

## 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

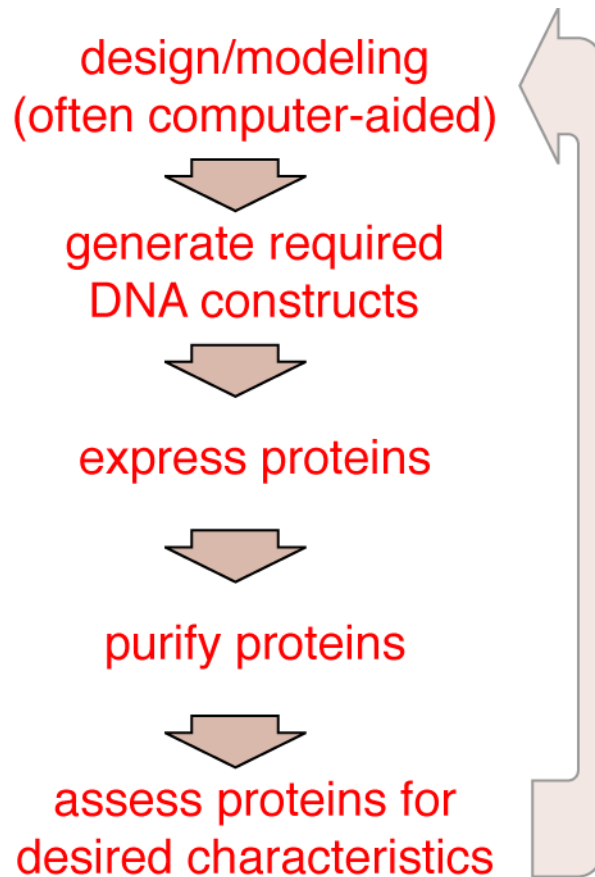
4. Prepare expression system
5. Induce protein
6. Characterize expression
7. Assess protein function

## **Lecture 7: High throughput engineering**

- I. General requirements for HT engineering
  - A. High throughput vs. rational design
  - B. Generating libraries
  
- II. Selection techniques
  - A. Phage display and related techniques
  - B. Selection for properties other than affinity

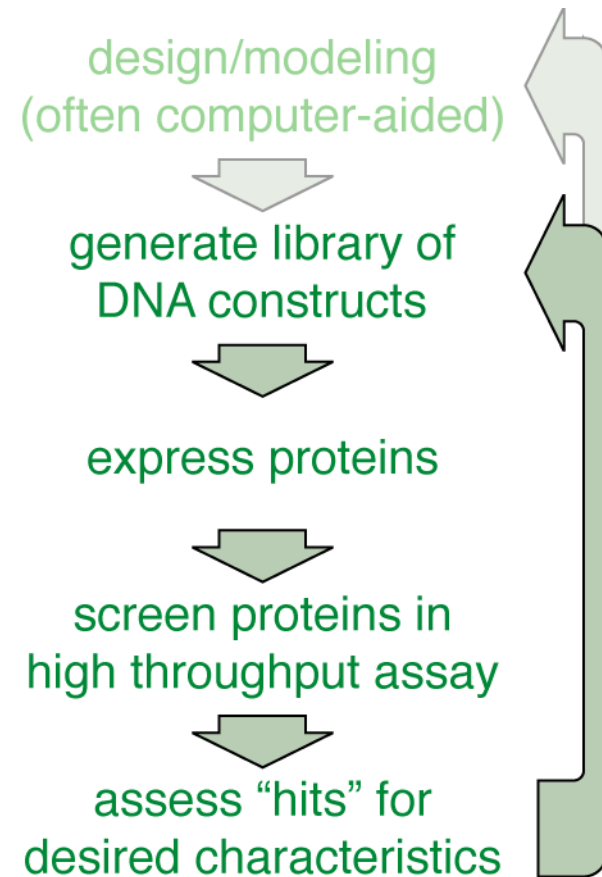
## Rational protein design:

Knowledge-based, deterministic engineering of proteins with novel characteristics



