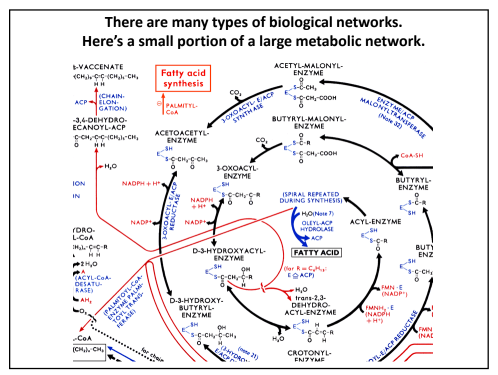
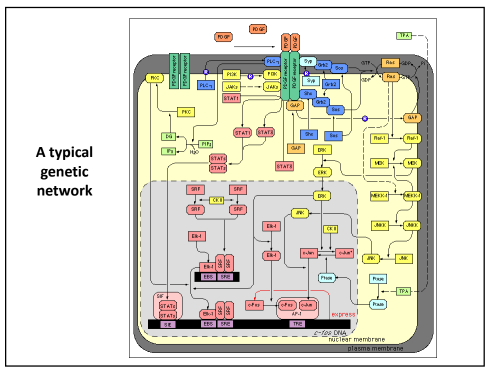
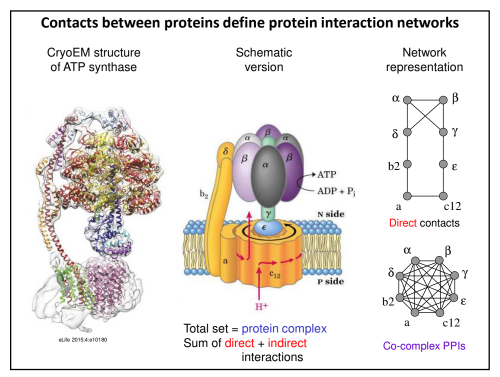
Network biology (& predicting gene function)

BCH394P/364C Systems Biology / Bioinformatics
Edward Marcotte, Univ of Texas at Austin

1







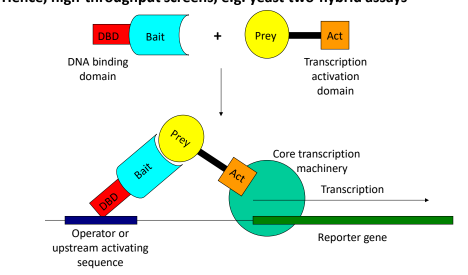
Let's look at some of the types of interaction data in more detail.

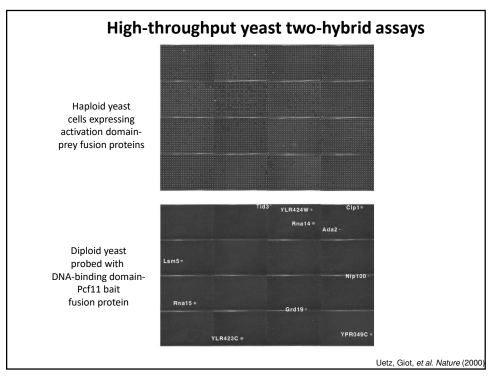
Some of these capture physical interactions, some genetic, some informational or logical.

