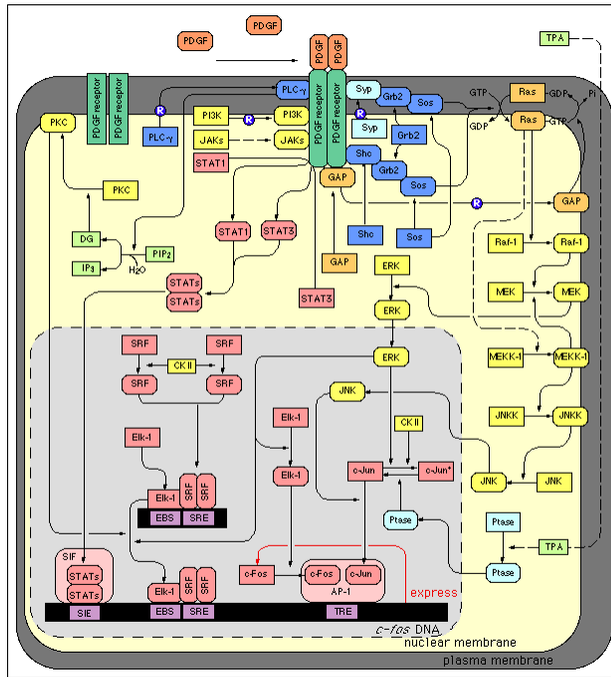




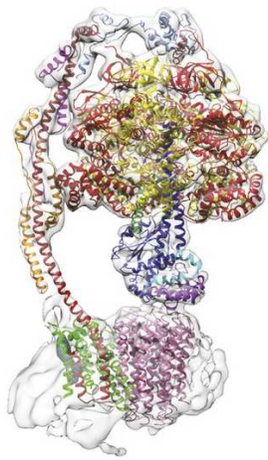
A typical genetic network



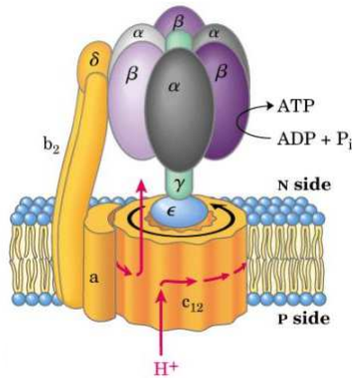
3

Contacts between proteins define protein interaction networks

CryoEM structure of ATP synthase

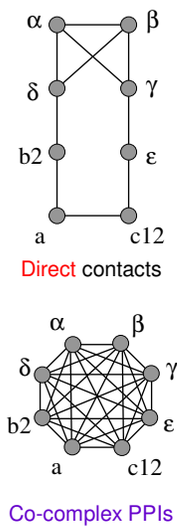


Schematic version



Total set = protein complex  
Sum of direct + indirect interactions

Network representation



4

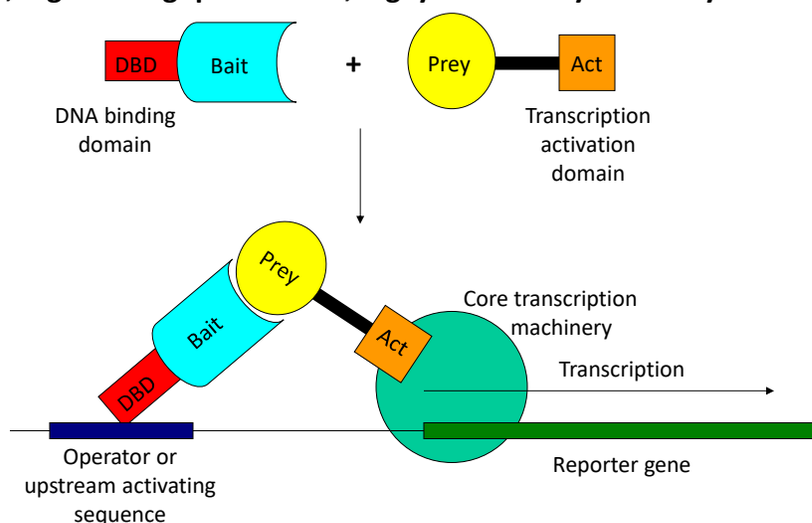
Let's look at some of the types of interaction data in more detail.

Some of these capture physical interactions, some genetic, some informational or logical.

5

### Pairwise protein interactions

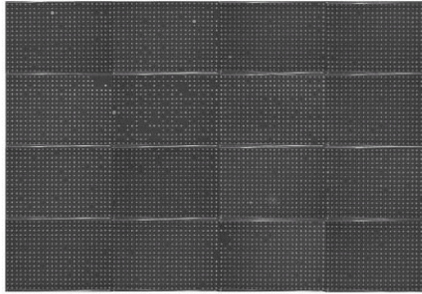
In general, purifying proteins one at a time, mixing them, and assaying for interactions is far too slow & laborious. We need something faster! Hence, high-throughput screens, e.g. yeast two-hybrid assays



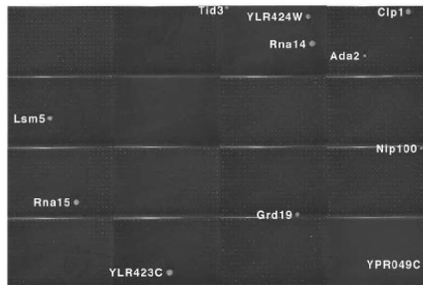
6

## High-throughput yeast two-hybrid assays

Haploid yeast cells expressing activation domain-prey fusion proteins



Diploid yeast probed with DNA-binding domain-Pcf11 bait fusion protein

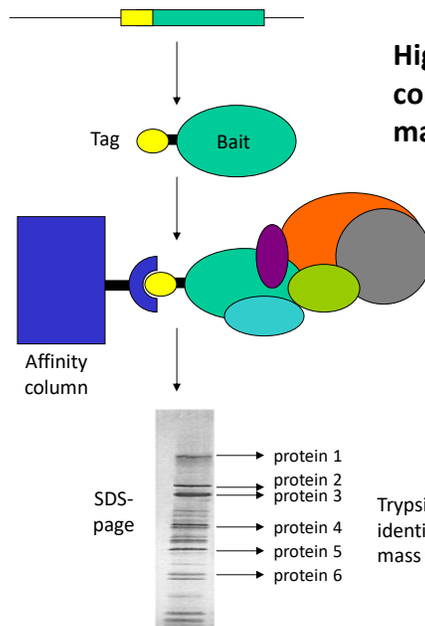


Uetz, Giot, *et al. Nature* (2000)

