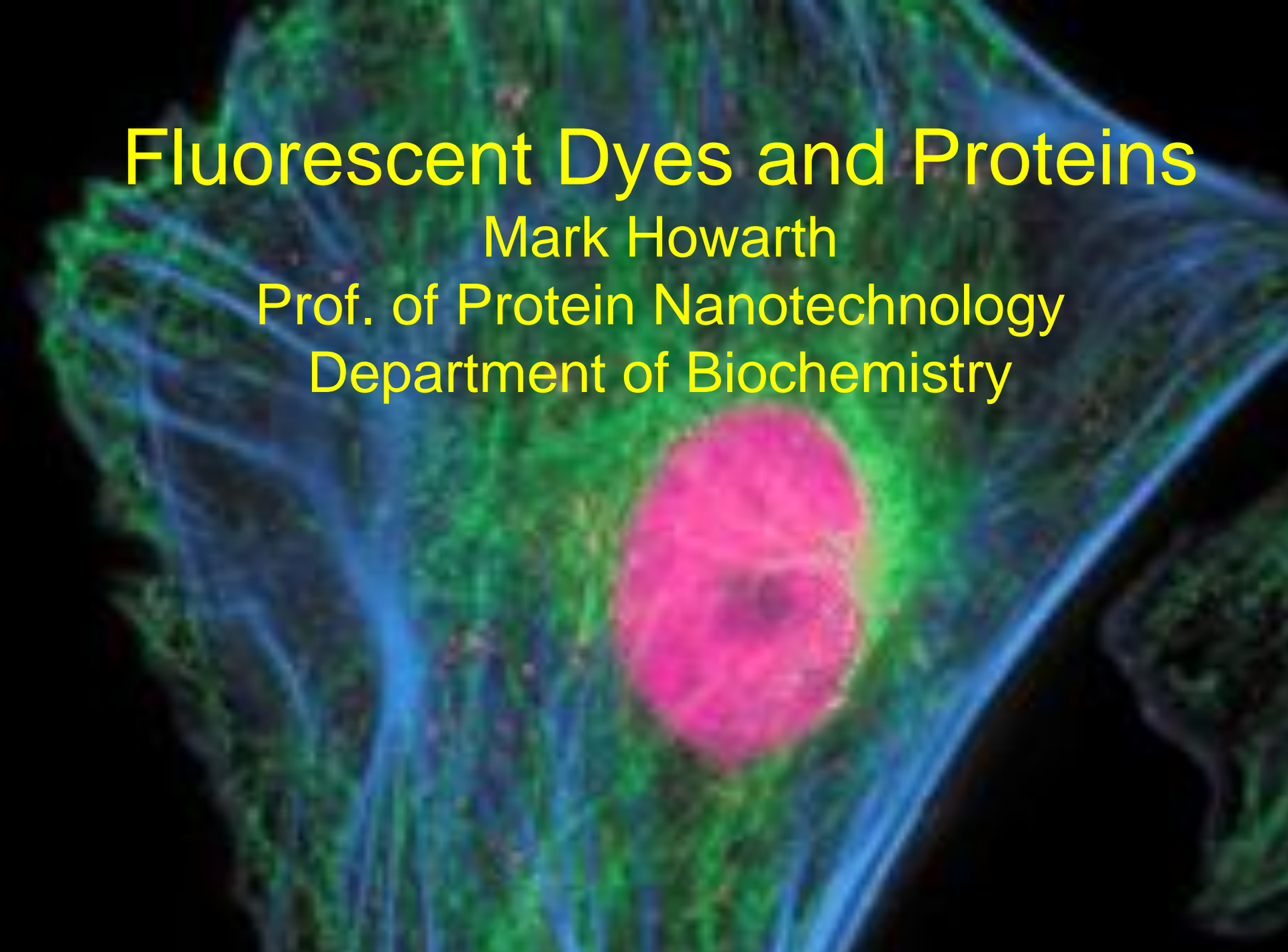


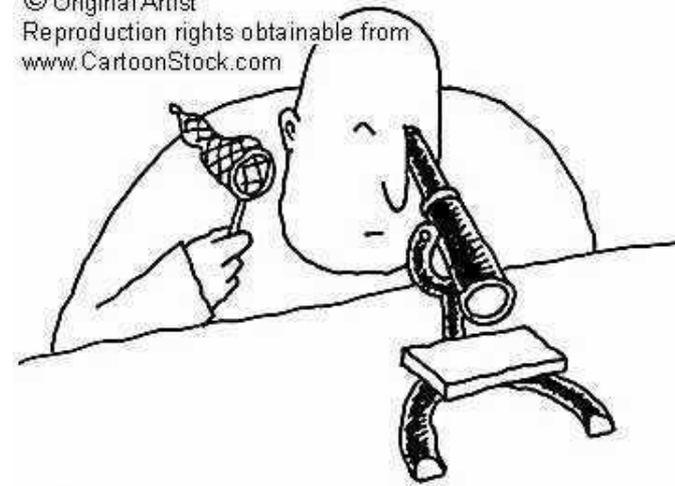
Fluorescent Dyes and Proteins

Mark Howarth

Prof. of Protein Nanotechnology

Department of Biochemistry

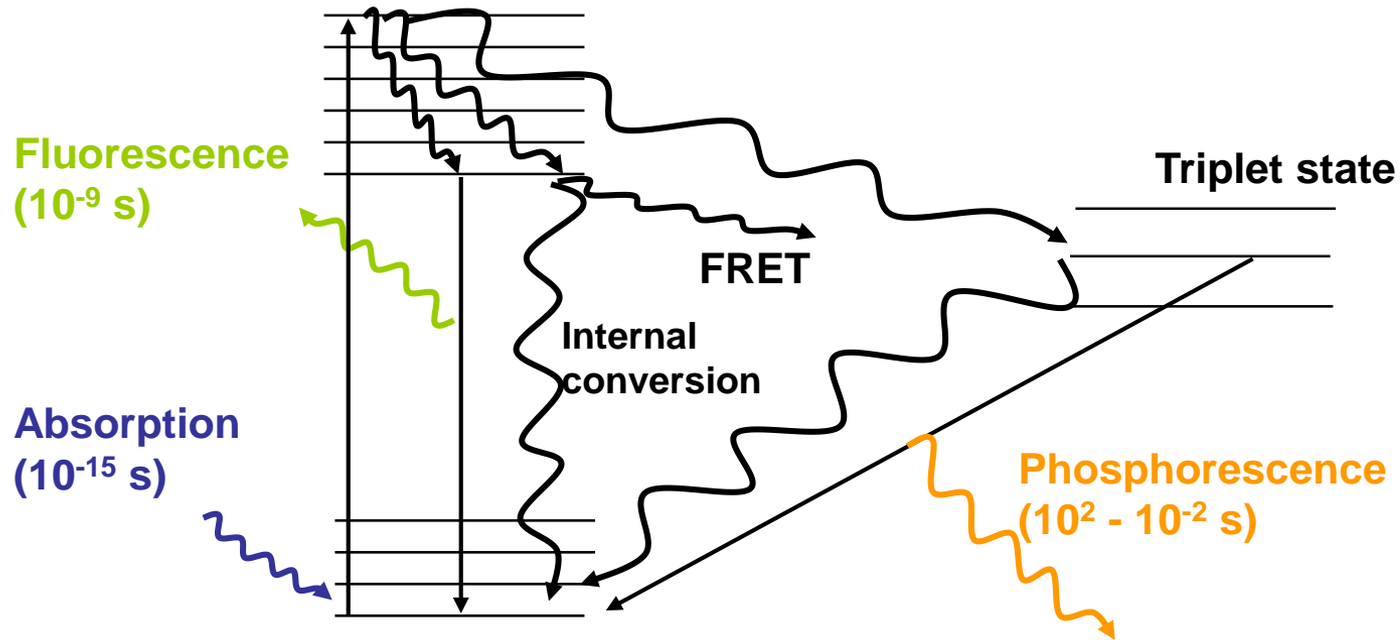




Overview

1. What kind of structures are fluorescent
2. How to make and target fluorescent probes
3. Fluorescent probes for cellular structure and function
4. Using light to control cells

Not all energy emitted as fluorescence



$$\text{Quantum yield} = \frac{\text{no. of fluorescent photons emitted}}{\text{no. of photons absorbed}}$$

e.g. EGFP QY=0.6 For every 10 photons absorbed, 6 are emitted.
(at optimal temp, pH etc.)