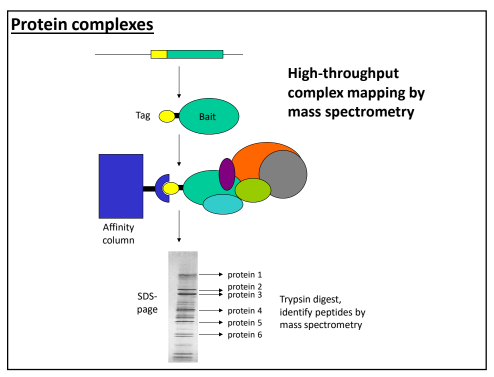
5

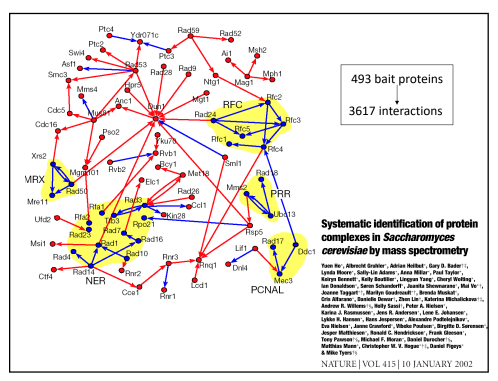
Pairwise protein interactions

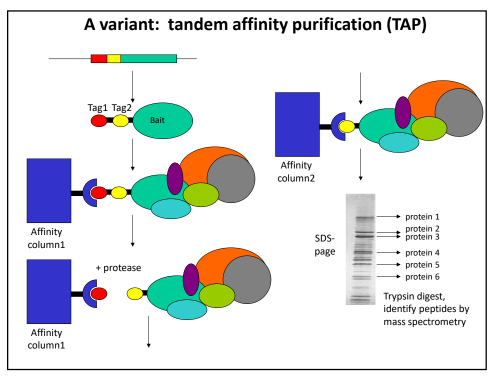
In general, purifying proteins one at a time, mixing them, and assaying for interactions is far too slow & laborious. We need something faster! Hence, high-throughput screens, e.g. yeast two-hybrid assays

