

## Module 2 overview

### *lecture*

1. Introduction to the module
2. Rational protein design
3. Fluorescence and sensors

### *lab*

1. Start-up protein eng.
2. Site-directed mutagenesis
3. DNA amplification

### **SPRING BREAK**

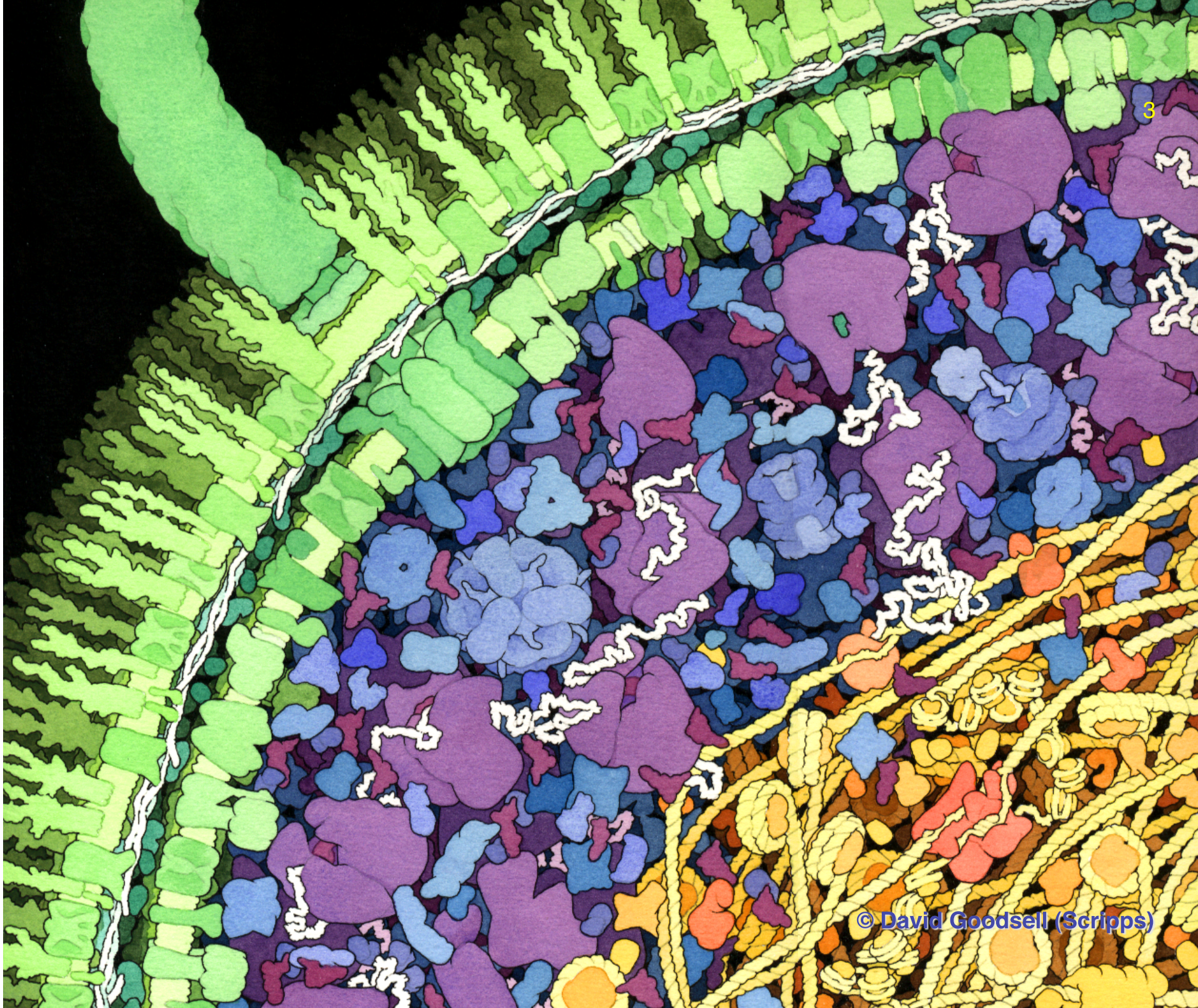
4. Protein expression
5. Purification and protein analysis
6. Binding & affinity measurements
7. High throughput engineering

4. Prepare expression system
5. Induce protein
6. Characterize expression
7. Assess protein function

## **Lecture 5: Protein purification**

- I. Standard purification methods
  - A. Harvesting and lysis
  - B. Protein separation techniques
  
- II. Assessing purified proteins
  - A. Electrophoresis
  - B. Mass spectrometry
  - C. Protein sequencing and AA analysis