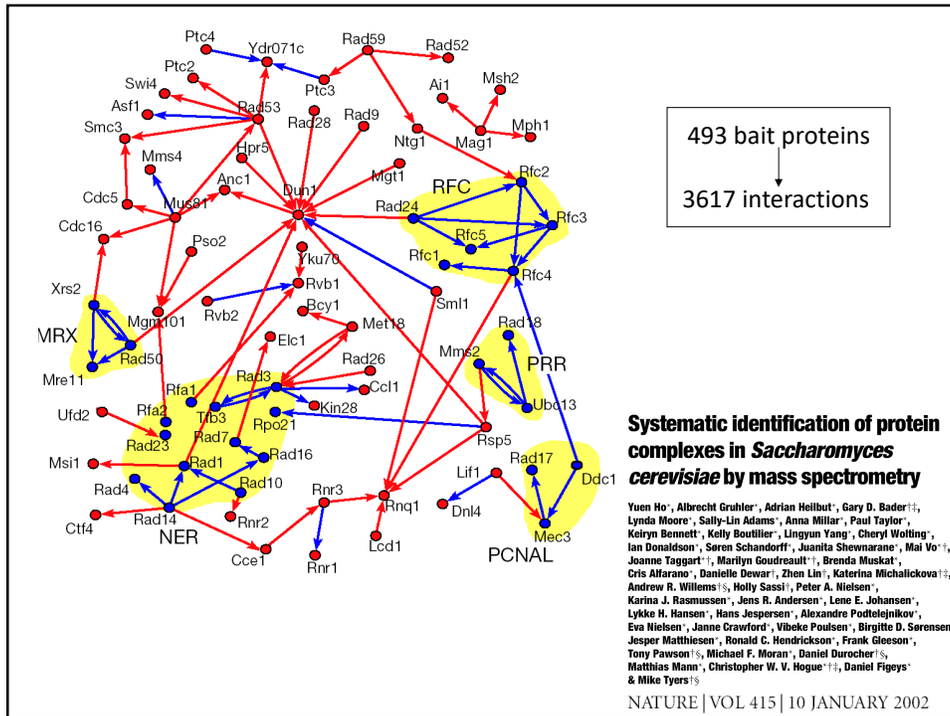
7

## Protein complexes

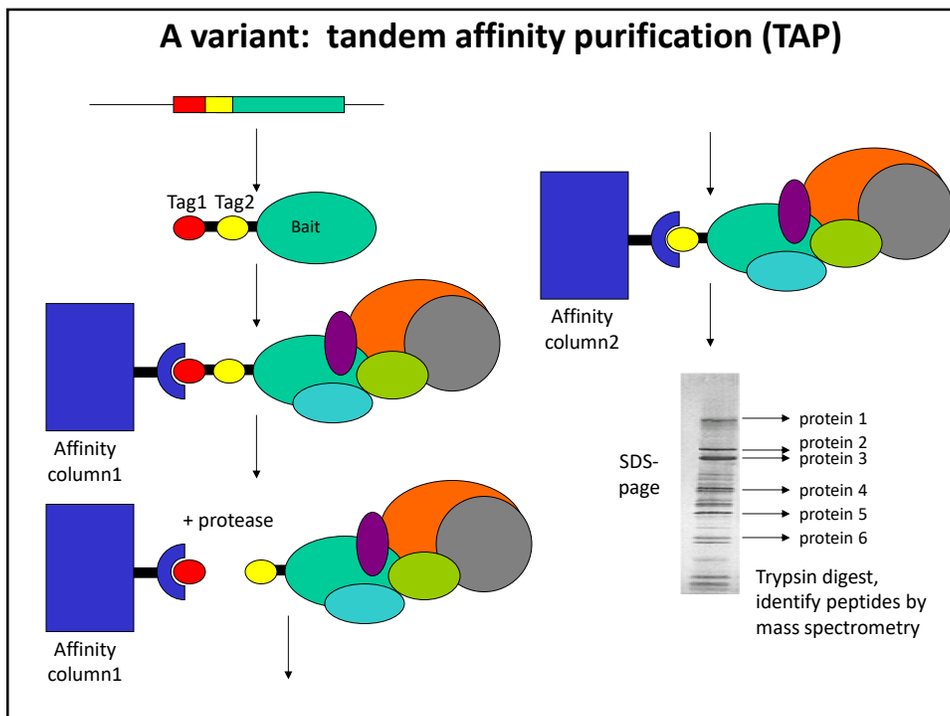
High-throughput complex mapping by mass spectrometry



