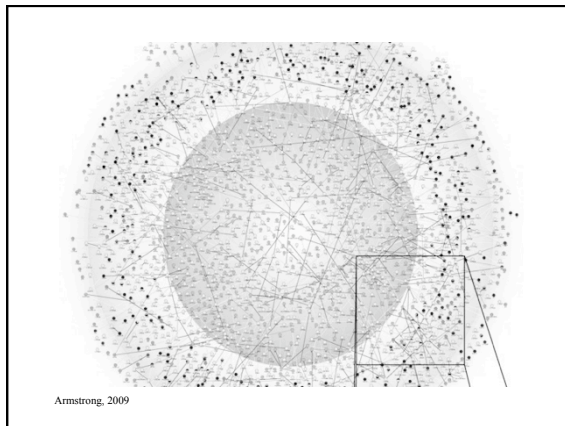
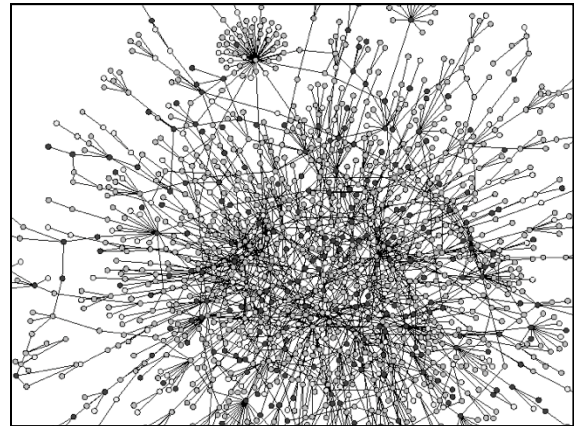
Interaction mapping schema from Rual et al 2005

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## Protein Networks

- Networks derived from high throughput yeast 2 hybrid techniques
  - yeast
  - *Drosophila melanogaster*
  - *C.elegans*
- Predictive value of reconstructed networks

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## Predictive value of networks

Jeong et al., (2001) *Lethality and Centrality in protein networks. Nature 411 p41*

- In the yeast genome, the essential vs. unessential genes are known.
- Rank the most connected genes
- Compare known lethal genes with rank order

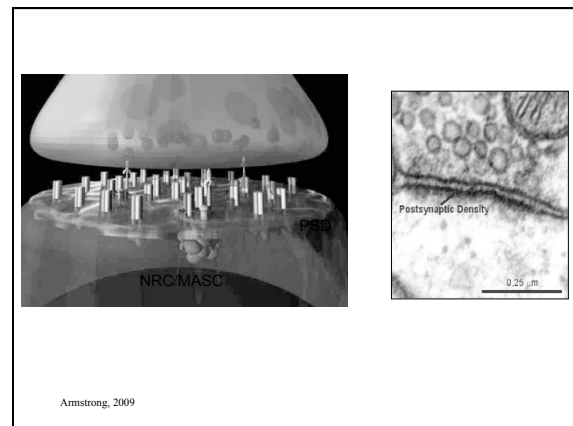
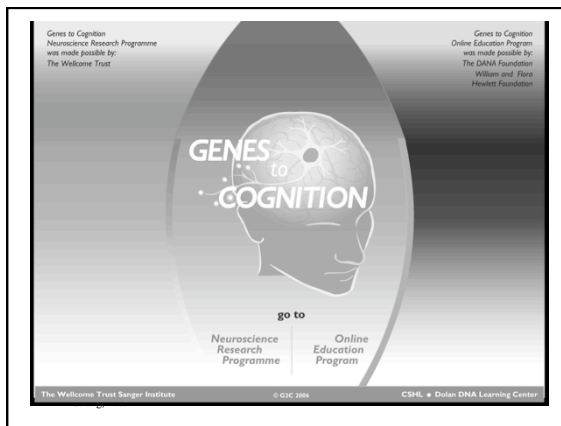
| $k$ | fraction | %lethal |
|-----|----------|---------|
| <6  | 93%      | 21%     |
| >15 | 0.7%     | 62%     |

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## A walk-through example...