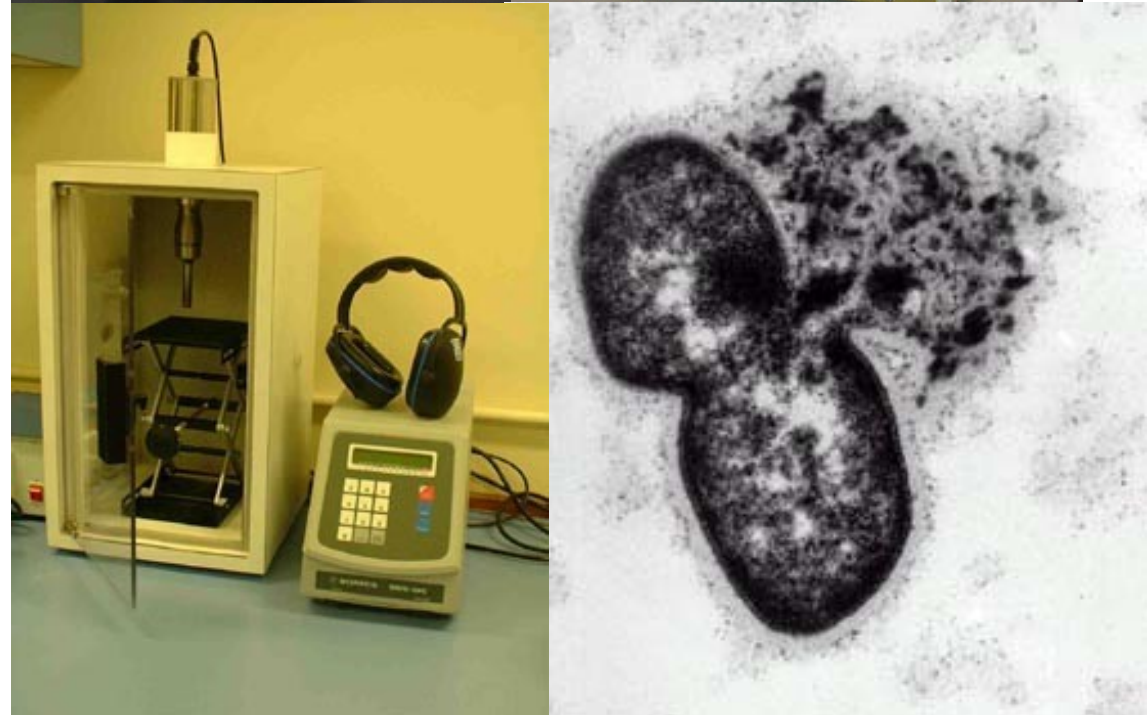




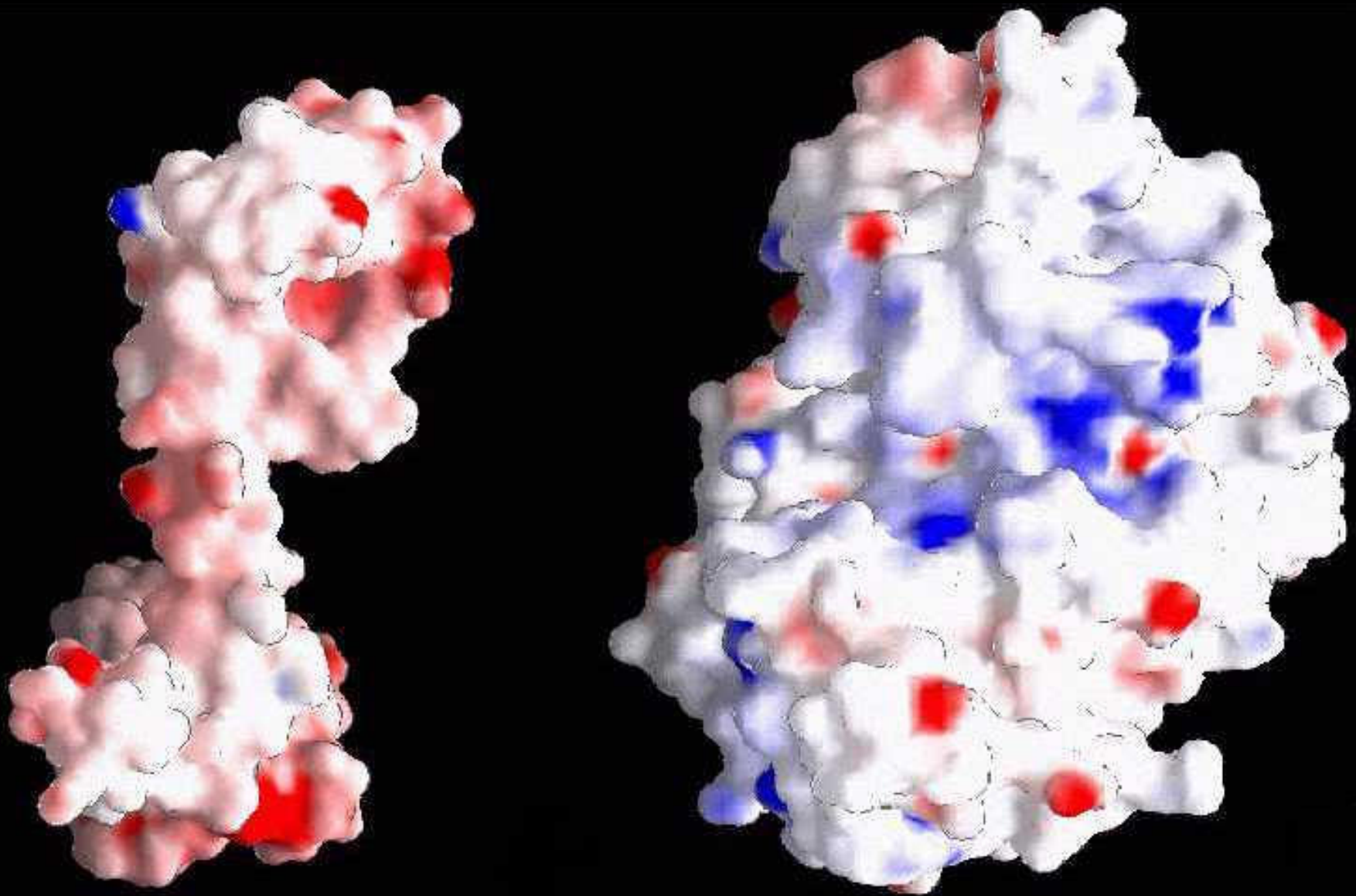
3



Once we've collected the cells, how do we get the proteins out?



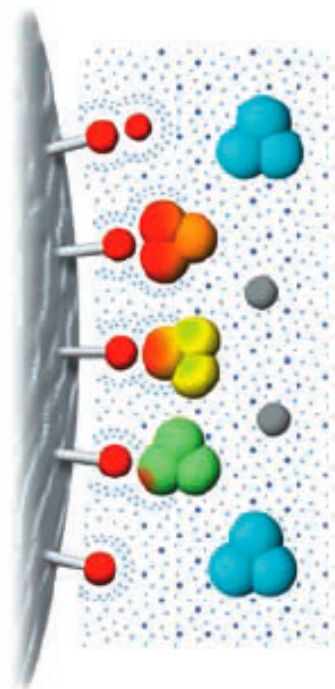
clockwise from top left:  
[lh6.ggpht.com](http://lh6.ggpht.com)  
[www.biomembranes.nl](http://www.biomembranes.nl)  
[bioinfo.bact.wisc.edu](http://bioinfo.bact.wisc.edu)  
[matcmadison.edu](http://matcmadison.edu)