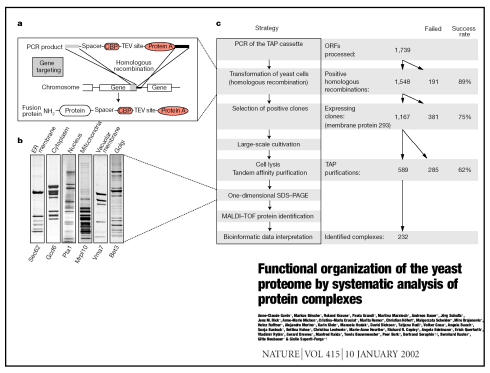


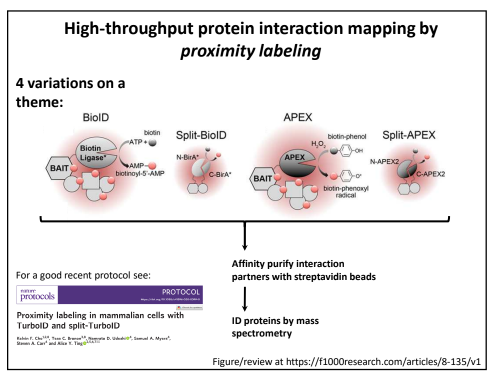


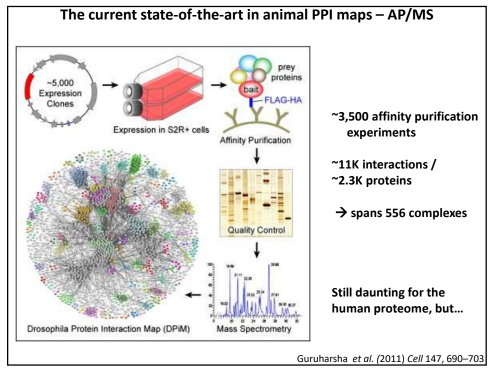


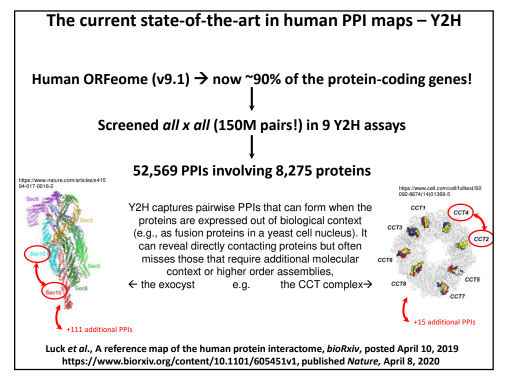


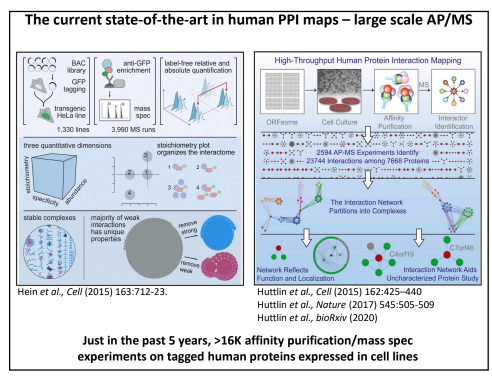


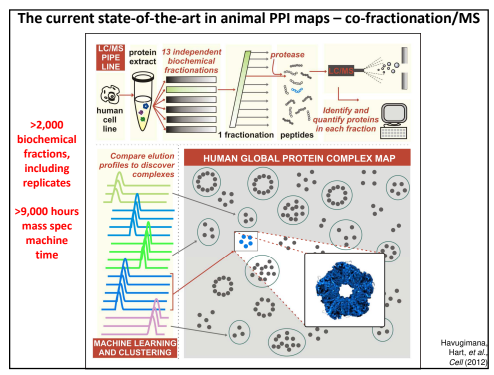


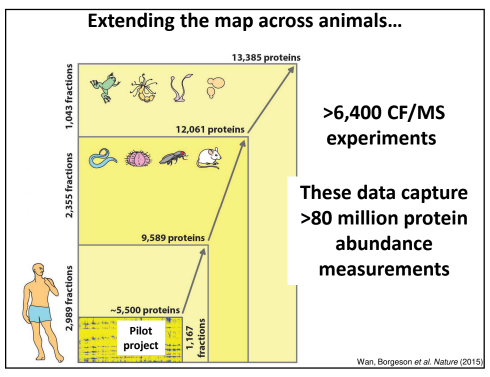


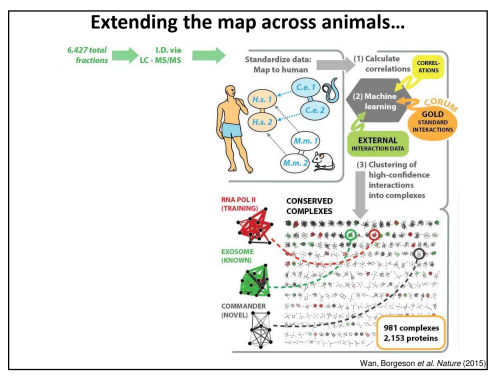


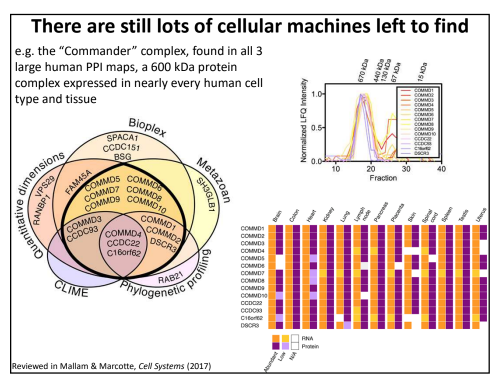


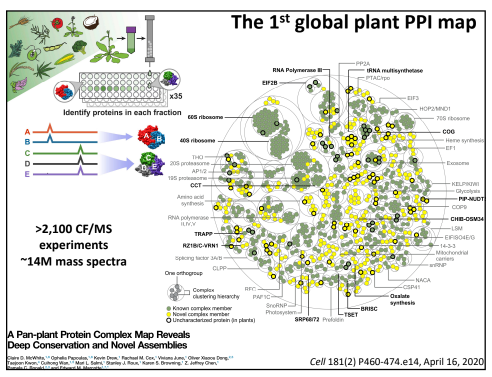






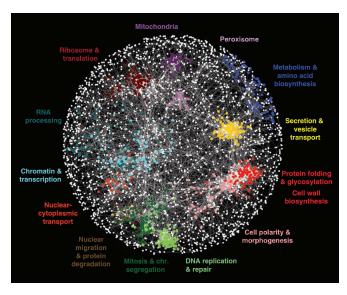






Genetic interactions

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

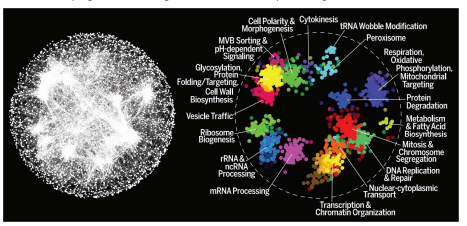


Costanzo et al., Science 327: 425 (2010)