What sort of molecules are fluorescent?

Organic fluorophores

especially

1. Intrinsic fluorophores (source of autofluorescence)
2. Dyes
3. Fluorescent proteins

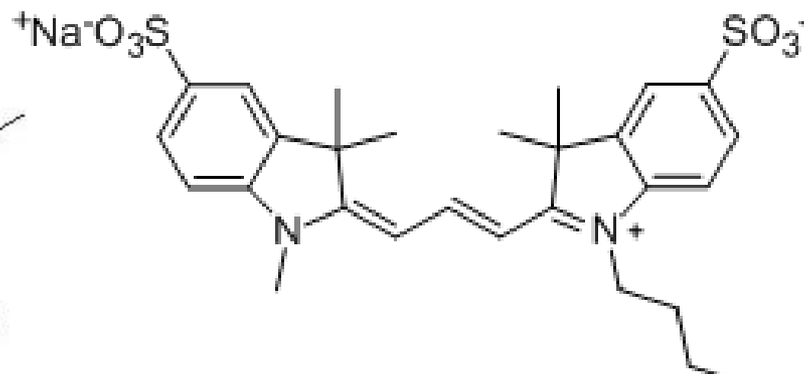
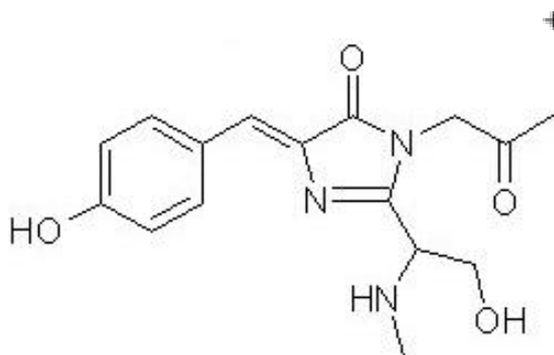
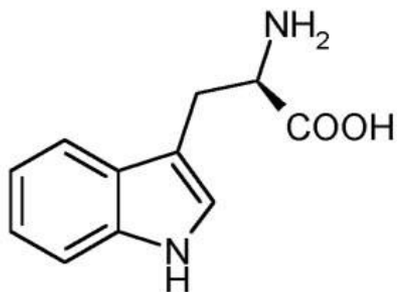
Inorganic fluorophores

especially

1. Lanthanides
2. Quantum dots

What sort of molecules are fluorescent?

1. Organic fluorophores



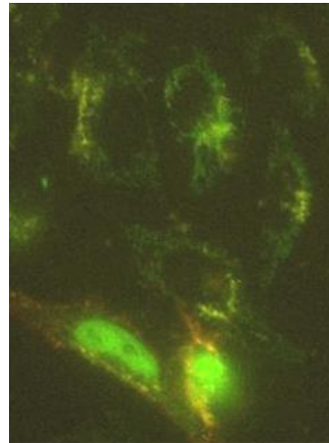
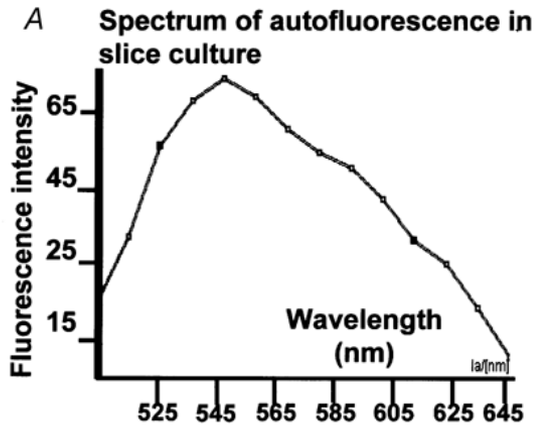
Chemical features:

1. Conjugation
2. Rigidity especially fused aromatic rings
3. Heteroatoms

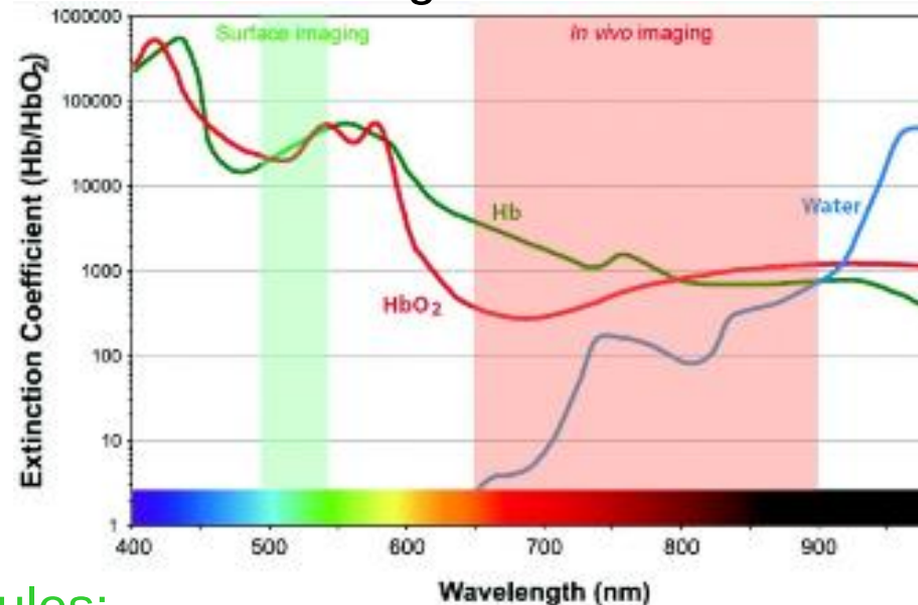
What sort of molecules are fluorescent?

1. Endogenous organic fluorophores

Isolated cells



Living animals



Most common autofluorescent molecules:

Flavins, NADH, NADPH, elastin, collagen, lipofuscin

Avoiding autofluorescence:

choose dye emitting in red with big Stokes shift

add quencher (Crystal violet)

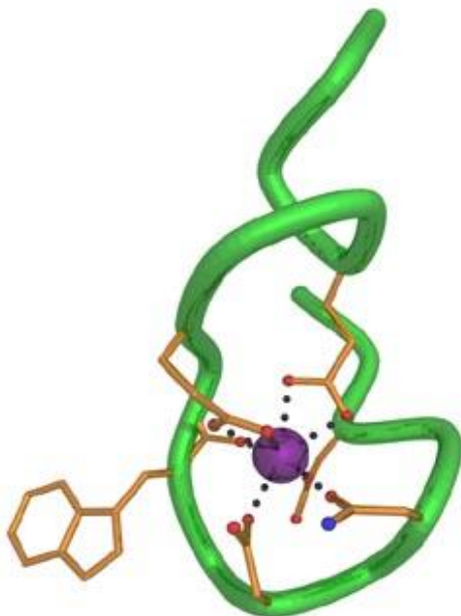
time-gate fluorescence

What sort of molecules are fluorescent?

2. Inorganic fluorophores

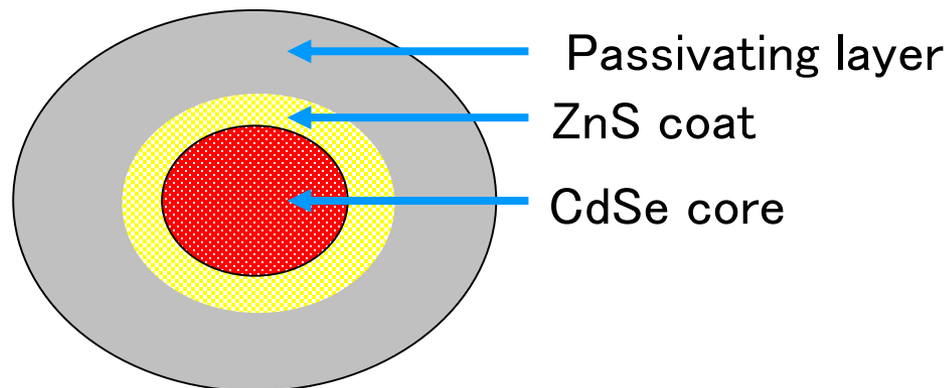
Lanthanides

Peptide sequence
binds Tb^{3+} and protects
from quenching by water



Curr Opin Chem Biol. 2010;14(2):247-54.
Lanthanide-tagged proteins--an illuminating
partnership. Allen KN, Imperiali B.

