#### Interaktionen von RNAs und Proteinen

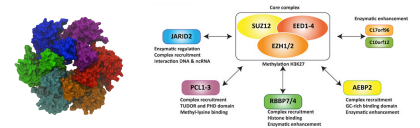
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**SS17** 

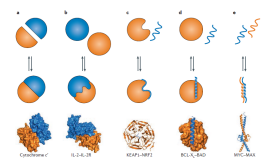
### (Multi) Protein Complexes

- two or more associated polypeptide chains
- with homologous structure/function
  formation of homo- and heterodimers to oligomers from monomers
  e.g. helicase
- with different structure/function
  built from (core) subunits and accessory proteins
  e.g. PRC2 complex
- protein complexes are a form of quaternary structure
- strong bonding is based on disulphid bridges, salt bridges, hydrophobic contacts, electon sharing



#### Protein-Protein Interactions

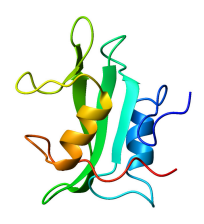
- define interaction interfaces/surfaces
- bonding is based on hydrophobic contacts, Van der Waals forces, hydrogen bonds



- a two globular proteins with preformed surfaces
- b two globular proteins with an induced binding surface
- c rigid globular protein with a peptide
- d flexible globular protein with a peptide
- e interaction of two peptides

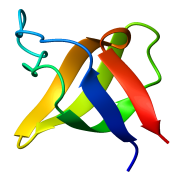
#### Protein-Protein Interaction Domain: SH2 Domain

- ▶ Src Homology 2
- around 100 aa
- $\triangleright$  2  $\alpha$ -helices and 7  $\beta$ -strands
- (one large  $\beta$ -sheet)
- known to identify a sequence of 3-6 aa
- high affinity to phosphorylated tyrosine
- ▶ function signaling
- found in about 100 human proteins



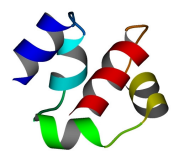
#### Protein-Protein Interaction Domain: SH3 Domain

- ► Src Homology 3
- beta-barrel fold: six β-strands forming two tightly packed anti-parallel β-sheets
- contacts proline-rich peptide sequence: -X-P-p-X-P- (X – aliphatic amino acid; p – sometimes proline)
- ▶ function signaling
- found in about 300 human proteins



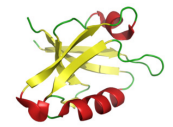
#### Protein-Protein Interaction Domain: SAM Domain

- ► Sterile Alpha Motif
- around 70 aa
- small five-helix bundle
- seems to possess the ability to bind RNA
- has two large interfaces
- can form dimers
- found in small group of genes



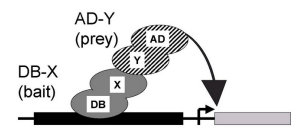
#### Protein-Protein Interaction Domain: PDZ domain

- ▶ 80-90 aa
- ▶ 5  $\beta$ -sheets, some helices
- binds to C-terminus of binding partner
   by adding a β strand to the β sk
  - by adding a  $\beta\text{-strand}$  to the  $\beta\text{-sheet}$
- multiple PDZs per protein increase specificity
- ▶ 260 PDZ in 180 human genes



# Yeast Two-Hybrid (Y2H)

test if X binds Y express fusion protein X-DB (DB ... DNA binding domain) express fusion protein Y-AD (AD ... activation domain) long linkers between X and DB, and Y and AD



### Yeast Saccharomyces cerevisiae

- genome size:  $12.5 \times 10^6$ bp
- ▶ about 5770 genes
- about 6100 proteins (about 2000 uncharacterized)
- ▶ 5100 soluble: 47% cytoplasm, 27% nucleus
- haploid and diploid living forms

## High-Throughput Yeast Two-Hybrid screening (HT-Y2H)

- ▶ a *bait* library (e.g. 192 proteins)
- theoretically any number of bait
- ▶ a *prey* library (e.g. about 6000 proteins)
- bait encodes fusion protein: DB and protein X
- prey encodes fusion protein: AD and protein Y
- array of prey-expressing haploid cells
- ▶ library of bait-expressing haploid cells
- mate bait with prey
- retrieve diploid cells expressing reporter gene due to protein-protein interaction (PPI)

It is difficulty to sample **all** possible binary combinations of proteins using the library screening methods.