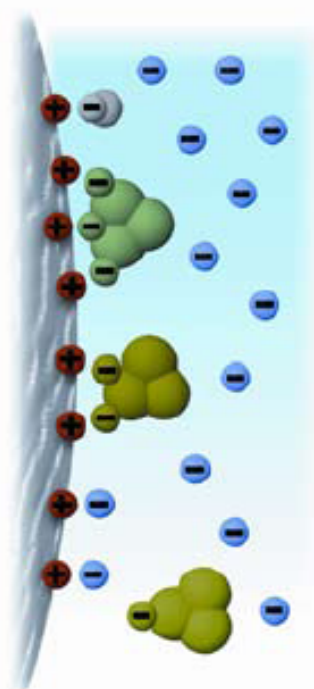
# Separation techniques



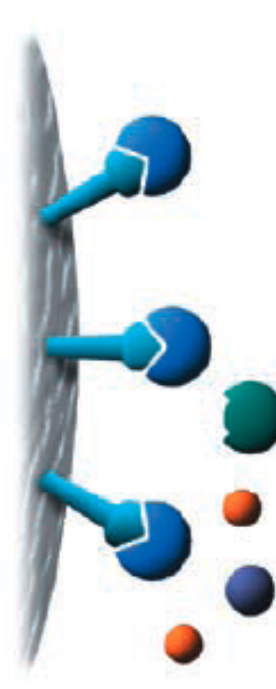
Gel filtration



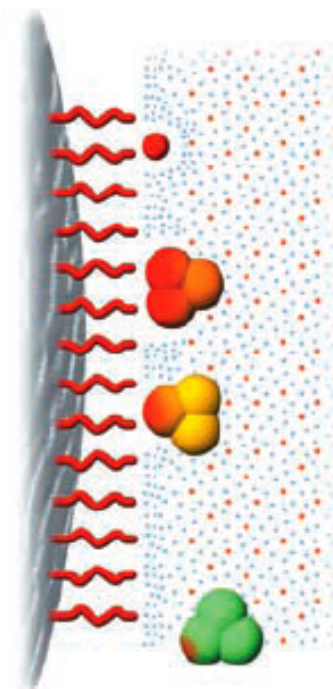
Hydrophobic interaction



Ion exchange



Affinity



Reversed phase

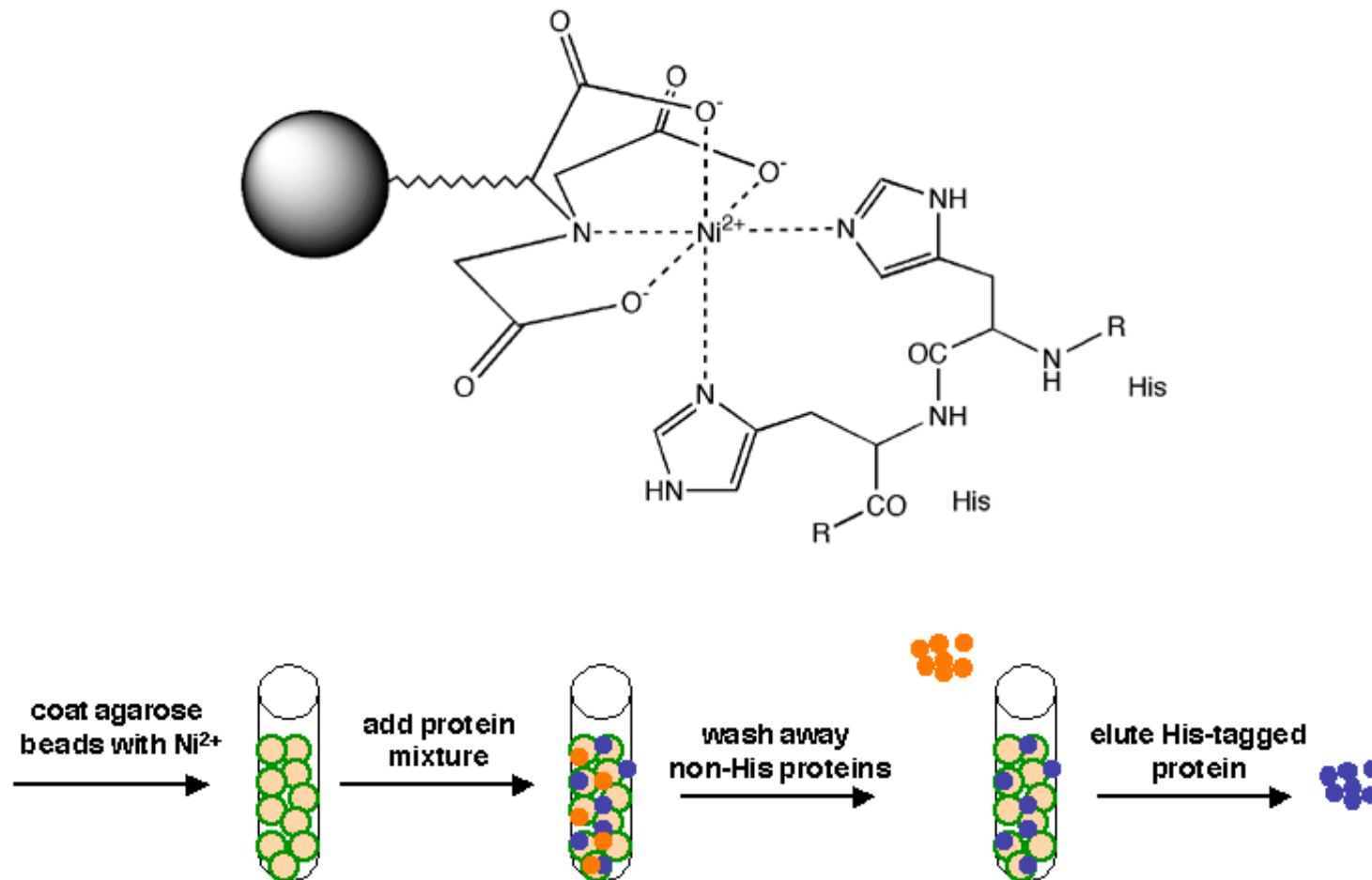
most common,  
in addition to  
affinity

*e.g.* Ni-NTA





# Nickel affinity purification with Ni-NTA agarose





Many other tags can be used for protein purification:

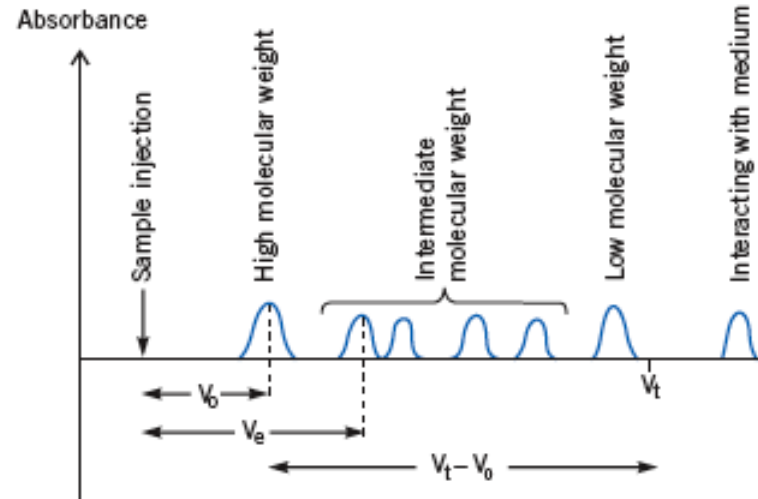
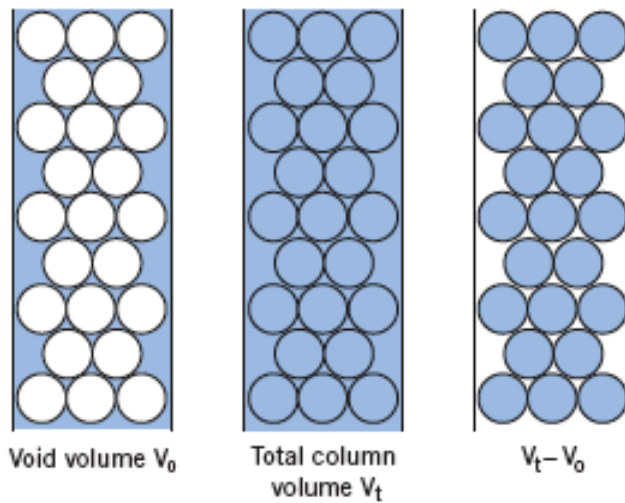
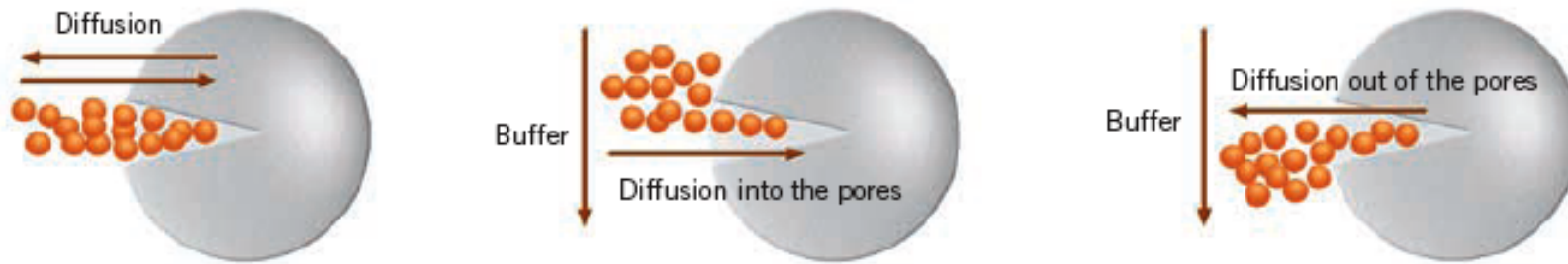
<i>tag</i>	<i>residues</i>	<i>matrix</i>	<i>elution condition</i>
poly-His	~6	Ni-NTA	imidazole, low pH
FLAG	8	anti-FLAG antibody	low pH, 2-5 mM EDTA
c-myc	11	anti-myc antibody	low pH
strep-tag	8	modified streptavidin	2.5 mM desthiobiotin
CBP	26	calmodulin	EGTA, EDTA
GST	211	glutathione	reduced glutathione
MBP	396	amylose	10 mM maltose

Tags may be chosen because they

- interfere minimally with protein structure/function
- improve recombinant protein expression or solubility
- offer most convenient purification methods

All tags may be cleaved from expressed proteins using specific proteases, if desired.