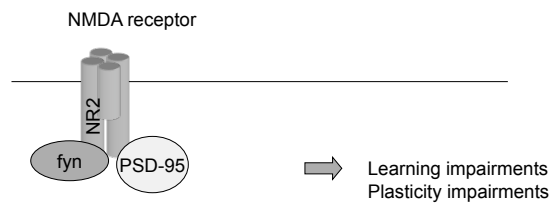
See linked papers on for further methodological details

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## Genetic evidence for postsynaptic complexes



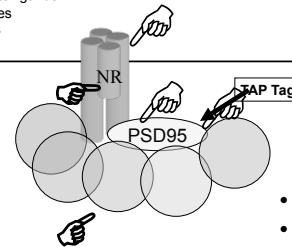
Grant, et al. Science, 258, 1903-10. 1992  
 Migaud et al. Nature, 396, 433-439. 1998  
 Sprengel et al. Cell 92, 279-89. 1998

Armstrong, 2009

## Proteomic characterisation of NRC / MASC

(MAGUK Associated Signaling Complex)

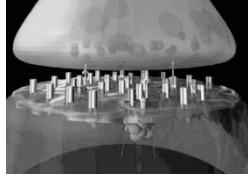
- glutamate ligands
- antibodies
- peptides
- TAP Tag



- ~2 MDa
- 77 proteins (2000)
- 186 (2005)

Husi et al. Nature Neuroscience, 3, 661-669. 2000.  
 Husi & Grant. J. Neurochem, 77, 281-291. 2001  
 Collins et al. J. Neurochem. 2005

Armstrong, 2009



|                       |      |
|-----------------------|------|
| Post Synaptic Density | 1124 |
| ER:microsomes         | 491  |
| Spliceosome           | 311  |
| NRC/MASC              | 186  |
| Nucleolus             | 147  |
| Peroxisomes           | 181  |
| Mitochondria          | 179  |
| Phagosomes            | 140  |
| Golgi                 | 81   |
| Choroplasts           | 81   |
| Lysosomes             | 27   |
| Exosomes              | 21   |

Armstrong, 2009  
Grant. (2006) Biochemical Society Transactions. 34, 59-63. 2006

## Literature Mining

- 680 proteins identified from protein preps
- Many already known to interact with each other
- Also interact with other known proteins
  - Immunoprecipitation is not sensitive (only finds abundant proteins)
- Literature searching has identified a group of around 4200 proteins
  - Currently we have extensive interaction data on 1700

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## Annotating the DB

- How do we find existing interactions?
  - Search PubMed with keyword and synonym combinations
  - Download abstracts
  - Sub-select and rank-order using regex's
  - Fast web interface displays the most 'productive' abstracts for each potential interaction

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## Keyword and synonym problem

- PSD-95:
  - DLG4, PSD-95, PSD95, Sap90, Tip-15, Tip15, Post Synaptic Density Protein - 95kD, PSD 95, Discs, large homolog 4, Presynaptic density protein 95
- NR2a:
  - Glutamate [NMDA] receptor subunit epsilon 1 precursor (N-methyl D-aspartate receptor subtype 2A) (NR2A) (NMDAR2A) (hNR2A) NR2a
- Protein interactions:
  - interacts with, binds to, does not bind to....