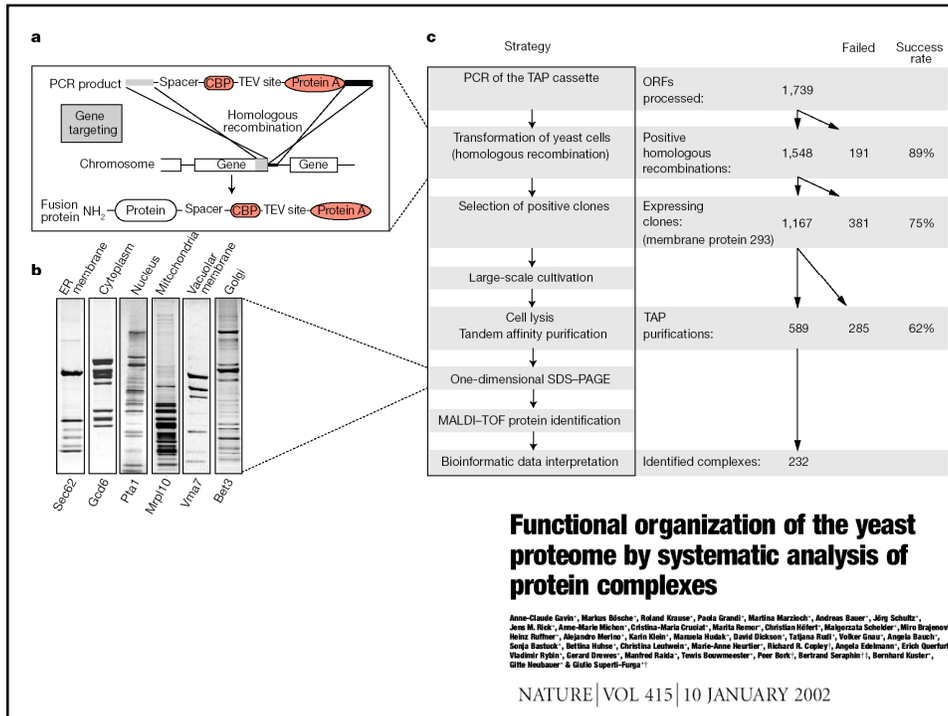
8



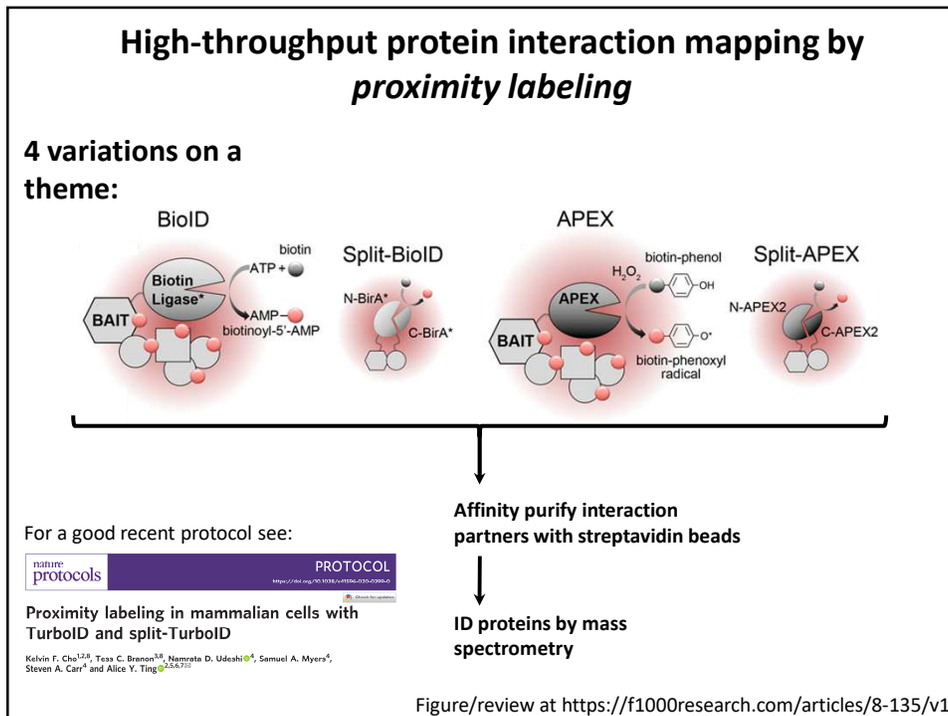
9



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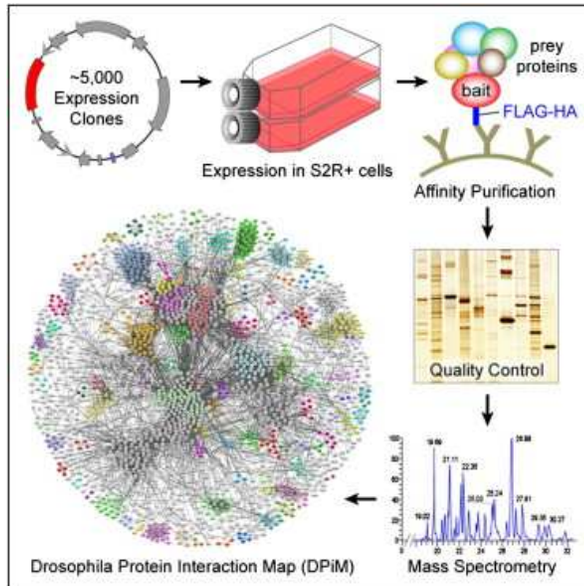


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### The current state-of-the-art in animal PPI maps – AP/MS



~3,500 affinity purification experiments

~11K interactions /  
~2.3K proteins

→ spans 556 complexes

Still daunting for the human proteome, but...

Guruharsha *et al.* (2011) *Cell* 147, 690–703

13

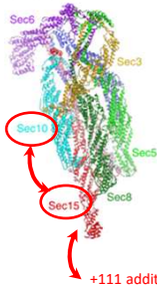
### The current state-of-the-art in human PPI maps – Y2H

Human ORFeome (v9.1) → now ~90% of the protein-coding genes!

Screened *all x all* (150M pairs!) in 9 Y2H assays

52,569 PPIs involving 8,275 proteins

<https://www.nature.com/articles/s41594-017-0016-2>

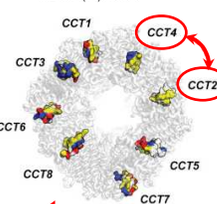


+111 additional PPIs

Y2H captures pairwise PPIs that can form when the proteins are expressed out of biological context (e.g., as fusion proteins in a yeast cell nucleus). It can reveal directly contacting proteins but often misses those that require additional molecular context or higher order assemblies,

← the exocyst e.g. the CCT complex →

[https://www.cell.com/cell/fulltext/S0092-8674\(14\)01369-5](https://www.cell.com/cell/fulltext/S0092-8674(14)01369-5)

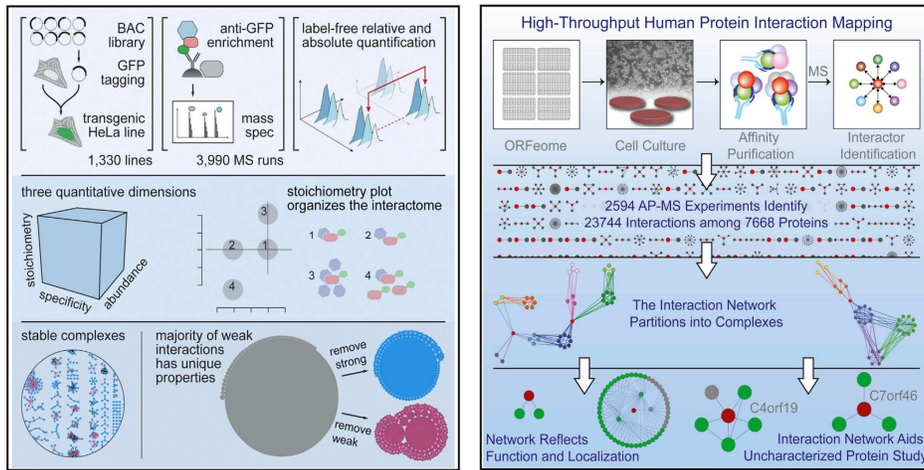


+15 additional PPIs

Luck *et al.*, A reference map of the human protein interactome, *bioRxiv*, posted April 10, 2019  
<https://www.biorxiv.org/content/10.1101/605451v1>, published *Nature*, April 8, 2020

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## The current state-of-the-art in human PPI maps – large scale AP/MS



Hein *et al.*, *Cell* (2015) 163:712-23.