## Gel filtration (size exclusion chromatography) principle

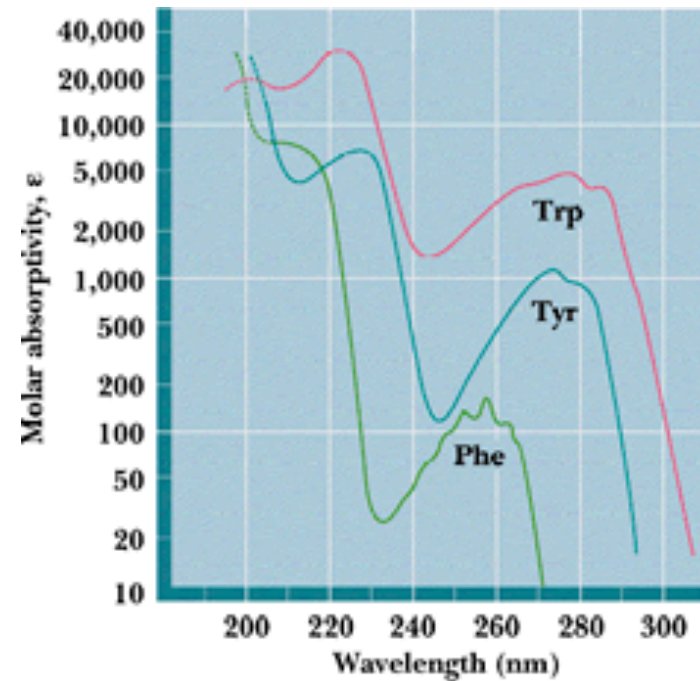




## Quantification of purified proteins

use **Beer-Lambert law**:

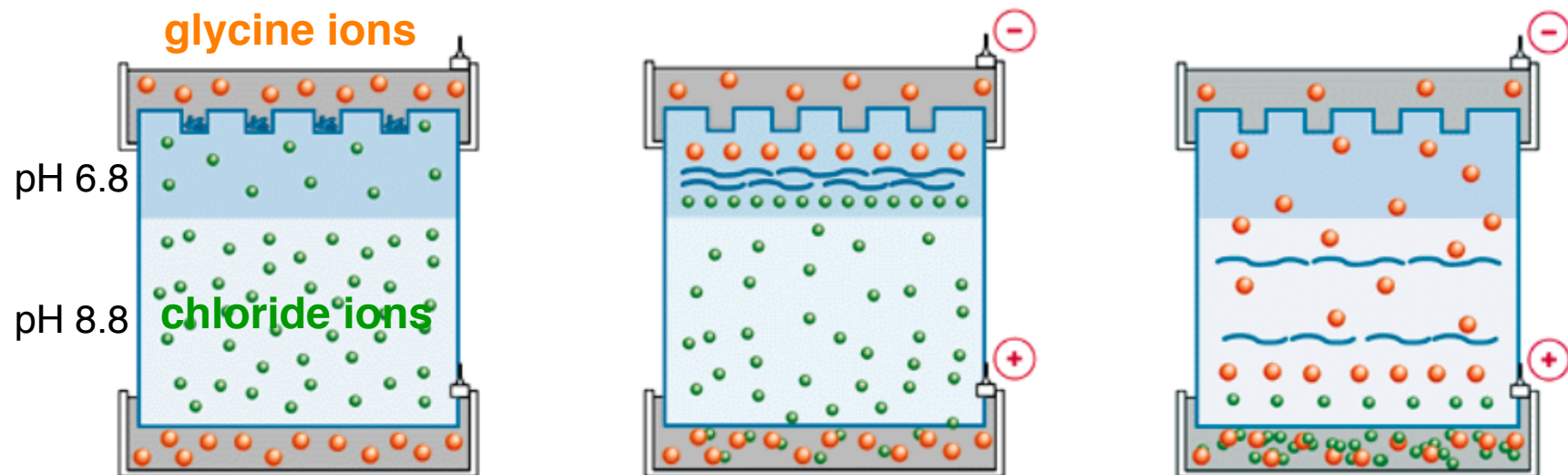
$$A_{280} = \epsilon_{280}Cl$$



## Assessing proteins for identity and purity

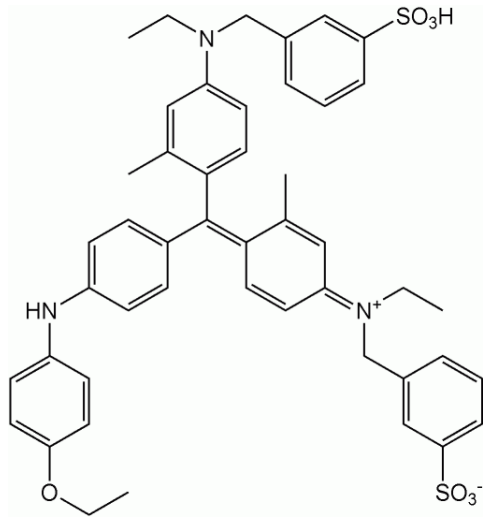
Most standard technique is sodium dodecylsulfate polyacrylamide gel electrophoresis (SDS-PAGE):