## “Irrational” high throughput protein engineering:

Selection for desired properties from libraries of random variants

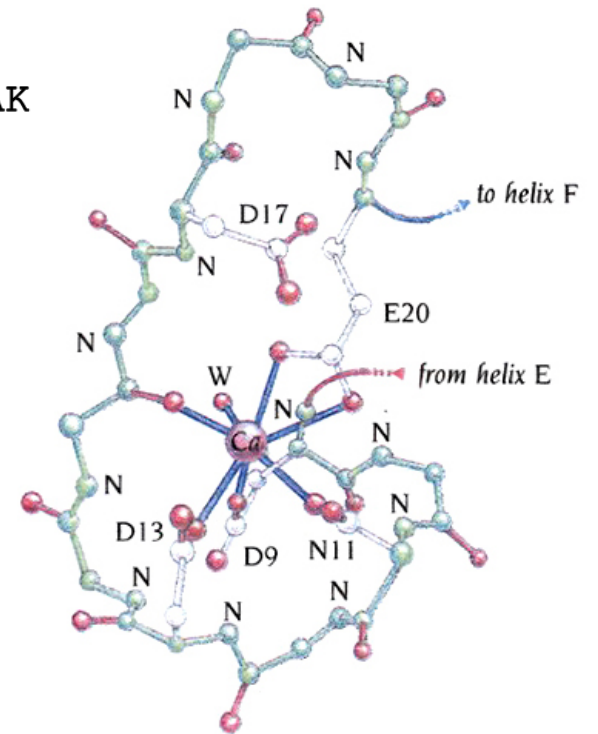


1  
 TDQLTEEQIAEFKEAFSLFDKDGDGTITTKELGTVMRSLGQNPTEAELQD  
 P WP P D ID  
 R HW R R  
 R

class mutants  
 2008-10

51  
 MINEVDADGNGTIYFPEFLTMMARKMKDSEEEIREAFRVFDKDGNGYI  
 H HPE D K P G PHHD D  
 I K E K A K P  
 H P  
 K W  
 D

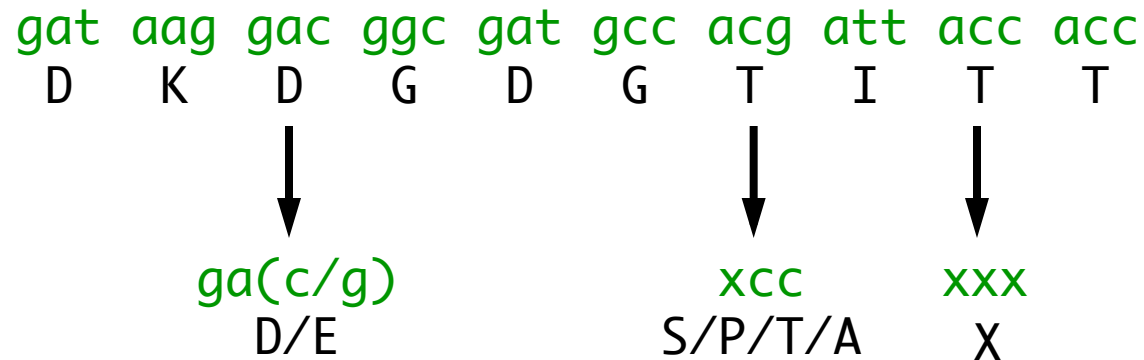
101  
 SAAQLRHVMTNLGEKLTDEEVDEMIREADIDGGQVNYEEFVQMMTAK  
 L RS S K P G R F D  
 T W  
 L



## Methods for generating mutant **genetic libraries**:

- site-directed mutagenesis with degenerate primers
- error-prone PCR
- gene shuffling

## Degenerate primers



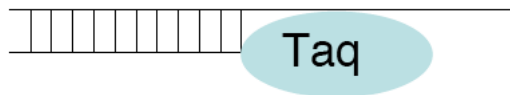
- not all combinations of AA's possible at each position
- number of combinations expands exponentially
- degenerate primers synthesized by split-pool method
- standard primer design criteria must be considered

# PCR polymerase and conditions may be chosen to **promote mutations**

Polymerase	Template doublings ( $d$ ) <sup>a</sup>	<i>lacI</i> <sup>-</sup> plaques <sup>b</sup> (% $\pm$ SD)	Mutation load <sup>c</sup> (per kilobase) ( $\pm$ SD)	Error rate <sup>d</sup> (per base) ( $\times 10^{-6} \pm$ SD)
Pfu-Pol (exo <sup>+</sup> )	12.3	0.61 $\pm$ 0.09	0.017 $\pm$ 0.002	1.4 $\pm$ 0.2
Pfu-Pol (exo <sup>-</sup> )	11.8	20 $\pm$ 1.7	0.58 $\pm$ 0.05	49 $\pm$ 4
Taq-Pol	11.6	3.9 $\pm$ 0.16	0.12 $\pm$ 0.006	10 $\pm$ 0.5