Quantum dots



+ bright, photostable, narrow emission
- large (~20 nm), expensive,
hard to target specifically

Michalet X, et al. Quantum dots for live cells, in
vivo imaging, and diagnostics. Science. 2005
307(5709):538-44.

How good is a fluorophore?

1. Excitation and emission appropriate

background worse in UV + with small Stokes shift
good match to filters on your microscope
look at other fluorophores at same time

2. Bright

see small numbers of fluorophores,
low self-quenching, high QY and absorbance

3. Stable to photobleaching

exciting light damages fluorophore

4. Non-toxic

5. Environment-insensitive (especially to pH)

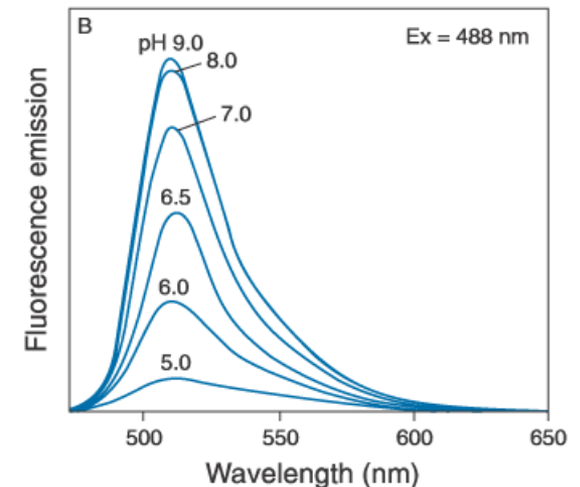
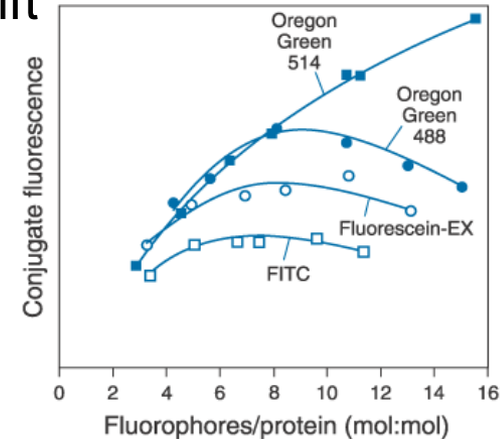
6. Little non-specific binding

7. Small

8. Little blinking

(9. Cost)

Green dye
self-quenching



Fluorescein pH sensitivity

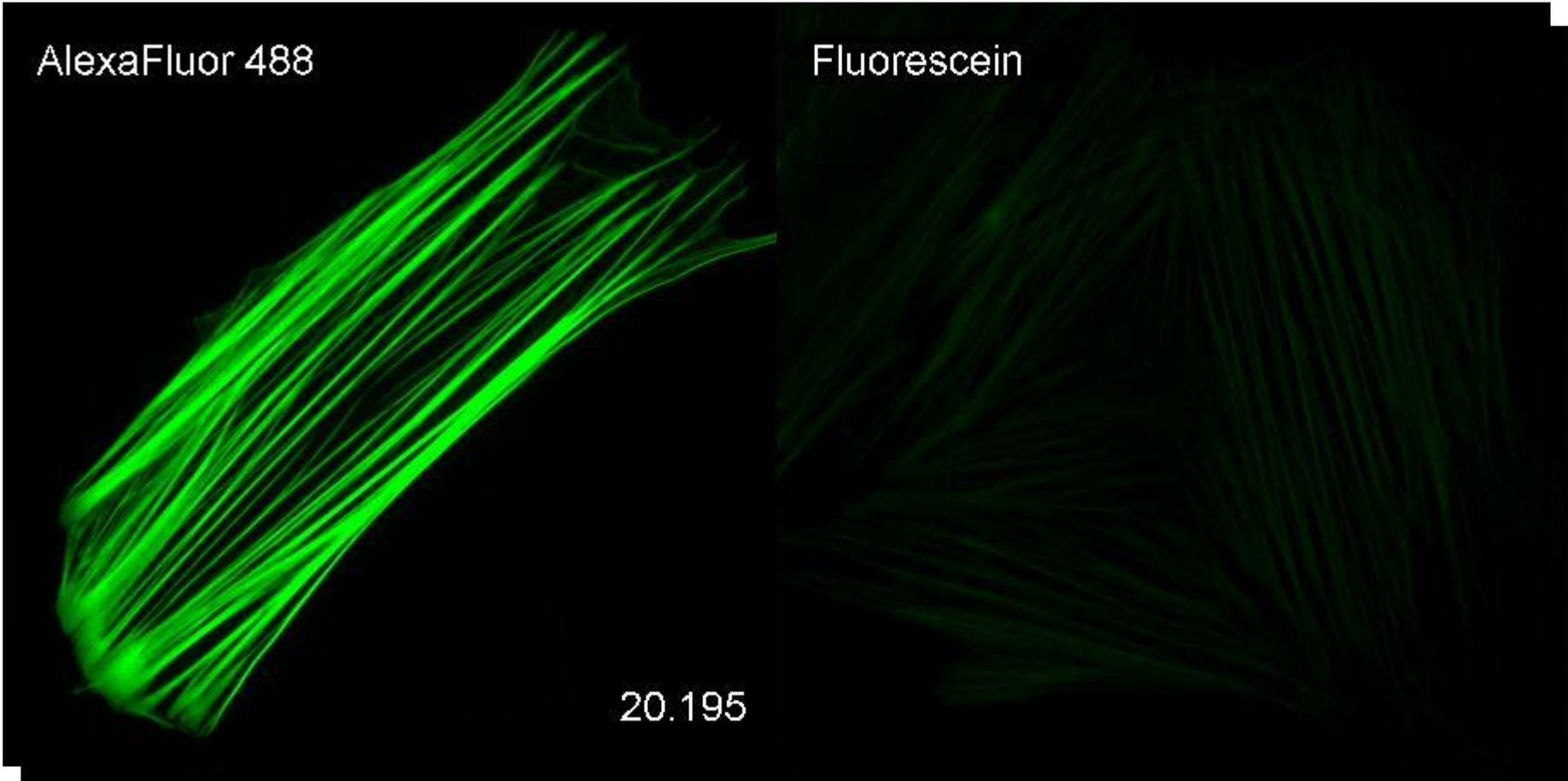
Alexa Fluor 488 vs Fluorescein Bleaching