- basis is the tendency of proteins to unfold in SDS and bind a fixed amount SDS per protein (1.4 g/g)
- negative charge of SDS overwhelms protein charges
- proteins have same charge to mass ratio, but are differentially retarded by the separation gel
- stacking layer “focuses” proteins before separation layer

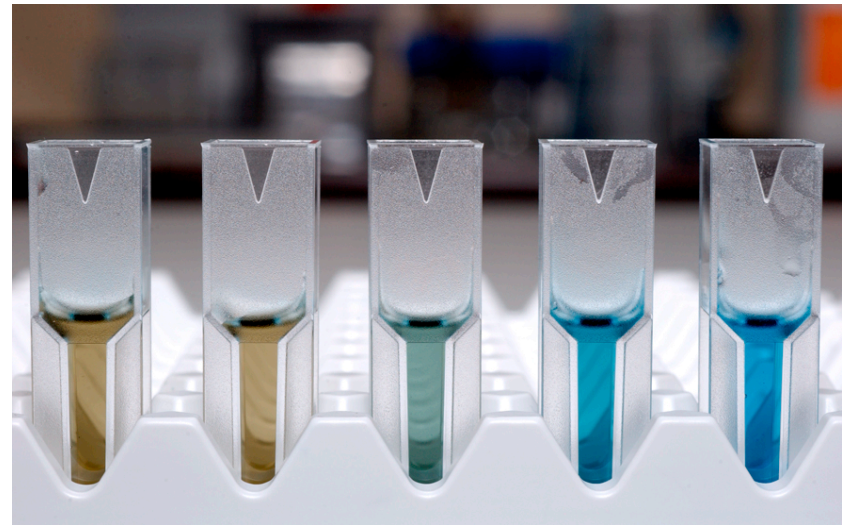
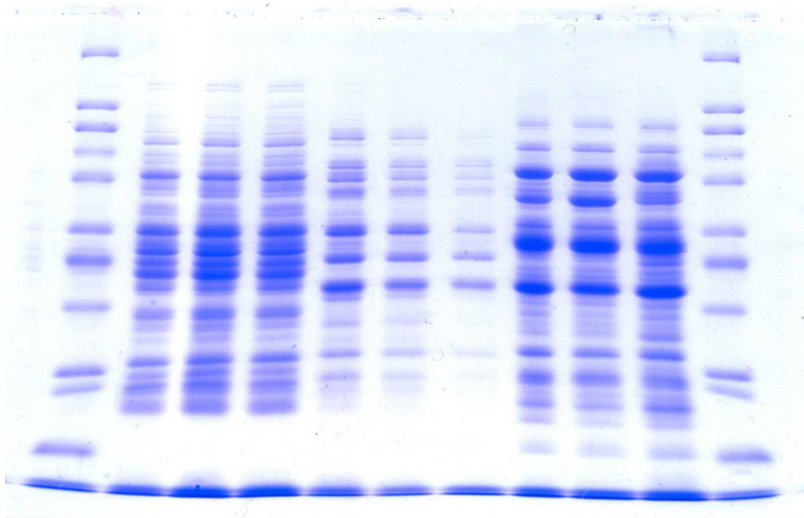