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`.\+sand\s.+sinteract`

(1..N characters) (space) and (1..N characters) interact

`.\+s((is)|(was))\sbound\sto\s.+s`

(1..N characters) (space) (is or was) (space) bound (space) to (1..N characters) (space)

`.\+sbinding\s of\s.+s((and)|(to))\s.+s`

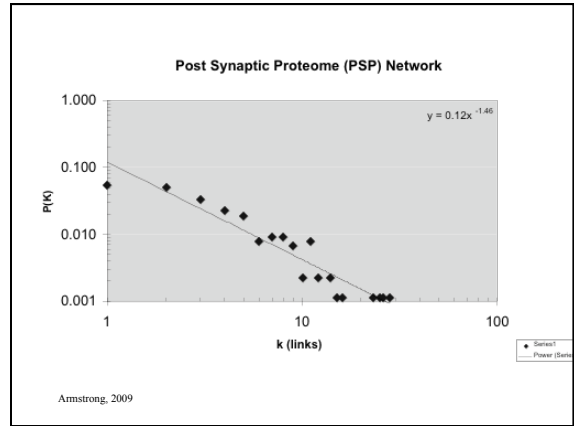
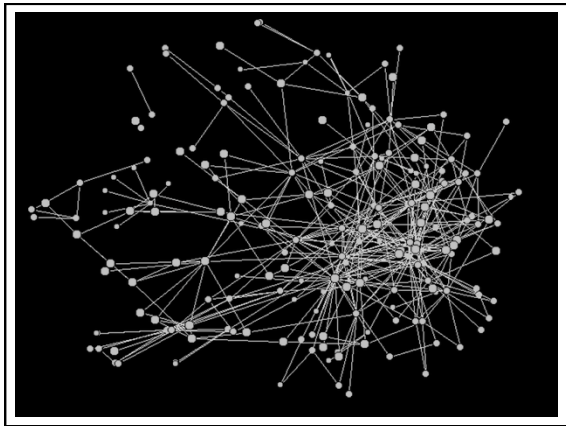
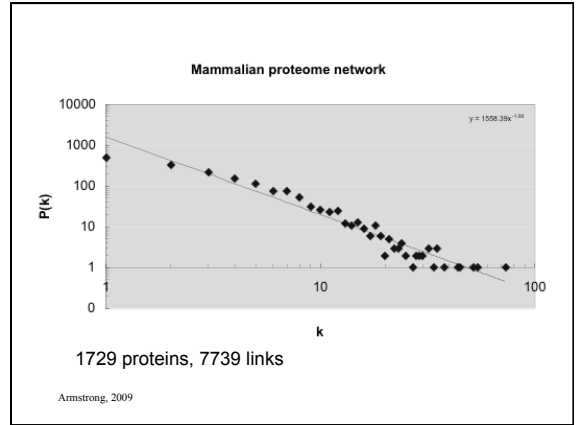