Huttlin *et al.*, *Cell* (2015) 162:425-440

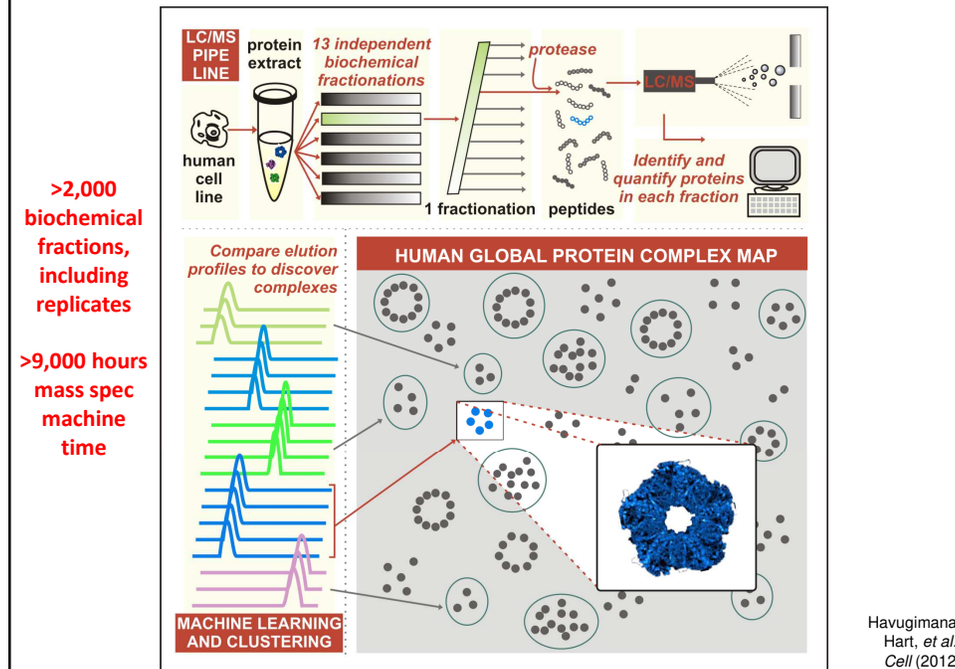
Huttlin *et al.*, *Nature* (2017) 545:505-509

Huttlin *et al.*, *bioRxiv* (2020)

**Just in the past 5 years, >16K affinity purification/mass spec experiments on tagged human proteins expressed in cell lines**

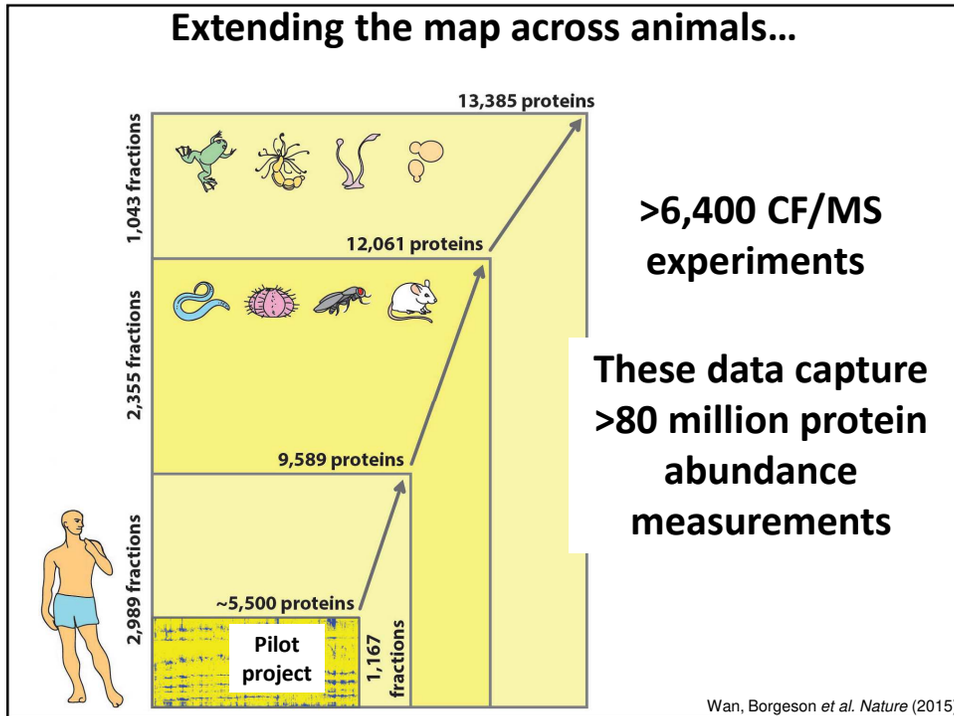
15

## The current state-of-the-art in animal PPI maps – co-fractionation/MS

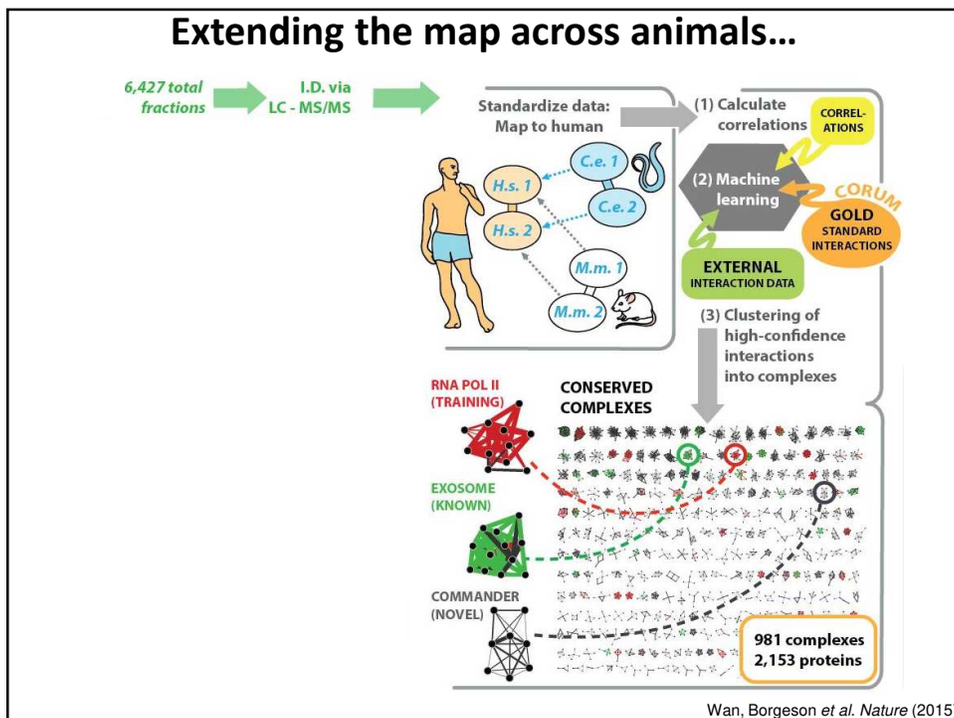


Havugimana,  
Hart, *et al.*,  
*Cell* (2012)