## Coomassie brilliant blue staining

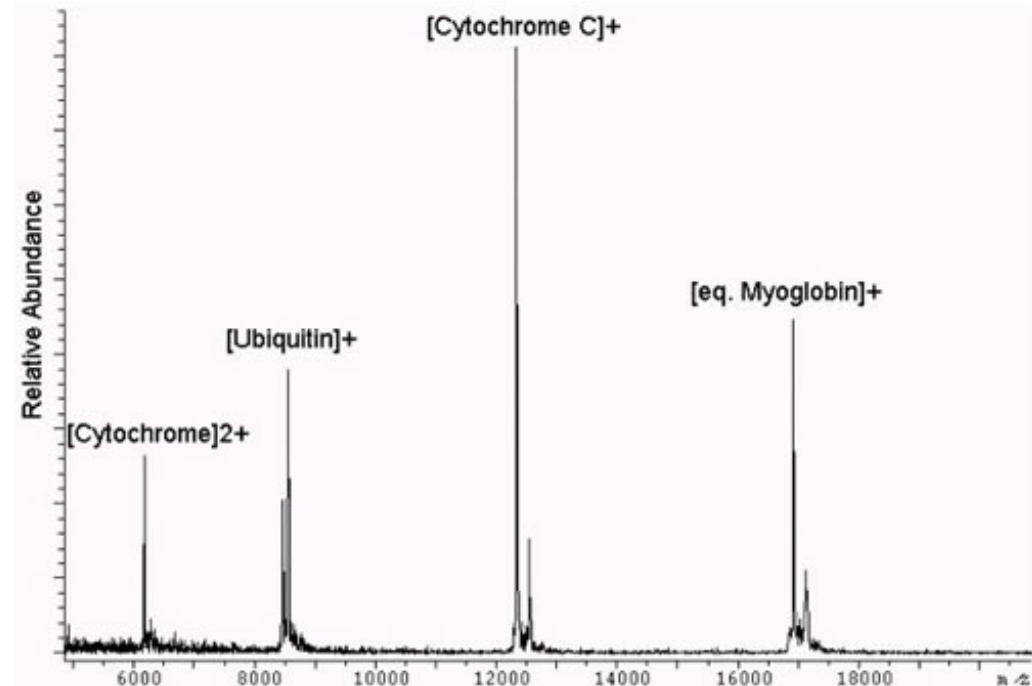
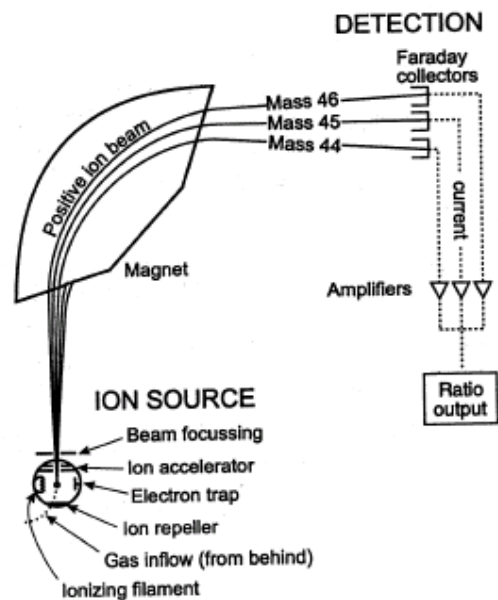


- binds proteins primarily via aromatic residues and arginine
- undergoes absorbance shift from 465 nm (brownish) to 595 nm (blue)
- basis for **Bradford Assay**; can be used to quantify proteins over ~3 kD



SDS-PAGE gives an approximate MW and purity estimate, but how can we be sure the protein we've purified is the correct one?

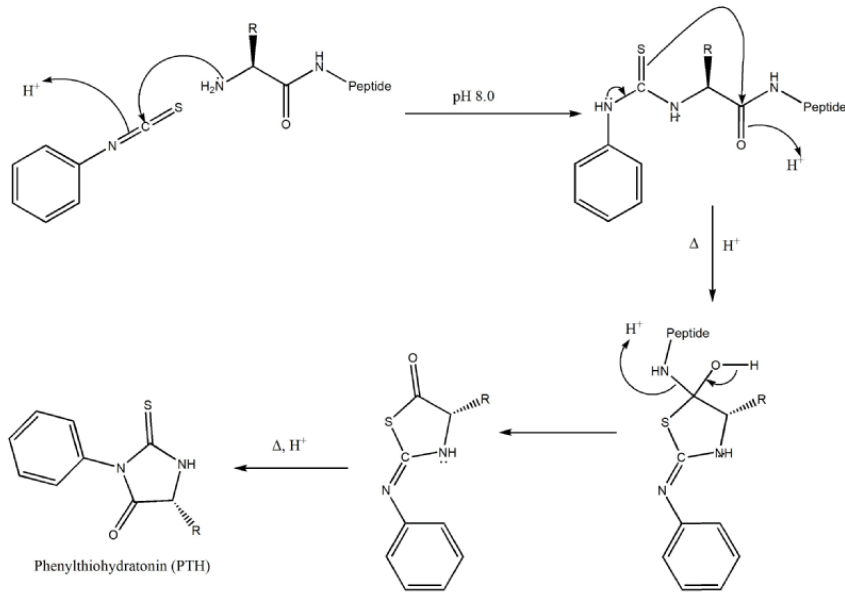
- **activity assay** if one is available
- knowledge of exact mass: **mass spectrometry**
- **N-term. sequencing** and **AA analysis**, if necessary



en.wikipedia.org/wiki/Mass\_spectrometry  
 www.kcl.ac.uk/ms-facility/images/maldispec2.jpg



## N-terminal sequencing (Edman degradation)



- products identified by chromatography or electrophoresis
- typically ~5 cycles practical for routine N-term. sequencing

[en.wikipedia.org/wiki/Edman\\_degradation](http://en.wikipedia.org/wiki/Edman_degradation)