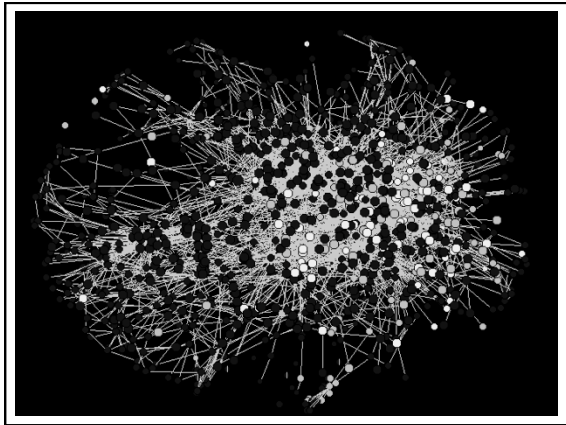
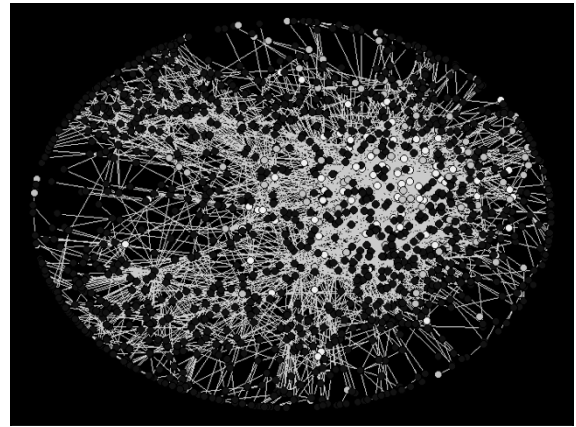
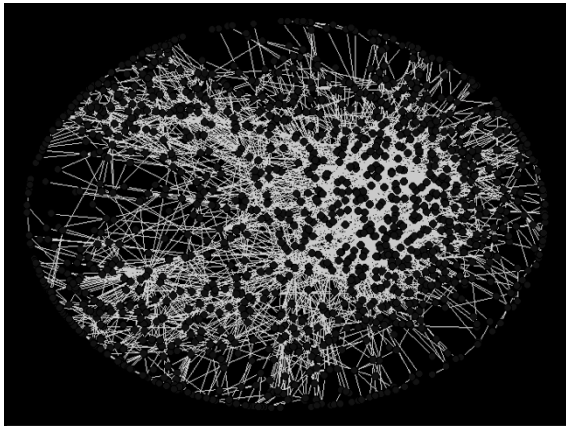
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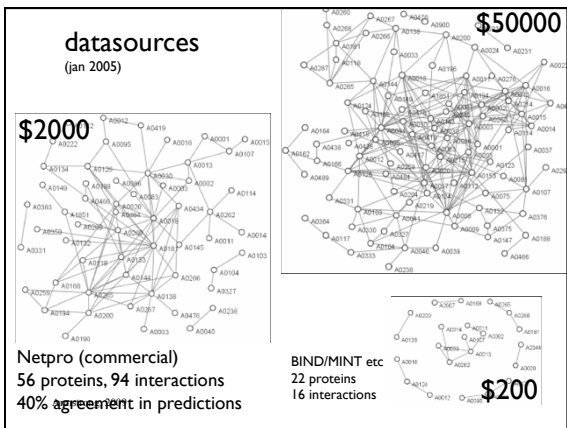
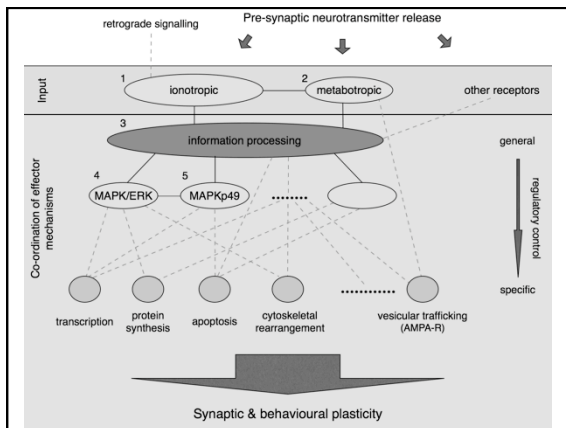
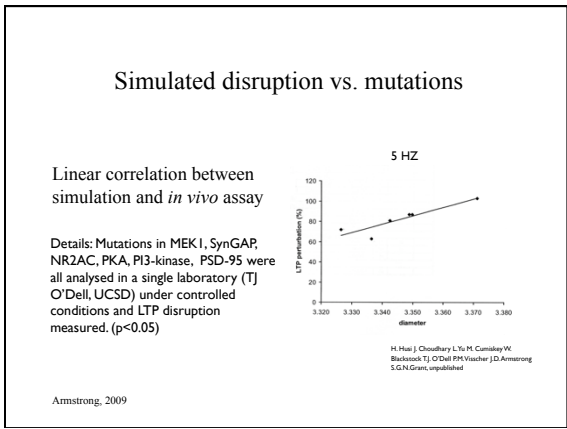
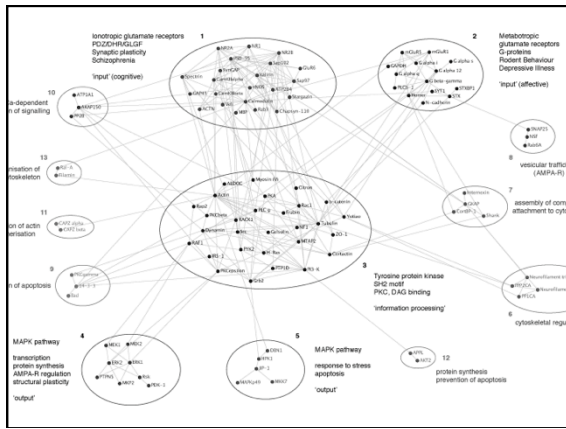
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## Annotating the DB