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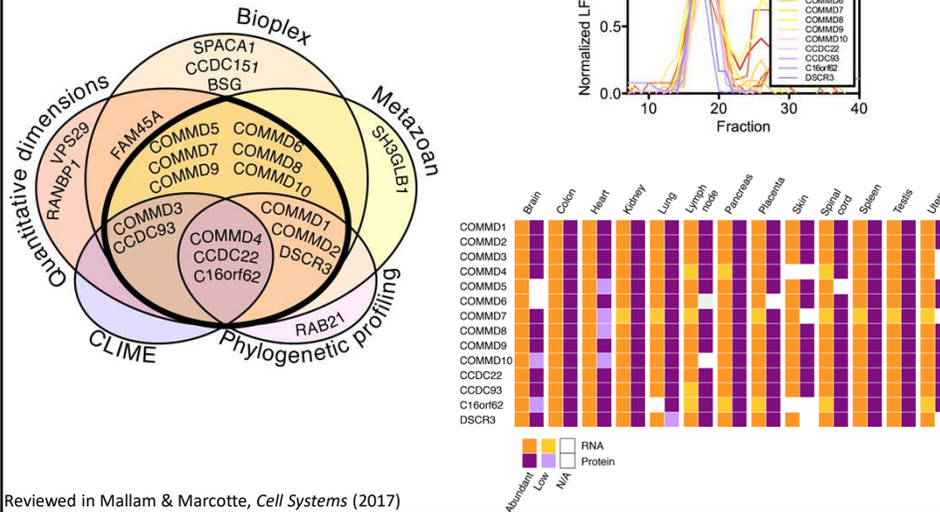
17



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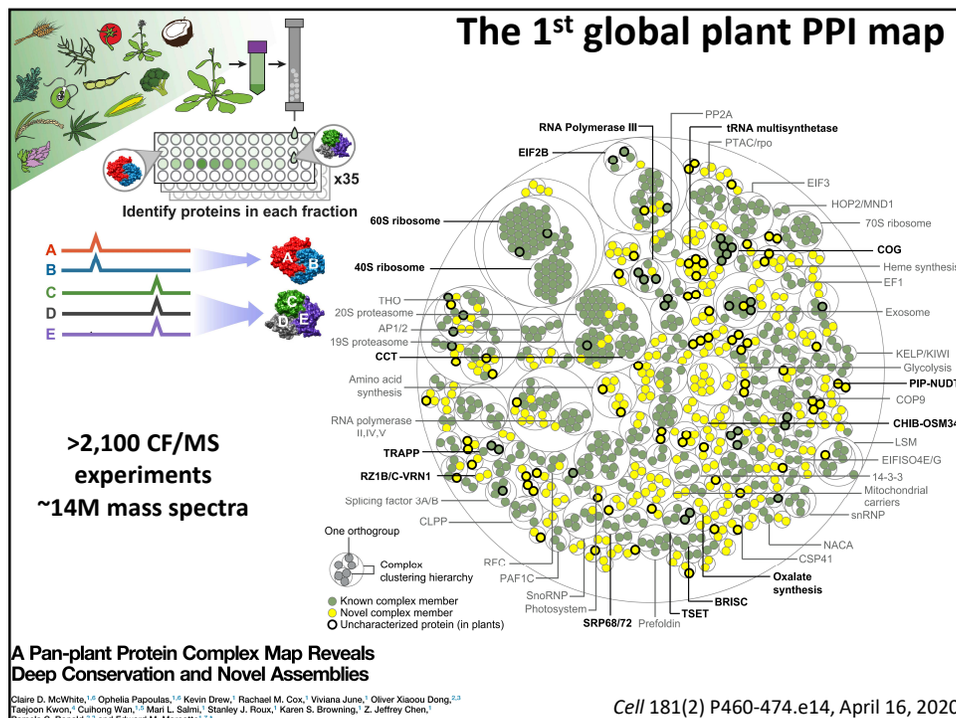
## There are still lots of cellular machines left to find

e.g. the “Commander” complex, found in all 3 large human PPI maps, a 600 kDa protein complex expressed in nearly every human cell type and tissue



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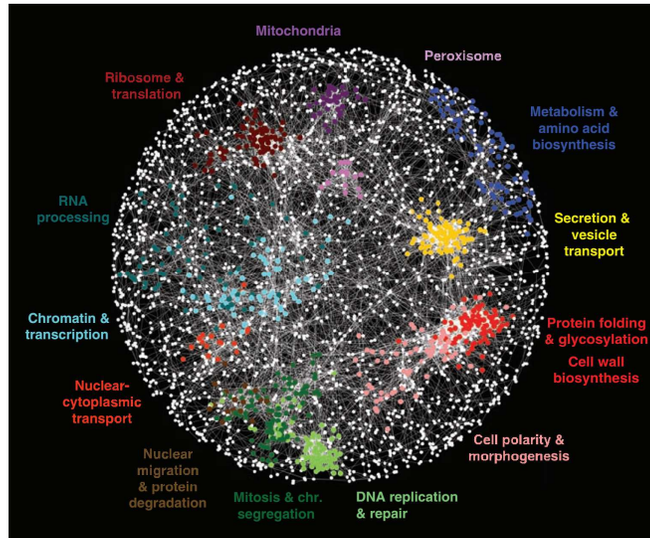
## The 1<sup>st</sup> global plant PPI map



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## Genetic interactions

5.4 million gene-gene pairs assayed for synthetic genetic interactions in yeast

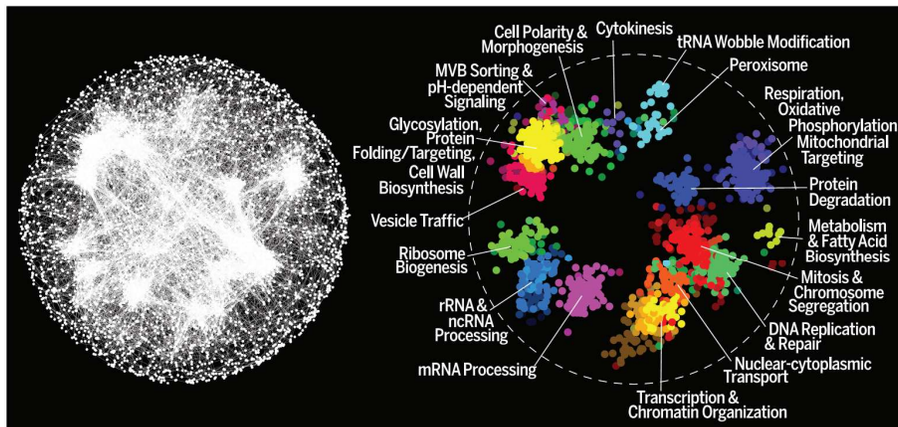


Costanzo *et al.*, *Science* 327: 425 (2010)

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## Genetic interactions, the 2016 version

23 million gene-gene pairs assayed for synthetic genetic interactions in yeast, identifying ~550,000 negative and ~350,000 positive genetic interactions