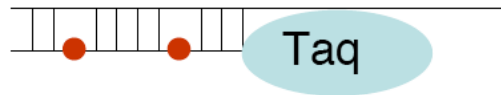
error rate = mutation load  $\div$  template doublings

## normal PCR



dCTP, dTTP  
dGTP, dATP  
Mg<sup>2+</sup>

## error-prone PCR



dCTP, dTTP  $\uparrow$   
dGTP, dATP  $\downarrow$   
Mg<sup>2+</sup>  $\uparrow$   
Mn<sup>2+</sup>

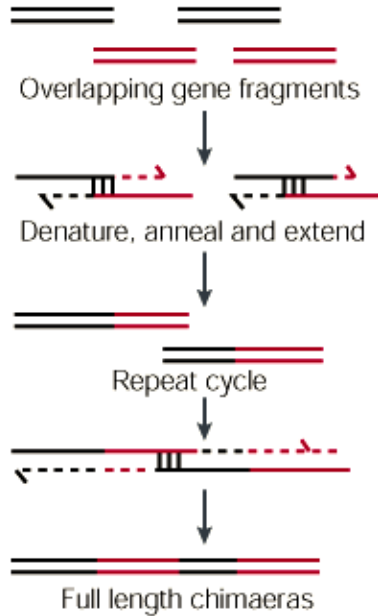
*some mutations are more likely than others*

Mutation	Pfu-Pol(exo <sup>-</sup> ) D473G <sup>a</sup>	Taq-Pol (Mn <sup>2+</sup> / unbalanced dNTPs) <sup>b</sup>	Taq-Pol (Mn <sup>2+</sup> / unbalanced dNTPs) <sup>c</sup>	Taq-Pol (unnatural mutagenic bases) <sup>d</sup>
A $\rightarrow$ T/T $\rightarrow$ A	28	40.9	11.4	0.2
A $\rightarrow$ C/T $\rightarrow$ G	7.4	7.3	3.3	8.4
A $\rightarrow$ G/T $\rightarrow$ C	19.2	27.6	60.9	78.3
G $\rightarrow$ A/C $\rightarrow$ T	22	13.6	18.1	13.2
G $\rightarrow$ C/C $\rightarrow$ G	7.3	1.4	4.3	0.7
G $\rightarrow$ T/C $\rightarrow$ A	10.3	4.5	1.8	0.0
Insertion	2.9	0.3	Not given	$\sim$ 0
Deletion	2.9	4.2	Not given	$\sim$ 0

**Gene shuffling** techniques mimic diversity due to meiotic recombination:

- fragments of homologous genes combined using “**sexual PCR**”
- diversity may arise from error prone PCR or multiple genes