- How do we find existing interactions?
  - Search PubMed with keyword and synonym combinations
  - Download abstracts
  - Sub-select and rank-order using regex's
  - Fast web interface displays the most 'productive' abstracts for each potential interaction
  - Learn from good vs. bad abstracts

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## Synapse proteome summary

- Protein parts list from proteomics
- Literature searching produced a network
- Network is essentially scale free
- Hubs more important in cognitive processes
- Network clusters show functional subdivision
- Overall architecture resembles bow-tie model
- Expensive...

Armstrong, 2009

### Protein (and gene) interaction databases

- BioGRID - A Database of Genetic and Physical Interactions
- DIP - Database of Interacting Proteins
- MINT - A Molecular Interactions Database
- IntAct - EMBL-EBI Protein Interaction
- MIPS - Comprehensive Yeast Protein-Protein interactions
- Yeast Protein Interactions - Yeast two-hybrid results from Fields' group
- PathCasting - A yeast protein interaction database by Curagen
- SPIID - Bacillus subtilis Protein Interaction Database
- AllFuse - Functional Associations of Proteins in Complete Genomes
- BRITe - Biomolecular Relations in Information Transmission and Expression
- ProMesh - A Protein-Protein Interaction Database
- The PIM Database - by Hybrigenics
- Mouse Protein-Protein interactions
- Human herpesvirus 1 Protein-Protein interactions
- Human Protein Reference Database
- BOND - The Biomolecular Object Network Databank. Former BIND
- MDSIP - Systematic identification of protein complexes in *Saccharomyces cerevisiae* by mass spectrometry
- Proteom - Database of protein-protein complexes enriched with the domain-domain structures
- Proteins that interact with GroEL and factors that affect their release
- DPIDB - DNA-Protein Interaction Database
- YPD<sup>TM</sup> - Yeast Proteome Database by Incyte

Source with links: <http://proteome.wayne.edu/PIIDB.html>

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**BioGRID** BETA  
General Repository for Interaction Datasets

home support contribute downloads mirrors about us

**Search the BioGRID**  
Example: Gankarrin 17A, Entrez-Gene 17A, SOD 17A, Gene Name [none]

Organism: All Organisms

Submit Your Search

**Having Problems Searching?**

**Interaction Statistics**

|                        |        |
|------------------------|--------|
| Total Raw              | 202054 |
| Total Raw Physical     | 140000 |
| Total Raw Genetic      | 62059  |
| Total Non-Redundant    | 128517 |
| Non-Redundant Physical | 92166  |
| Non-Redundant Genetic  | 40657  |

**Database Statistics**

|              |        |
|--------------|--------|
| Proteins     | 322372 |
| Publications | 28120  |
| Organisms    | 13     |

**Download Osprey**  
Osprey is a software platform for visualization of complex interaction networks. Osprey builds data-rich graphical representations from Gene Ontology (GO) annotated interaction data maintained by the BioGRID.  
<http://biodata.mshri.on.ca/osprey>

**Latest News**

**BioGRID version 2.0.36 release (1,831 physical and genetic interactions added)**  
Jul 16, 2008 @ 03:04:47  
The BioGRID's curated set of physical and genetic interactions has been updated to include an additional 1,831 interactions. These additions bring our total number of non-redundant interactions to 132,837 and raw interactions to 202,054. New interactions will be added in curation updates on a monthly basis. Please let us know if we have missed or incorrectly reported any interactions by sending an e-mail to [gridadmin@bioGRID.ca](mailto:gridadmin@bioGRID.ca).