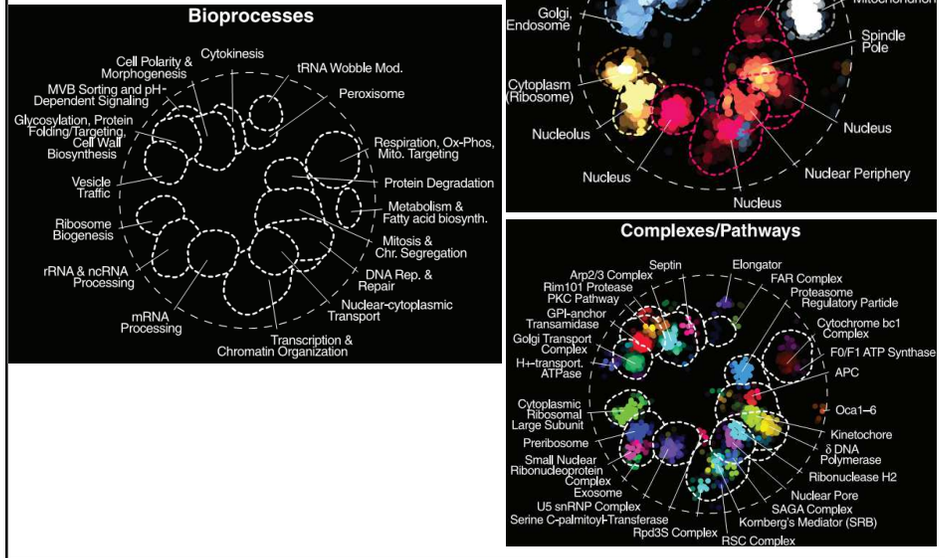


**A global network of genetic interaction profile similarities.** (Left) Genes with similar genetic interaction profiles are connected in a global network, such that genes exhibiting more similar profiles are located closer to each other, whereas genes with less similar profiles are positioned farther apart. (Right) Spatial

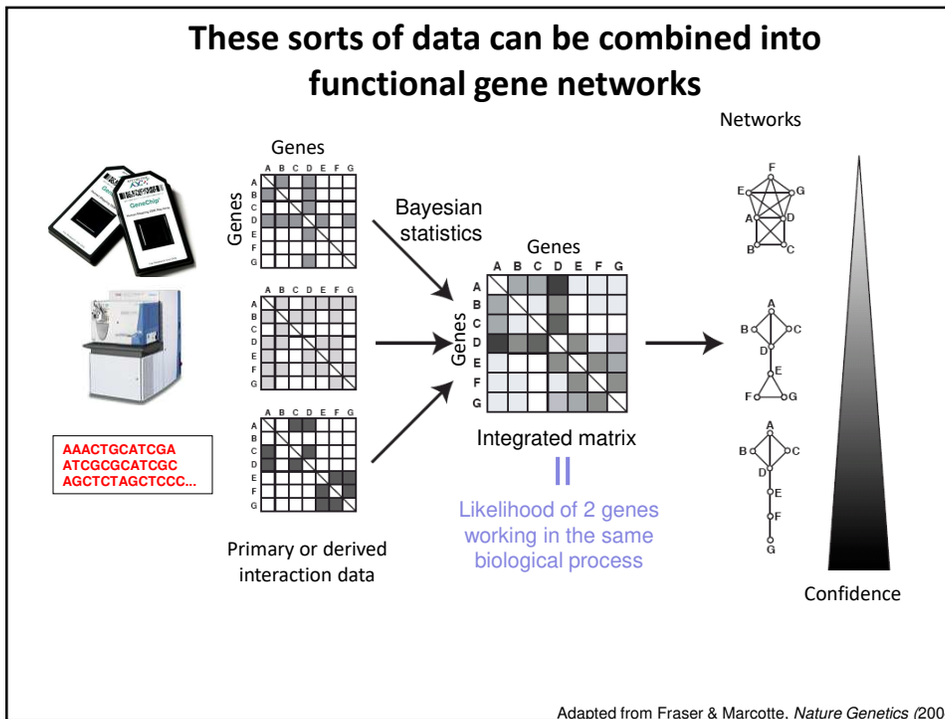
Costanzo *et al.*, *Science* 353: 1381 (2016)

22

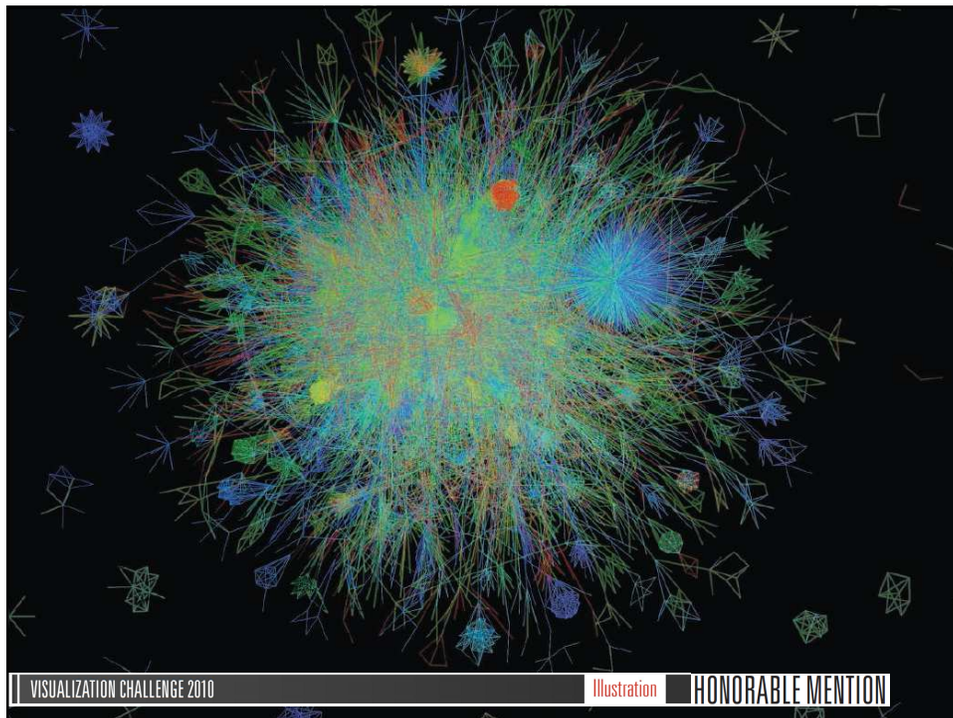
The global genetic interaction profile similarity network reveals a hierarchy of cellular function.