## High-Throughput Yeast Two-Hybrid screening (HT-Y2H)

#### How it works: see blackboard

- ▶ 1-30 positives per bait
- only 20% true positives
- reason: self activation of reporter gene
- solution: two independent screens per bait
- even better solution: varify with other method
- found 87 baits in 281 PPI
- very low resolution
- Problem: construction of artificial fusion proteins

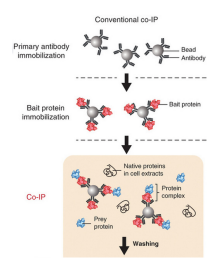
## co-Immunoprecipitation (co-IP)

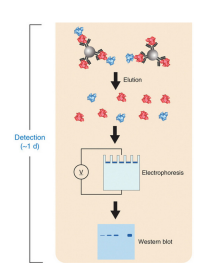
#### Co-Immunoprecipitation is not a high-throughput method.

It is used for validating PPI predictions, e.g. from HT-Y2H.

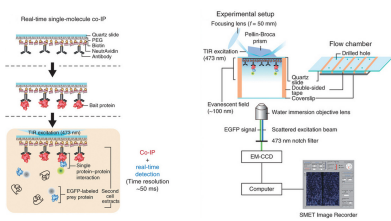
- antibodies against epitope on known protein (e.g. bait)
- pull the entire protein complex ("pull-down")
- works if proteins bind to each other tightly
- Problem: protein might hide epitope in complex
- Solution: antibody against different epitope on same protein
- Solution: antibody against proposed binding partner (double-check)
- Problem: no detection of transient interactions

## co-Immunoprecipitation (co-IP)





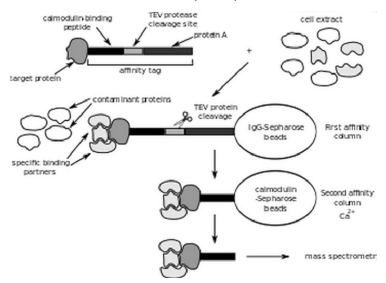
### Real-Time Single-Molecule co-IP



**Advantage**: suited for transient interactions, kinetics studies (time resolution of 50ms)

Disadvantage: requires EGFP labeling of prey

### Tandem Affinity Purification (TAP)



## Tandem Affinity Purification (TAP)

- target protein is tagged on C-terminus
- tag: calmodulin binding peptide (CBP), cleavage site, protein
  A
- ▶ 1. purification: beads with IgG bind protein A, wash
- cleave at cleavage site, elute
- ▶ 2. purification: beads with calmodulin bind CBP, wash
- analyse protein complexes with mass spectromentry

#### TAP and mass spectroscopy

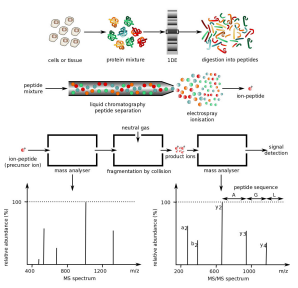
- using LC-MS, and MALDI/TOF-MS
- 4562 different tagged proteins
- 2708 proteins in 7123 PPI (2006)
- improvement!

### Mass Spectrometry - General Idea

#### Identify and quantify multiple proteins in one run

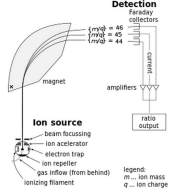
- starting with the pool of proteins
- coarse-grained protein separation by size (electrophoresis)
- protein fragmentation e.g. with trypsin
- fine-grained peptide separation with liquide chromatography (LC)
- electrospray ionization of peptides
- ▶ mass analysis → MS spectrum
- ► (collide peptides with neutral gas, even smaller fragments, mass analysis → MS/MS spectrum)
- bioinformatic analysis of MS spectrum

## Protein Mass Spectrometry



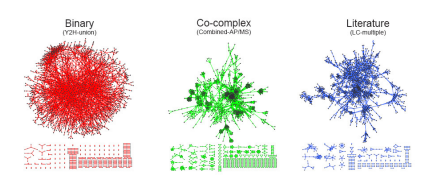
## Mass Spectrometry

- ▶ mass (m) charge (q) ratio
- peptide mass fingerprints are stored in databases
- ▶ for comparison



- some amino acids have identical masses
- utilize a sequence homology search in parallel
- ▶ problem: modifications change mass → missidentification

#### Interactome



Propose "highly significant clustering between essential proteins"

### Interactome as a Graph?

- only binary protein interactions are concidered
- intersting interactions might be transient
- only a static picture lacking dynamics and context
- A interacts with B and B interacts with C
  - at the same time?
  - in the same compartment or cell type?
  - Is there direct or indirect interaction of A and C?
  - Do A, B and C form a complex?
- would we see pathways?
- what can we infer from such a network?

#### Literature

Hong-Won Lee, Ji Young Ryu, Janghyun Yoo, Byungsan Choi, Kipom Kim, Tae-Young Yoon. *Real-time single-molecule coimmunoprecipitation of weak protein-protein interactions*. Nature Protocols 2013; 8, 2045-2060

#### For further reading:

https://www.intechopen.com/books/protein-protein-interactions-computational-and-experimental-tools