**BioGRID version 2.0.35 release (1,856 physical and genetic interactions added)**  
Dec 16, 2007 @ 03:33:50  
The BioGRID's curated set of physical and genetic interactions has been updated to include an additional 1,856 interactions. These additions bring our total number of non-redundant interactions to 131,983 and raw interactions to 201,323. New interactions will be added in curation updates on a monthly basis. Please let us know if we have missed or incorrectly reported any interactions by sending an e-mail to [gridadmin@bioGRID.ca](mailto:gridadmin@bioGRID.ca).

**BioGRID version 2.0.34 release (576 physical and genetic interactions added)**  
Nov 15, 2007 @ 02:47:55

IntAct : [www.ebi.ac.uk/intact](http://www.ebi.ac.uk/intact)

IntAct Home  
About IntAct  
Search IntAct  
FAQ  
Contact Us  
Privacy Policy  
Terms of Use  
Sitemap

Search IntAct  
To perform a search in the IntAct database use the search box in the top left corner.

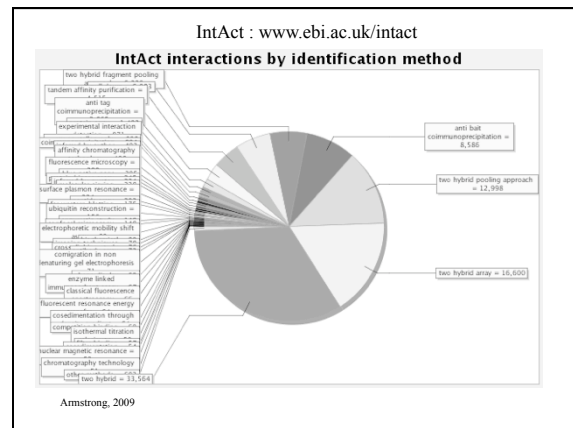
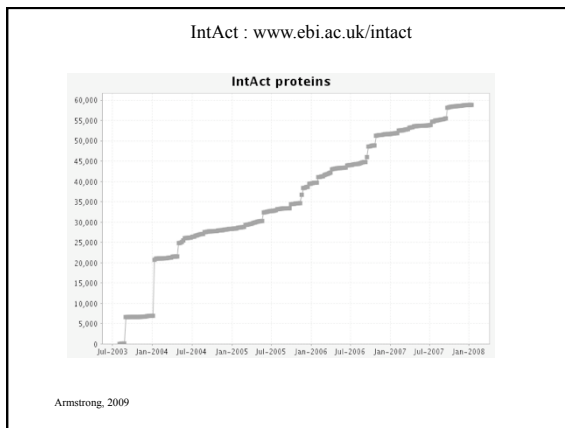
Search criteria:  
Gene name (ECED)  
UniProt ID (2008)  
Pubmed ID (2008)

Introductions  
IntAct provides a freely available, open access database system and analysis tools for protein interaction data. All interactions are derived from literature curation of third user submissions and are freely available.

Default of the month: January  
Using et al. [2008] [PubMed] [IntAct]

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All items available on this site, under the Creative Commons Attribution License. This means that you are free to copy, distribute, display and make derivative use of all content from the IntAct database provided appropriate credit is given.

Armstrong, 2009



**comparing two approaches**

- Pocklington et al 2006
  - Emphasis on QC and literature mining
  - Focussed on subset of molecules
- Rual et al 2005
  - Emphasis on un-biased measurements
  - Focussed on proteome wide models
- Both then look at disease/network correlations

Armstrong, 2009