21

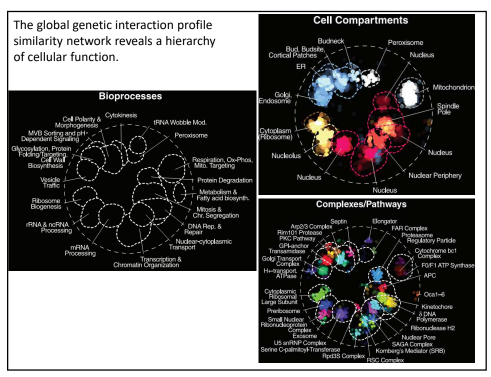
Genetic interactions, the 2016 version

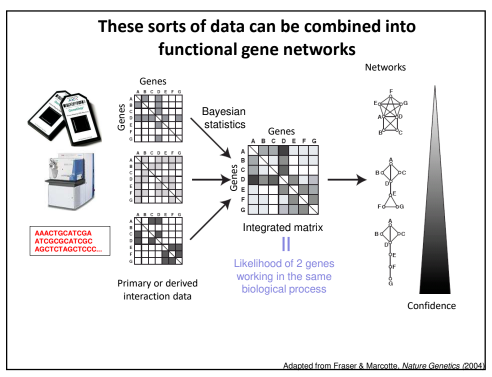
23 million gene-gene pairs assayed for synthetic genetic interactions in yeast, identifying ~550,000 negative and ~350,000 positive genetic interactions

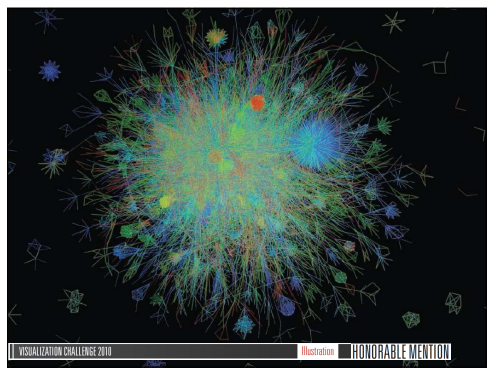


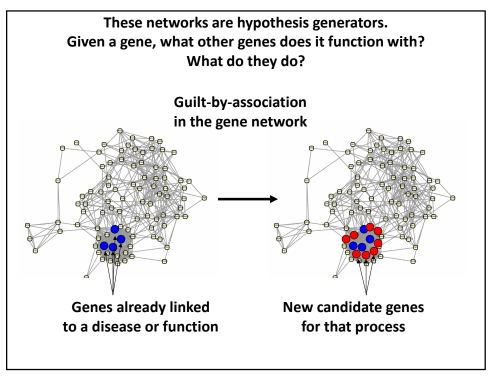
A global network of genetic interaction profile similarities. (Left) Genes with similar genetic interaction profiles are connected in a global network, such that genes exhibiting more similar profiles are located closer to each other, whereas genes with less similar profiles are positioned farther apart.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

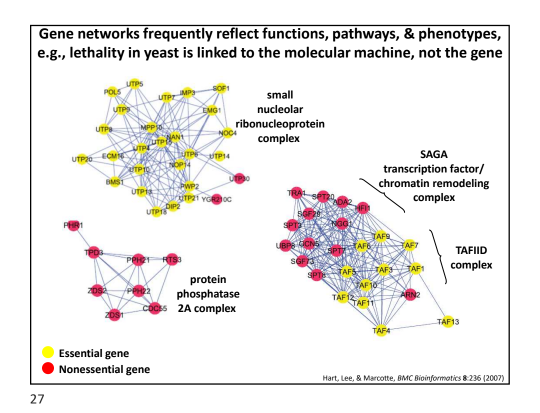
Costanzo et al., Science 353: 1381 (2016)