2x Real Time

AlexaFluor 488

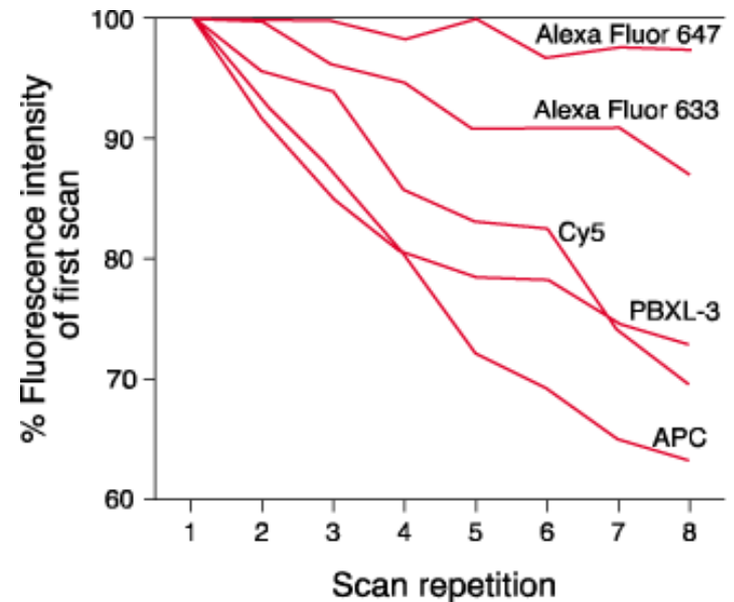
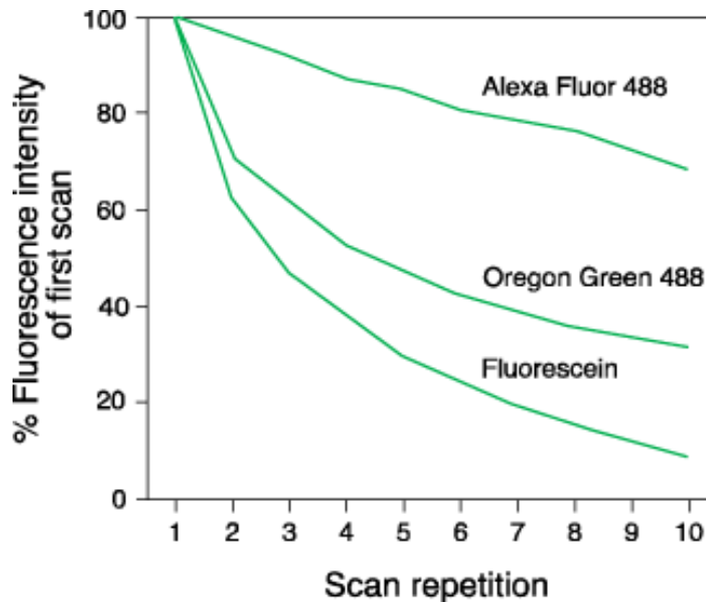
Fluorescein

20.195



Alexa Fluor Dyes – Photostability

Laser-scanning
cytometry
EL4 cells
biotin-anti-CD44
+ streptavidin
conjugates



Fluorescein is the commonest dye
but has poor photostability.

Also consider Atto dyes (Sigma) and Dyomics dyes

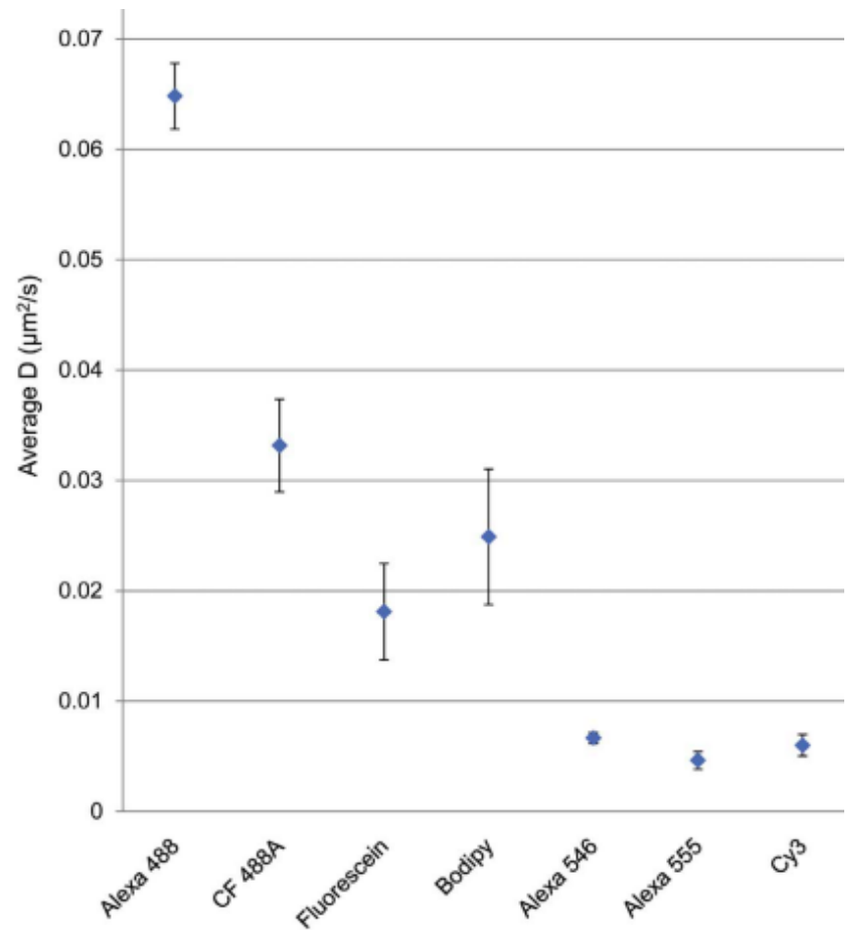
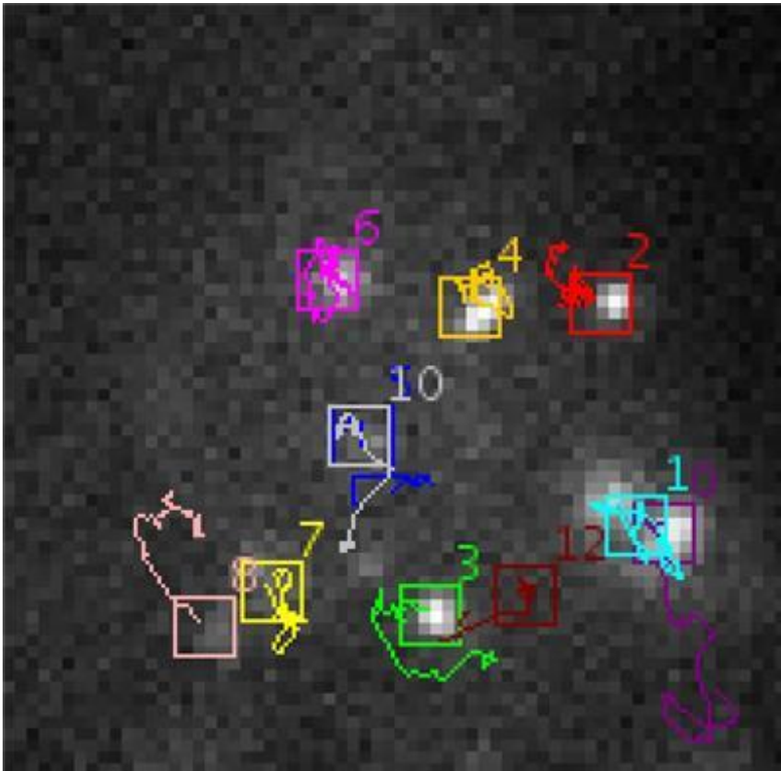
Dye affects non-specific binding and receptor mobility

Dye makes a big difference to non-specific binding

Diffusion coefficient for dye-labelled Affibody against EGFR
varies 10-fold with hydrophilic versus more hydrophobic dye!

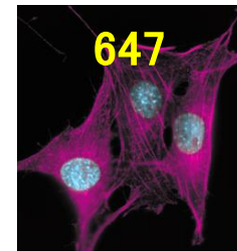
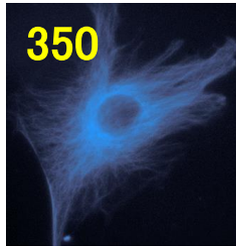
L. Zanetti-Domingues et al.