**a Recursive PCR and gene assembly**

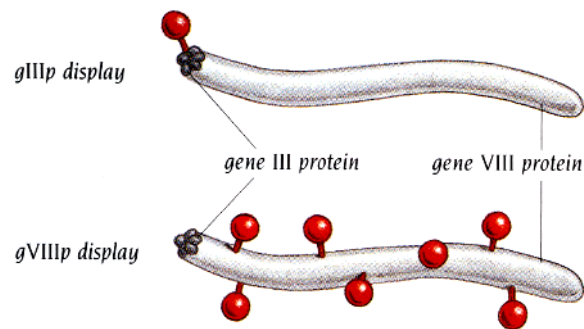


## How are libraries of mutant proteins screened?

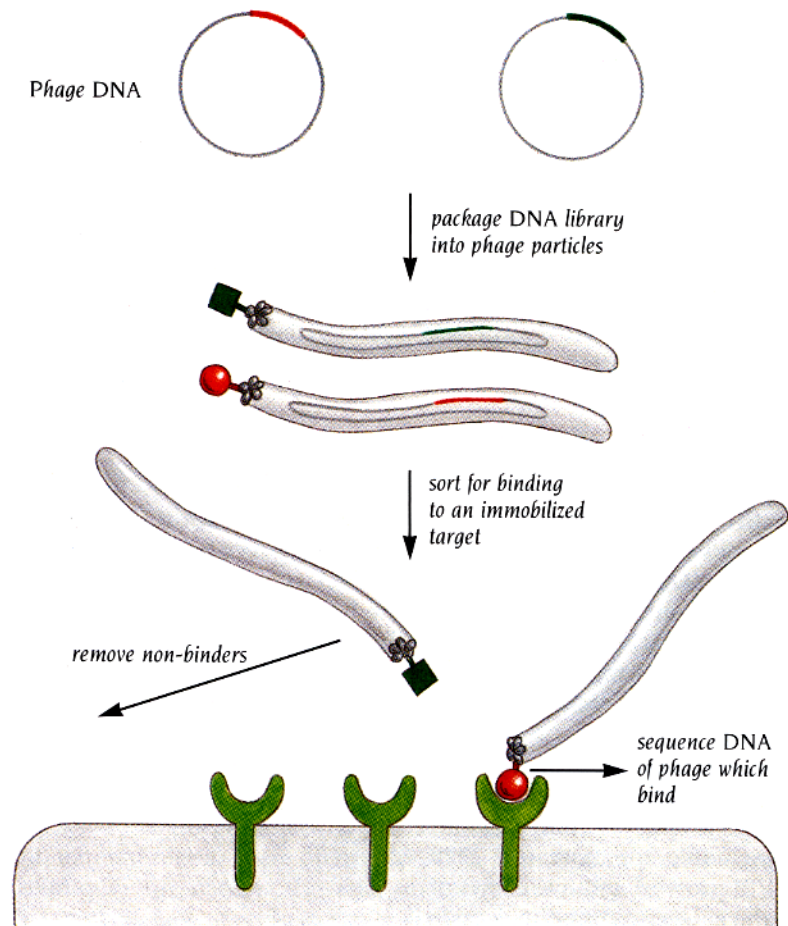
All major methods include a strategy to **keep DNA sequence info** associated with the proteins that are being screened.

Phage display is a versatile high throughput method to do this:

protein “displayed” on the coat of a bacteriophage, by fusing to a natural phage coat protein



**M13 phage  
(contains DNA)**



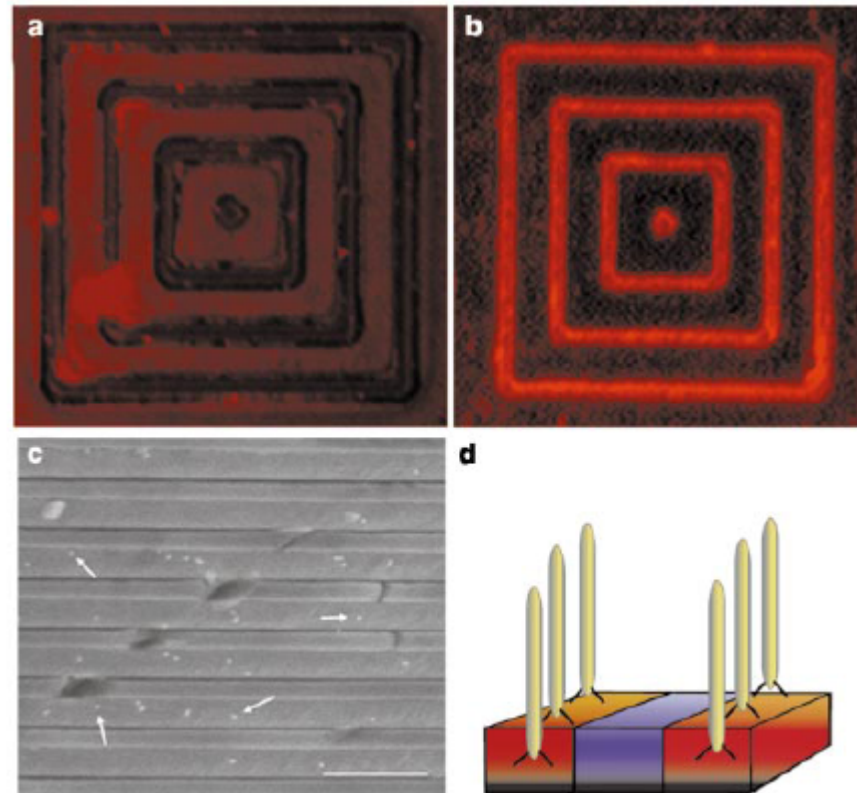


## Application: phage-displayed peptides that bind to GaAs

### selected sequences

G13-5	V	T	S	P	D	S	T	T	G	A	M	A
G12-5	A	A	S	P	T	Q	S	M	S	Q	A	P
G12-3	A	Q	N	P	S	D	N	N	T	H	T	H
G1-4	A	S	S	S	R	S	H	F	G	Q	T	D
G12-4	W	A	H	A	P	Q	L	A	S	S	S	T
G14-3	A	R	Y	D	L	S	I	P	S	S	E	S
G7-4	T	P	P	R	P	I	Q	Y	N	H	T	S
G15-5	S	S	L	Q	L	P	E	N	S	F	P	H
G14-4	G	T	L	A	N	Q	Q	I	F	L	S	S
G11-3	H	G	N	P	L	P	M	T	P	F	P	G
G1-3	R	L	E	L	A	I	P	L	Q	G	S	G