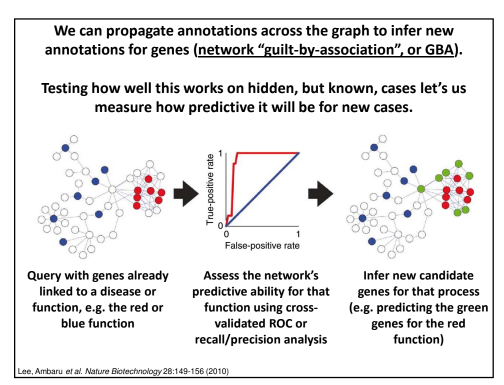


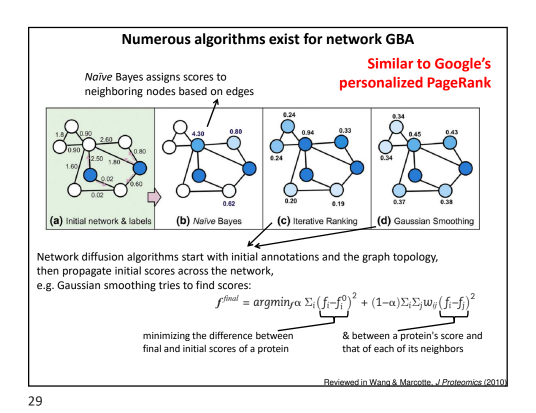












For example, predicting genes linked with worm phenotypes in genome-wide RNAi screens 100 Lifespan increased (Hansen) Egg osmotic integrity abnormal Body morphology defect Pace of development abnormal Pace of P lineage development abnormal Nonviable Dumpy True-positive rate (%) Growth defective Severe early pleiotropic defects Long Ruptured Clear Patchy coloration Protruding vulva Blistered Osmotic stress Small Distal tip cell migration Radiation sensitive Molting defect dsRNA uptake Meitoic maturation Polyglutamine toxicity Synthetic multivulva Suppressors of par-2 lethality Transposon silencing defective MAT-3 suppressors Lifespan increased (essentials) RNAi defective 100 50 False-positive rate (%) ROC curves! Here, indicating the likely predictive power of the 100 network for a system of Some very interest, independent of Sensitivity (%) poorly how big the system is. 50 predicted

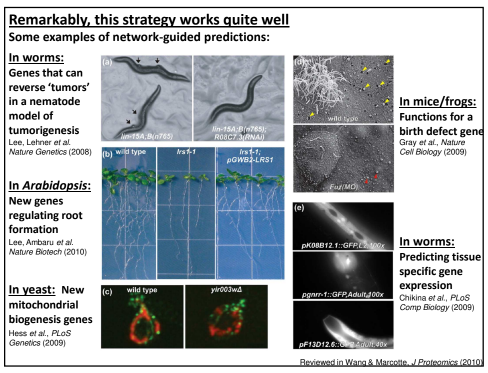
1-Specificity (%)

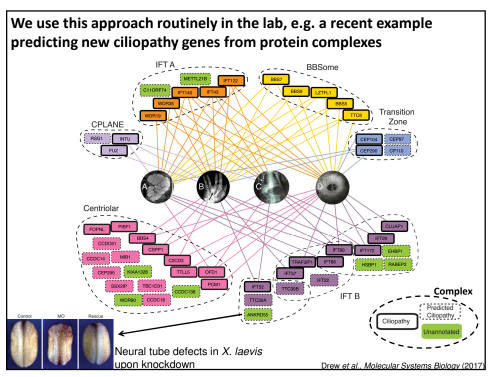
A poor ROC → no better than random guessing.

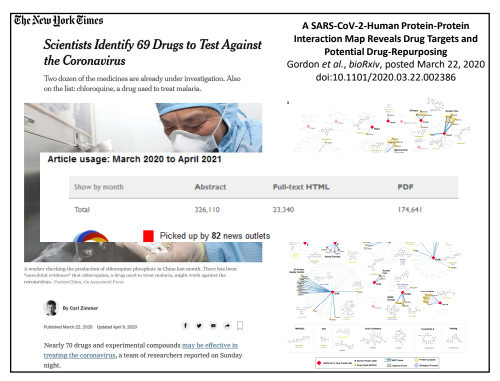
Lee, Lehner et al., Nat Genet, 40(2):181-8 (2008)

30

pathways:







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