PLoS one 2013



Multiplexing- four main colours

Excitation wavelengths:



Emission wavelengths:

Blue

green

orange/red

far red

350

400

450

500

550

600

650

700

DAPI/UV

FITC

TRITC

FAR RED

Alexa Fluor® 350
Coumarin, AMCA

Alexa Fluor® 488
Fluorescein (FITC)
Cy2

Alexa Fluor® 555
Rhodamine,
TAMRA, TRITC
Cy3

Alexa Fluor® 647
Cy5, APC

Alexa Fluor® 594
Texas Red, Cy3.5

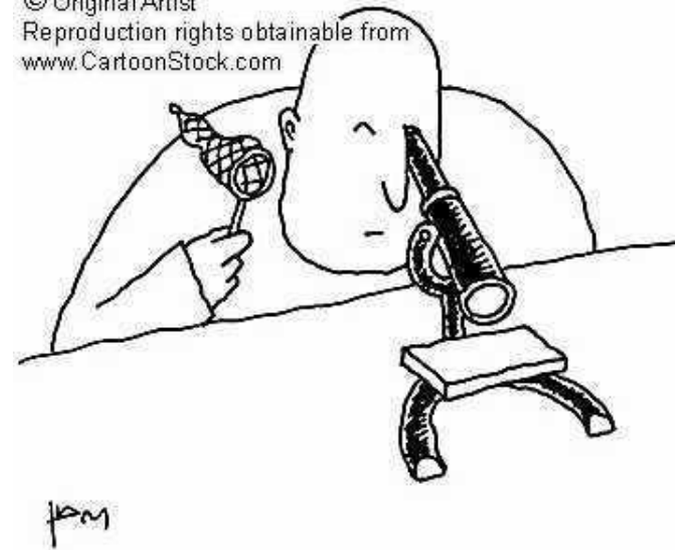
Colour Selection



Brightness



Photostability



Overview

1. What kind of structures are fluorescent
2. How to make and target fluorescent probes
3. Fluorescent probes for cellular structure and function
4. Using light to control cells

Antibodies for cellular imaging

Live cells

Label plasma membrane and secretory pathway

Penetrate plasma membrane

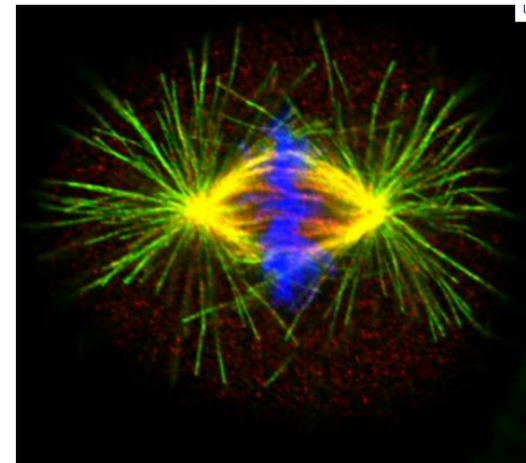
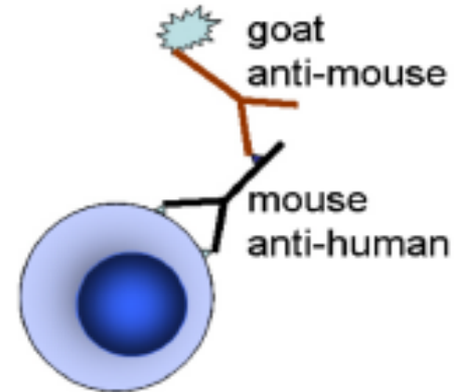
(microinjection, electroporation, pinosome lysis, streptolysin, cell permeable peptides, ester cage)

Get dynamics, avoid fixation artifacts

Fixed cells

Permeabilise

Still can give enormous amount of useful information



Not just antibodies for targeting

Other types of targeting agents:

Proteins

(especially antibodies, but also transferrin, insulin, EGF etc.)

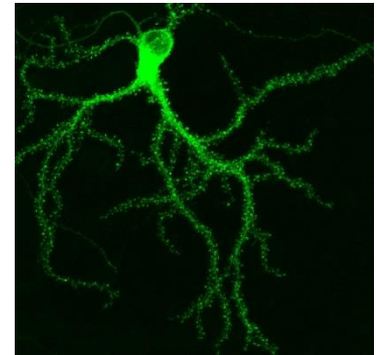
Peptides (MHC class I pathway, proteasome function)

RNA (mRNA, molecular beacons, aptamers, siRNA)

DNA

lipids, lipoproteins

drugs



How to dye: it is easy

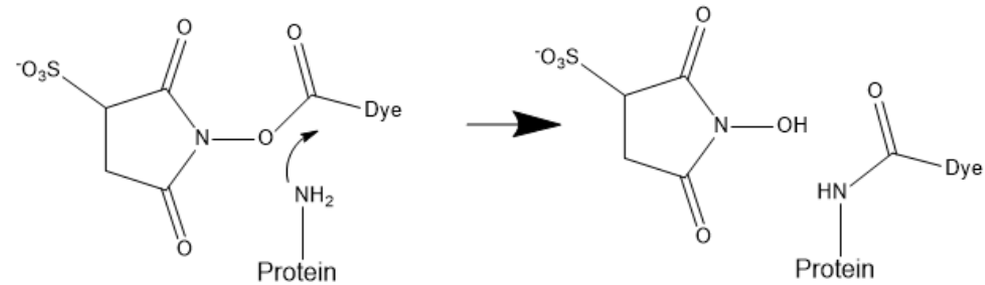
Multiple ways to modify proteins
(see Molecular Probes catalogue)

Most common ways are to modify:

1. Lysine

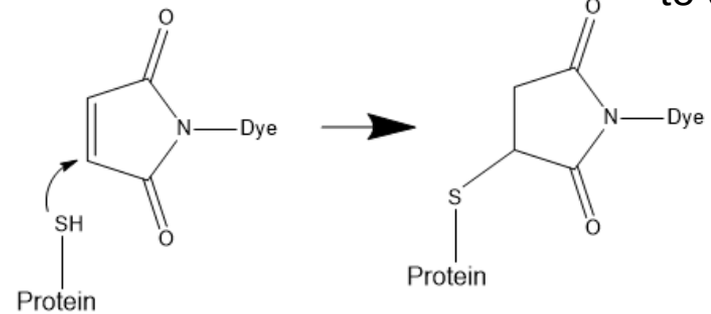
or

2. Cysteine



sulfoNHS-dye

Amide bond
to dye



maleimide-dye

Thioether bond
to dye

- A Add dye to protein for 3 hr
- B 1cm Sephadex column to remove most free dye (10 min)
- C Dialyse away rest of free dye (24 hr)

Site-specific protein labelling methods

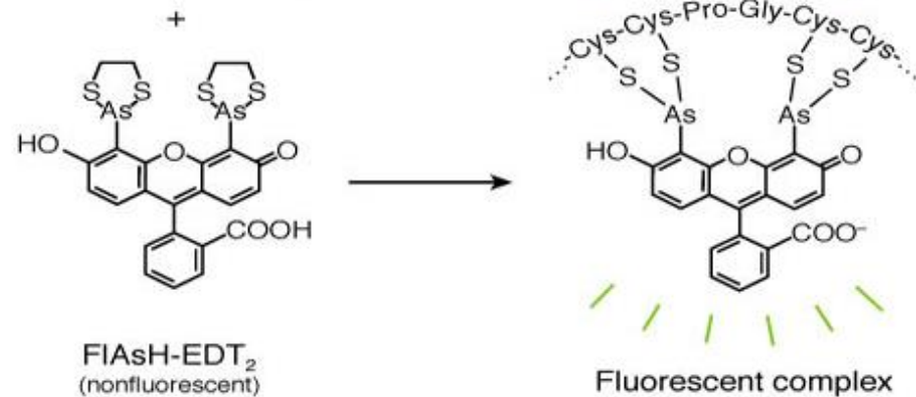
1. Binding domain

SNAP-tag (19 kDa NEB), HaloTag (34 kDa, Promega)