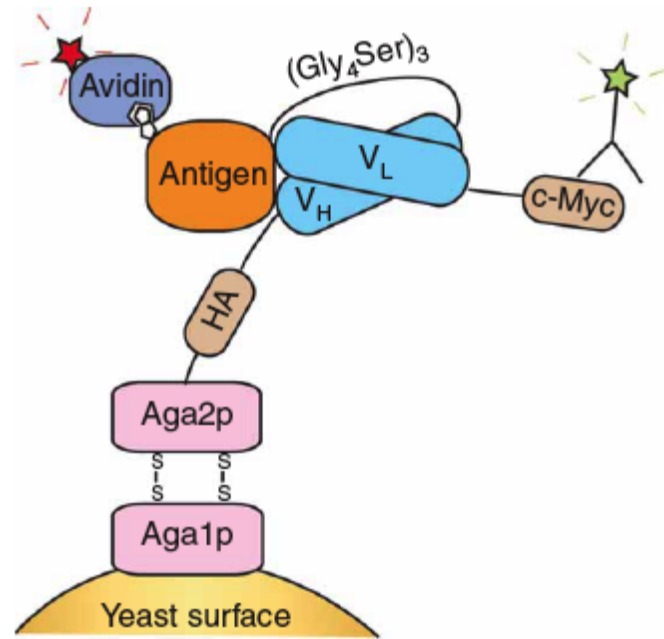
### phages patterned on target substrate



Whaley *et al.* (2000) *Nature* 405: 665-8.

Yeast display: similar to phage display, but with proteins fused to a *Saccharomyces* cell wall protein (DNA in yeast)

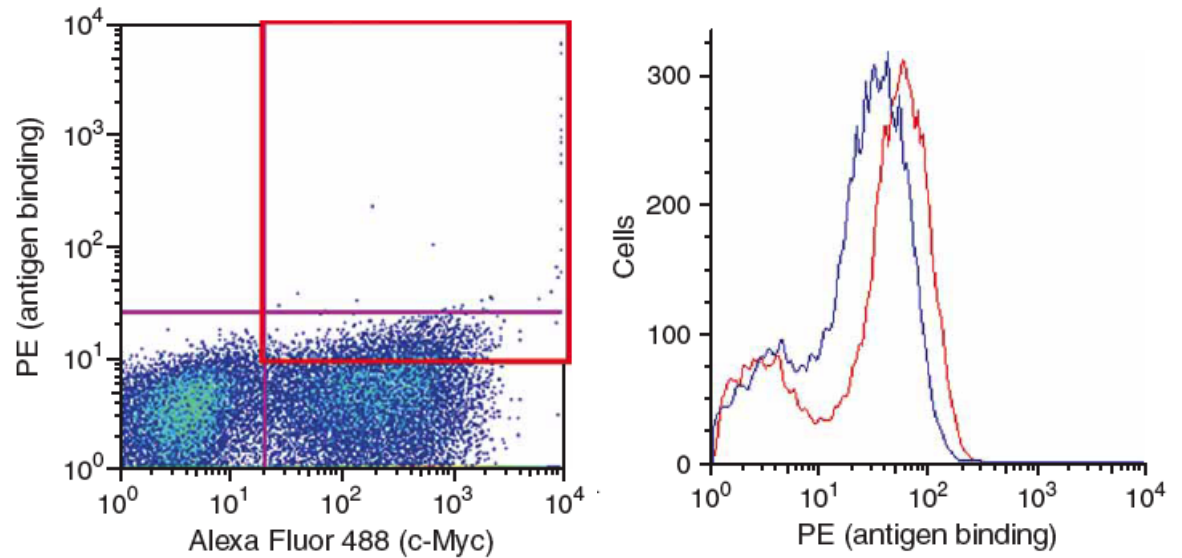
What would you expect advantages to be, compared with phage display?