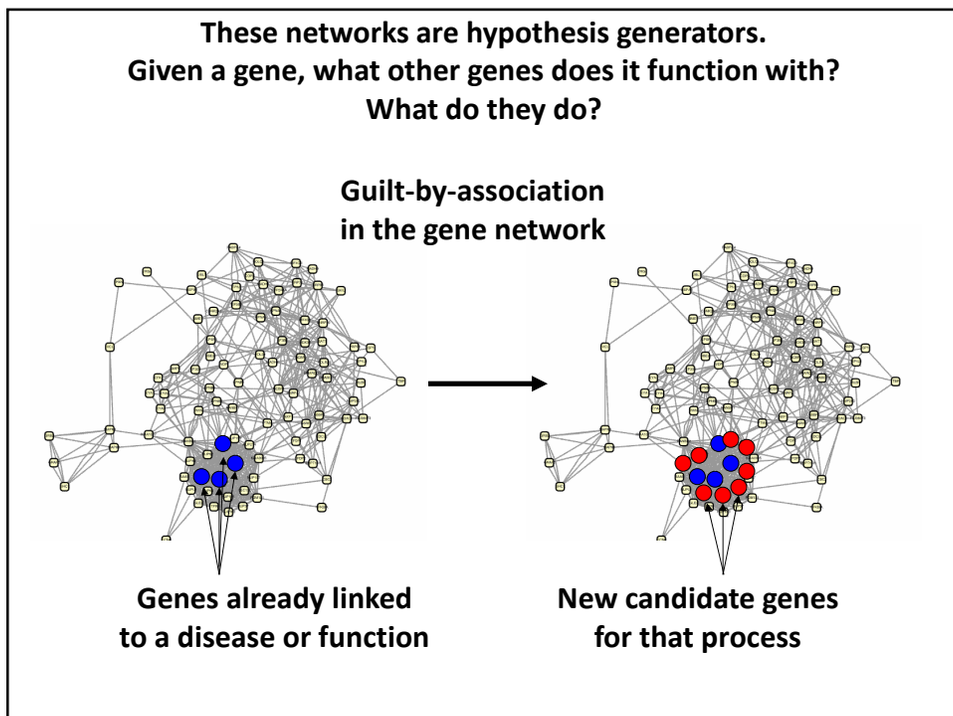
23



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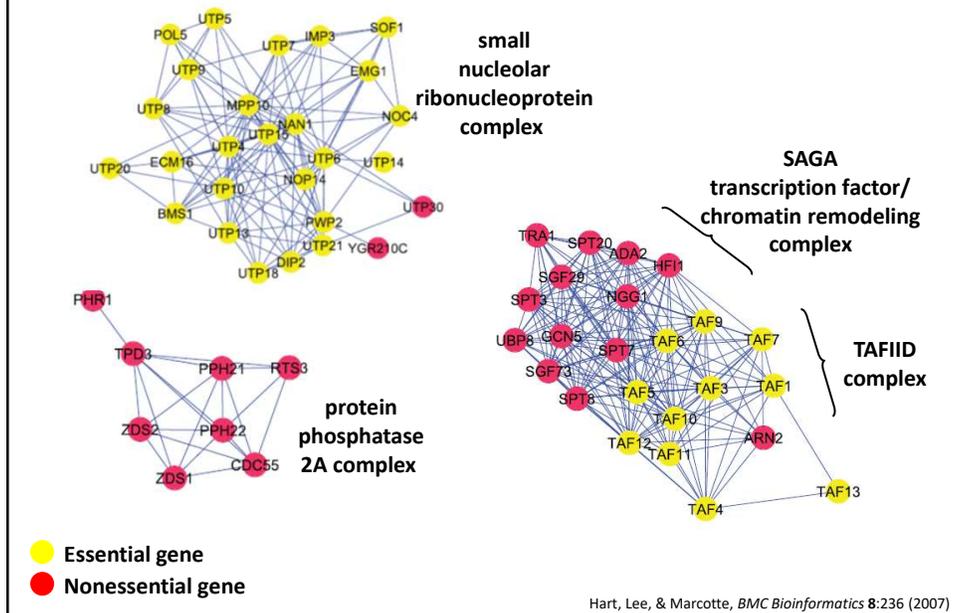


25



26

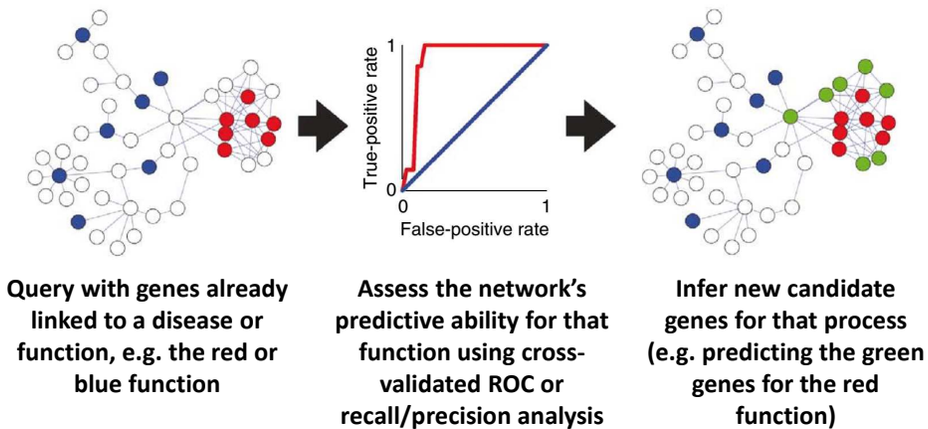
Gene networks frequently reflect functions, pathways, & phenotypes, e.g., lethality in yeast is linked to the molecular machine, not the gene



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We can propagate annotations across the graph to infer new annotations for genes (network “guilt-by-association”, or GBA).

Testing how well this works on hidden, but known, cases let’s us measure how predictive it will be for new cases.



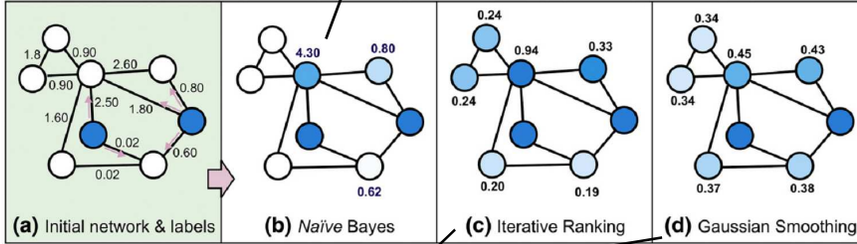
Lee, Ambaru *et al. Nature Biotechnology* 28:149-156 (2010)

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## Numerous algorithms exist for network GBA

Similar to Google's personalized PageRank

Naïve Bayes assigns scores to neighboring nodes based on edges



Network diffusion algorithms start with initial annotations and the graph topology, then propagate initial scores across the network, e.g. Gaussian smoothing tries to find scores:

$$f^{final} = \underset{f}{\operatorname{argmin}} \alpha \sum_i (f_i - f_i^0)^2 + (1-\alpha) \sum_i \sum_j w_{ij} (f_i - f_j)^2$$

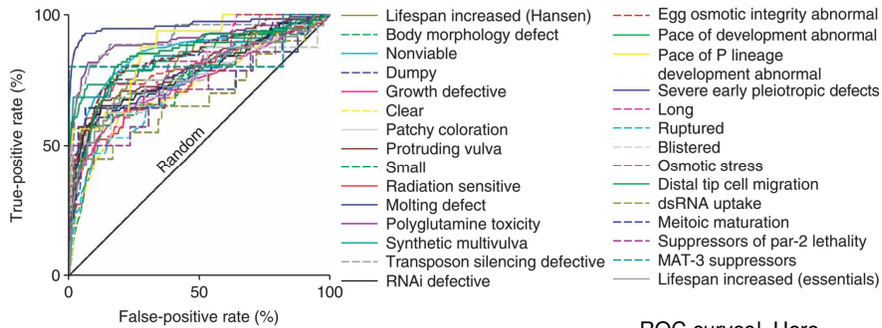
minimizing the difference between final and initial scores of a protein

& between a protein's score and that of each of its neighbors

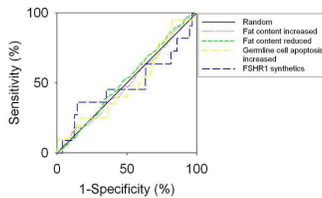
Reviewed in Wang & Marcotte, *J Proteomics* (2010)

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## For example, predicting genes linked with worm phenotypes in genome-wide RNAi screens



Some very poorly predicted pathways:



ROC curves! Here, indicating the likely predictive power of the network for a system of interest, independent of how big the system is.