2. Binding peptide

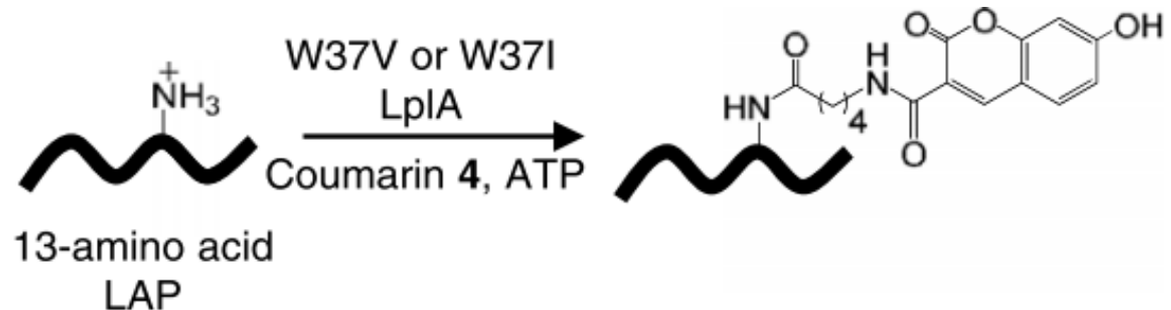
FIAsh (Invitrogen)

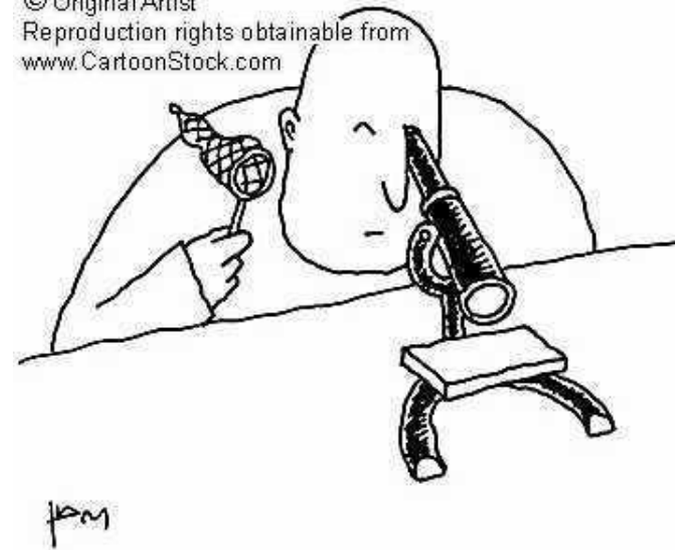


3. Enzymatic ligation to peptide

PRIME

AY Ting PNAS 2010





Overview

1. What kind of structures are fluorescent
2. How to make and target fluorescent probes
3. Fluorescent probes for cellular structure and function
4. Using light to control cells

Putting the signal in context: nuclear labelling

(follow DNA even when nucleus breaks down)

Fixed cells:

Intercalate into DNA

DAPI

(well away from other channels)

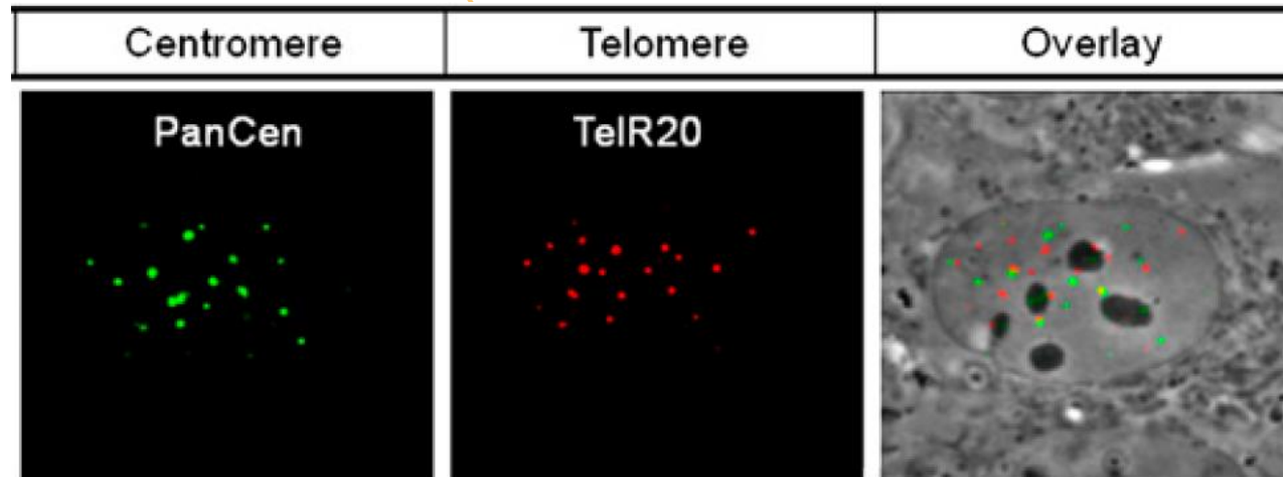
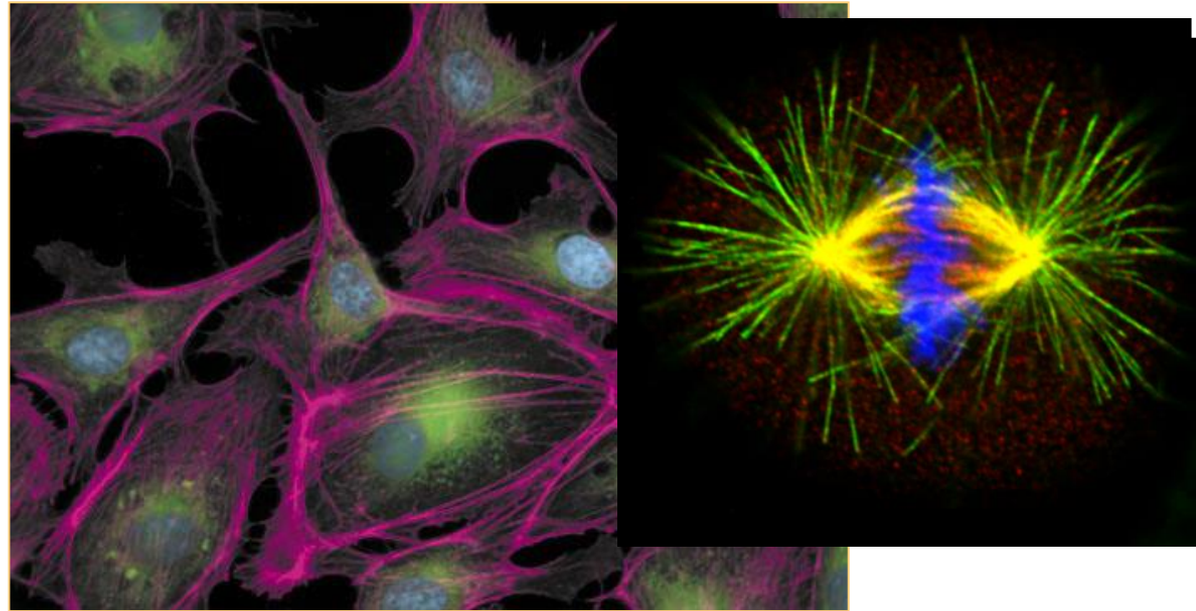
Hoechst 33342

Live cells:

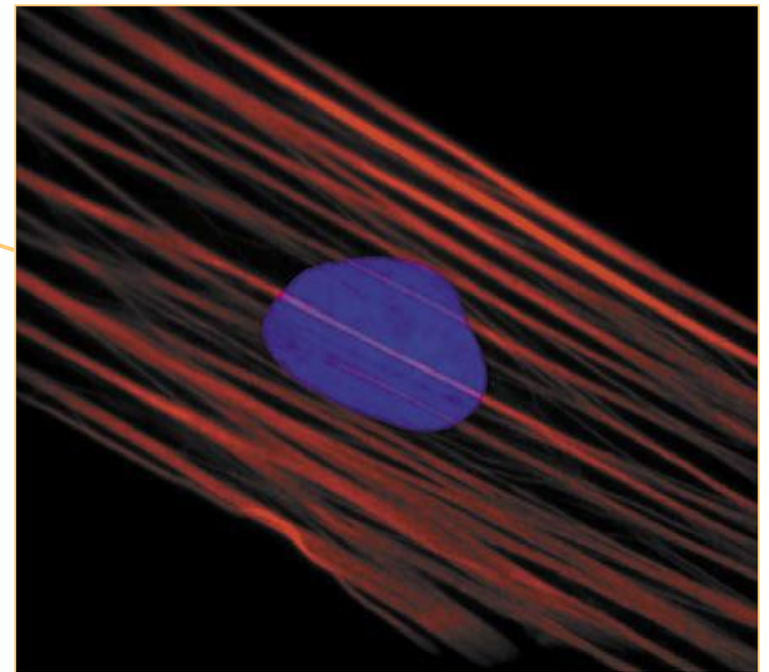
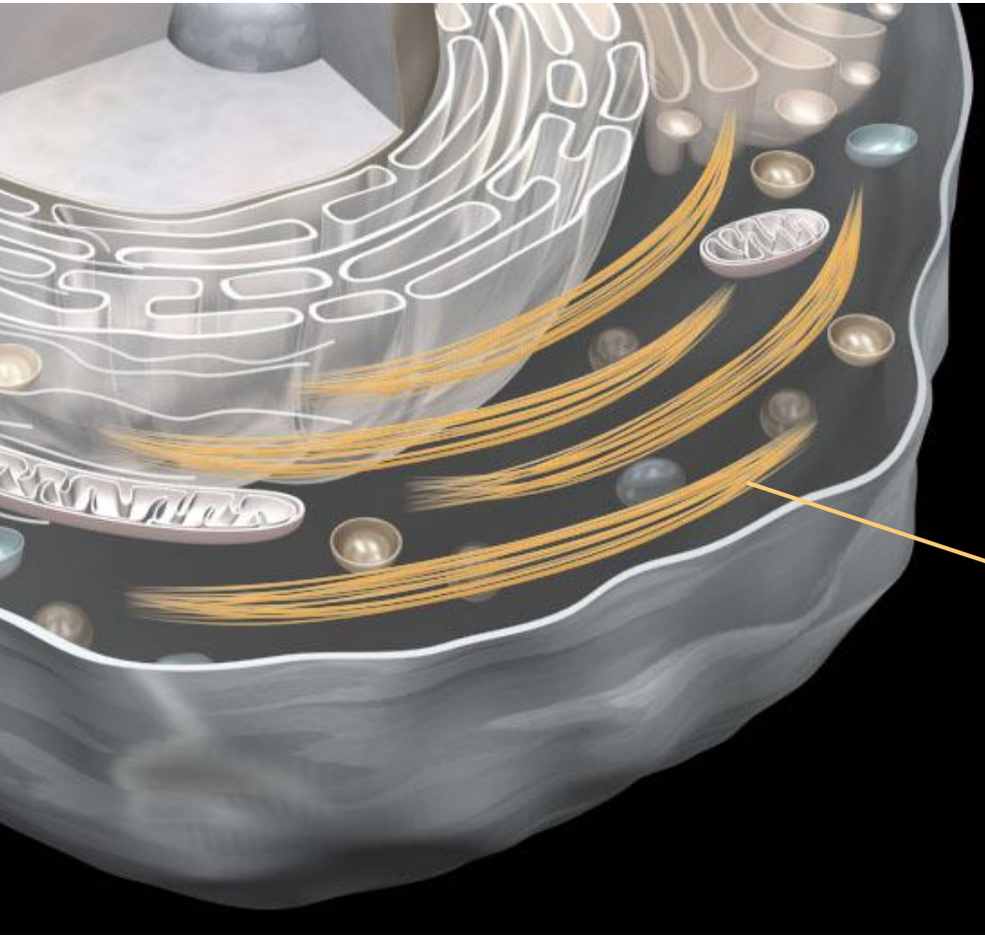
histone H2B-GFP

TALEN-XFP

CRISPR/Cas9-GFP (B. Chen et al. Cell 2013 155:1479)