In this example, a population of scFvs was screened for binding to an antigen

*left:* selection criterion for FACS assay

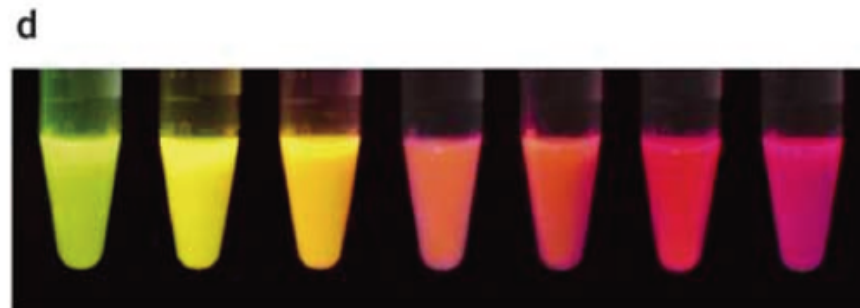
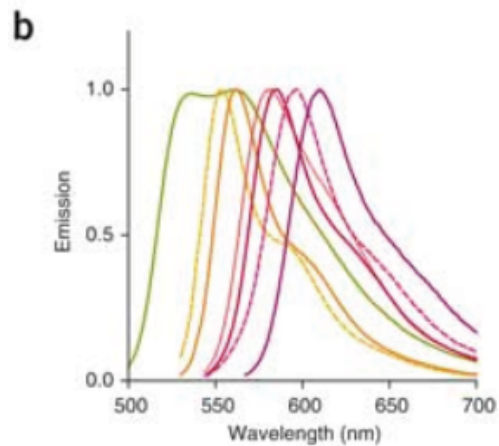
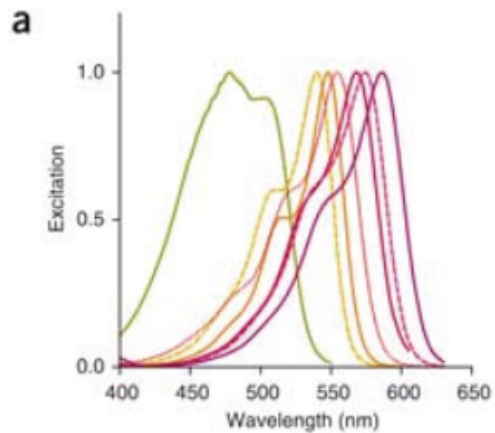
*right:* comparison of wt (blue) and selected (red) scFv binding



Why not more stringent selection?

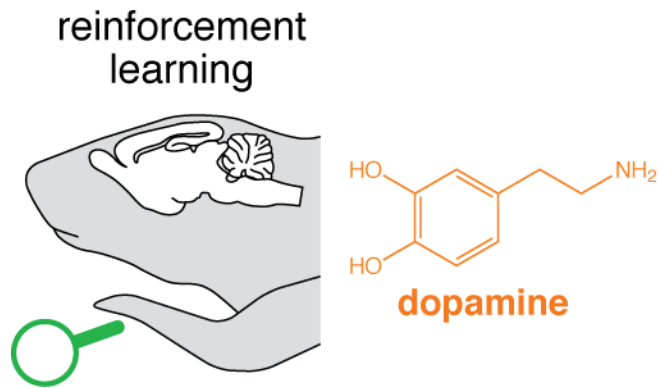
What about **properties other than affinity**?

A simple example: screen for dsRed variants with different excitation and emission wavelengths—**how could this be done**?



Shaner *et al.* (2004) *Nat. Biotechnol.* 22: 1567-72

# another example: neurotransmitter sensor for MRI



This screen only involved ~500 variants/round; **under what circumstances would you expect this level of throughput to be successful?**

## Which type of screening method to use?

<u>screen method</u>	<u>throughput</u>	<u>other notes</u>
SELEX	$10^{15}$	selection of DNA/RNA
ribosome display	$10^{15}$	<i>in vitro</i> protein synthesis
phage display	$10^{11}$	best for small proteins/peptides
yeast display	$10^8$	compatible w/eukar. proteins
plate assays	$< 10^5$	versatile but more complex

### number of variants in a protein library

x residues =  $20^x$  possible variants

12 residues =  $4 \times 10^{15}$  variants

lesson: impossible to cover **sequence space** except with short sequences (or few positions) and only the most high throughput techniques

Good luck with your papers!