A poor ROC  $\rightarrow$  no better than random guessing.

Lee, Lehner *et al.*, *Nat Genet*, 40(2):181-8 (2008)

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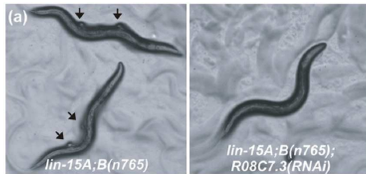
**Remarkably, this strategy works quite well**

Some examples of network-guided predictions:

**In worms:**

Genes that can reverse 'tumors' in a nematode model of tumorigenesis

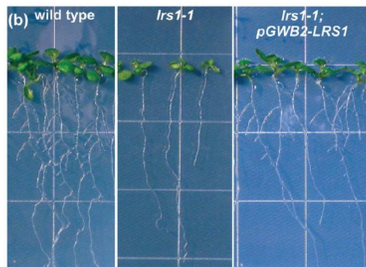
Lee, Lehner *et al.* *Nature Genetics* (2008)



**In Arabidopsis:**

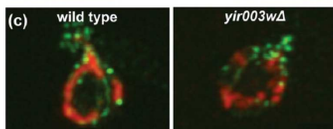
New genes regulating root formation

Lee, Ambaru *et al.* *Nature Biotech* (2010)



**In yeast:** New mitochondrial biogenesis genes

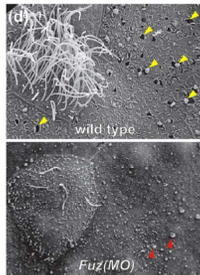
Hess *et al.*, *PLoS Genetics* (2009)



**In mice/frogs:**

Functions for a birth defect gene

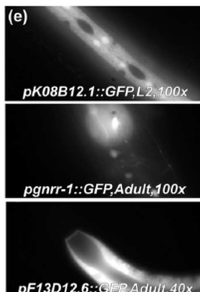
Gray *et al.*, *Nature Cell Biology* (2009)



**In worms:**

Predicting tissue specific gene expression

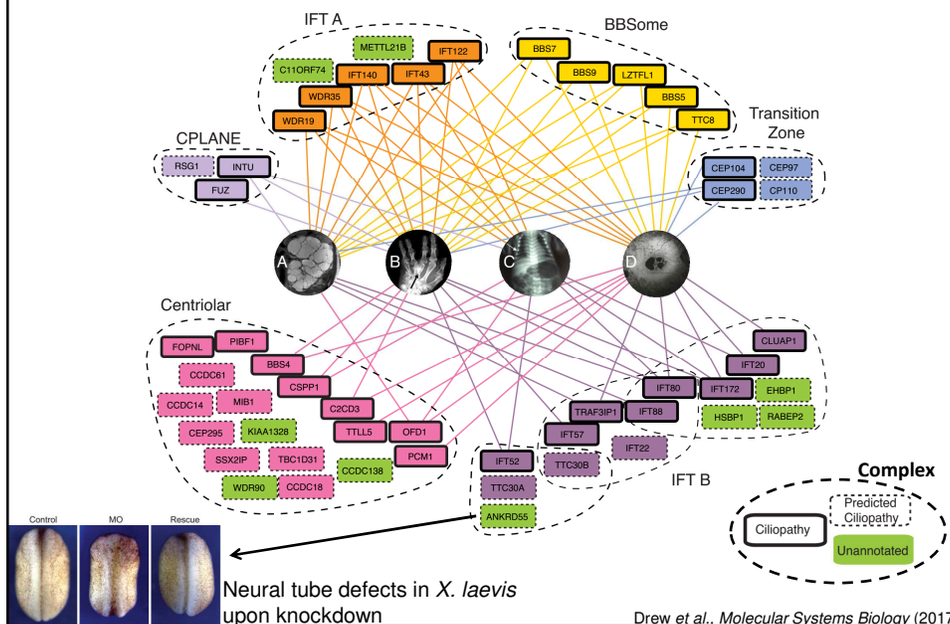
Chikina *et al.*, *PLoS Comp Biology* (2009)



Reviewed in Wang & Marcotte. *J Proteomics* (2010)

31

**We use this approach routinely in the lab, e.g. a recent example predicting new ciliopathy genes from protein complexes**



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**The New York Times**

## Scientists Identify 69 Drugs to Test Against the Coronavirus

Two dozen of the medicines are already under investigation. Also on the list: chloroquine, a drug used to treat malaria.

**A SARS-CoV-2-Human Protein-Protein Interaction Map Reveals Drug Targets and Potential Drug-Repurposing**  
 Gordon *et al.*, *bioRxiv*, posted March 22, 2020  
 doi:10.1101/2020.03.22.002386



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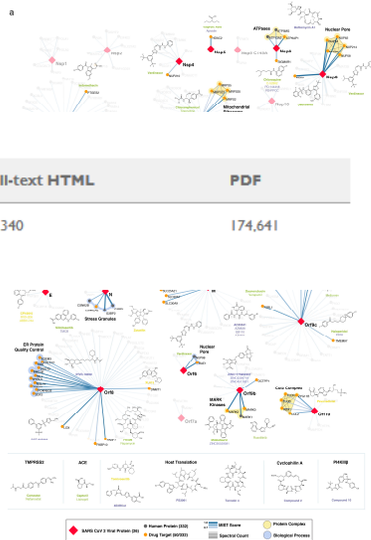
**Picked up by 82 news outlets**

A worker checking the production of chloroquine phosphate in China last month. There has been "anecdotal evidence" that chloroquine, a drug used to treat malaria, might work against the coronavirus. FeatureChina, via Associated Press

By Carl Zimmer

Published March 22, 2020 Updated April 9, 2020

Nearly 70 drugs and experimental compounds [may be effective in treating the coronavirus](#), a team of researchers reported on Sunday night.



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# Live demo of STRING, BioGRID, GeneMania, functional networks and Cytoscape

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