Putting the signal in context: actin labelling



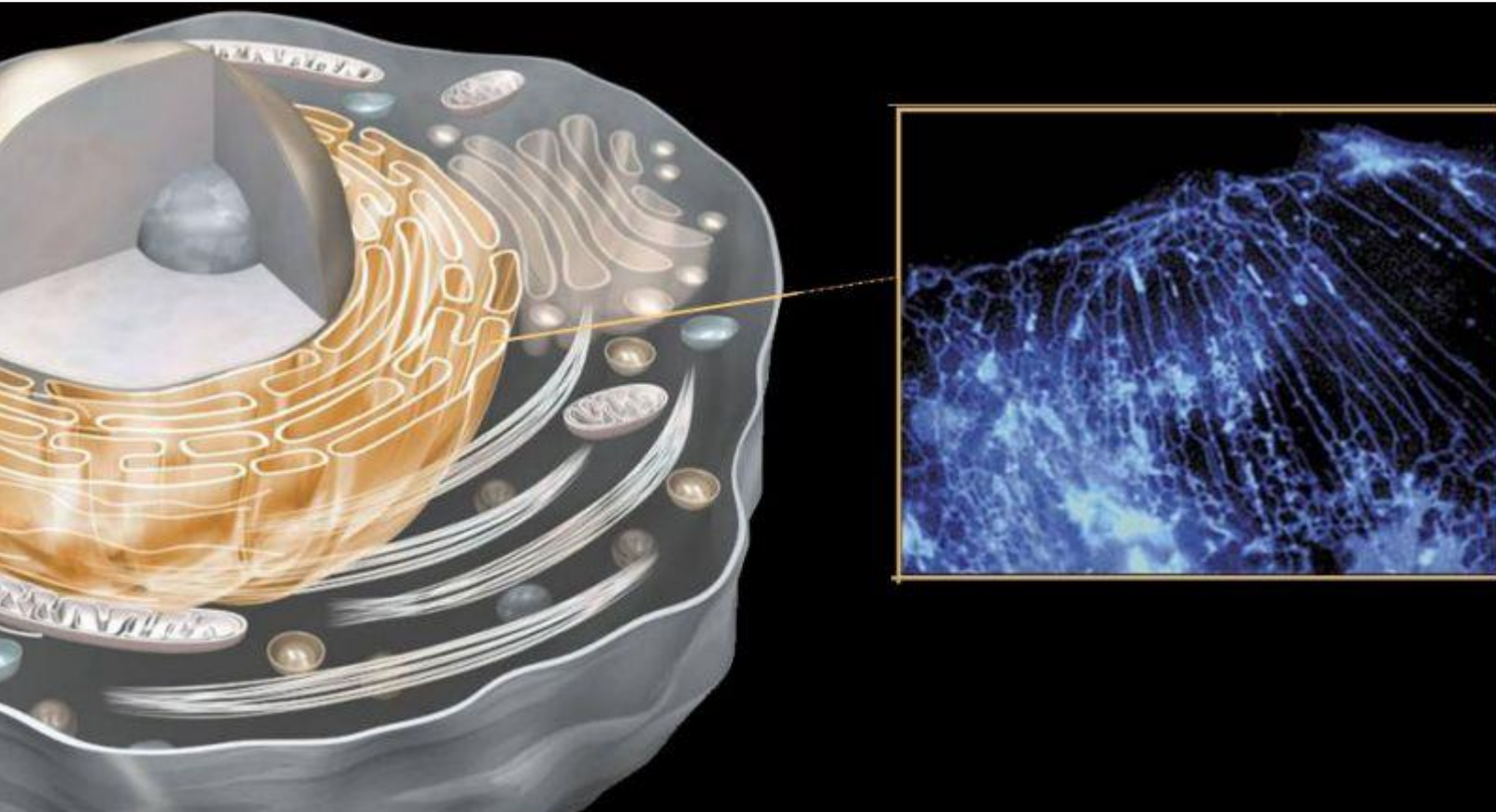
Fixed cells: phalloidin-dye

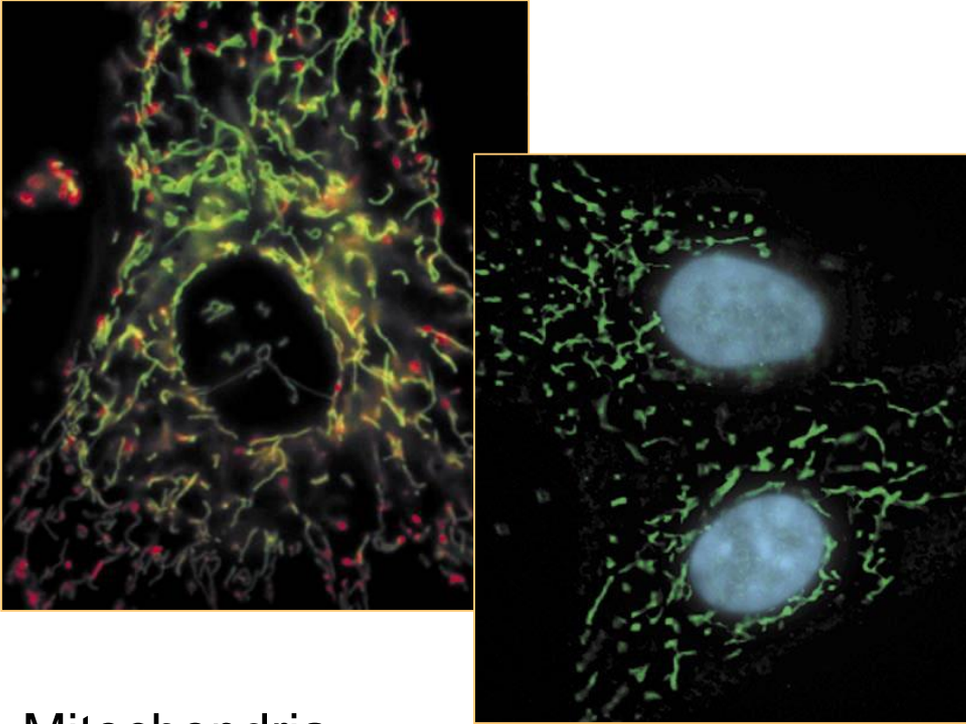
Endoplasmic Reticulum

ER-Tracker™ Blue-White DPX

antibody to calnexin

Live cells: ss-GFP-KDEL





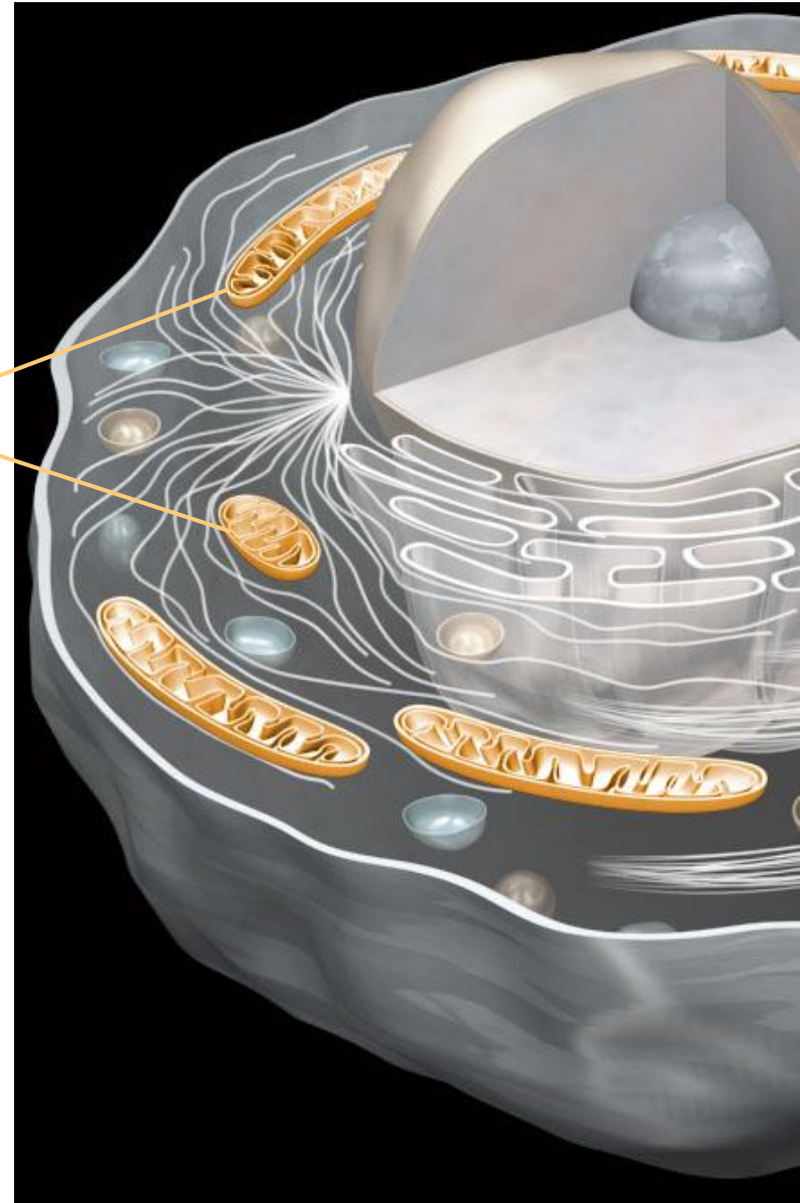
Mitochondria

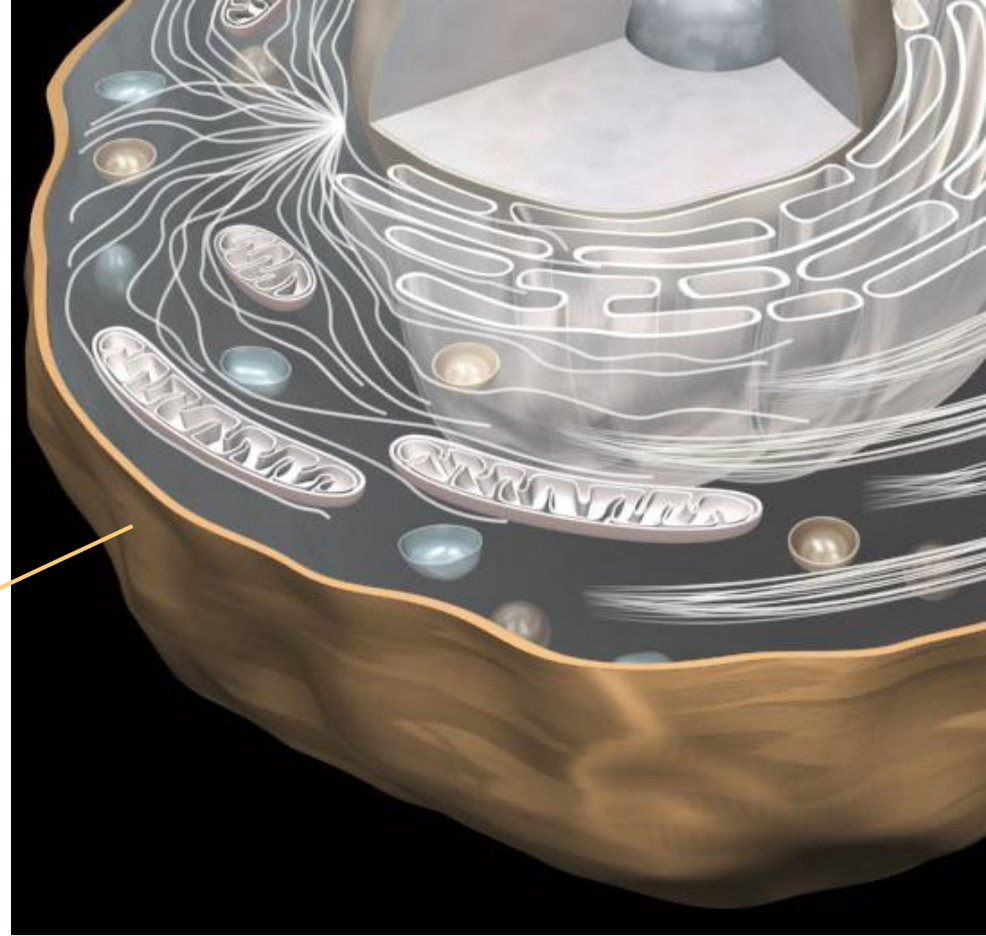
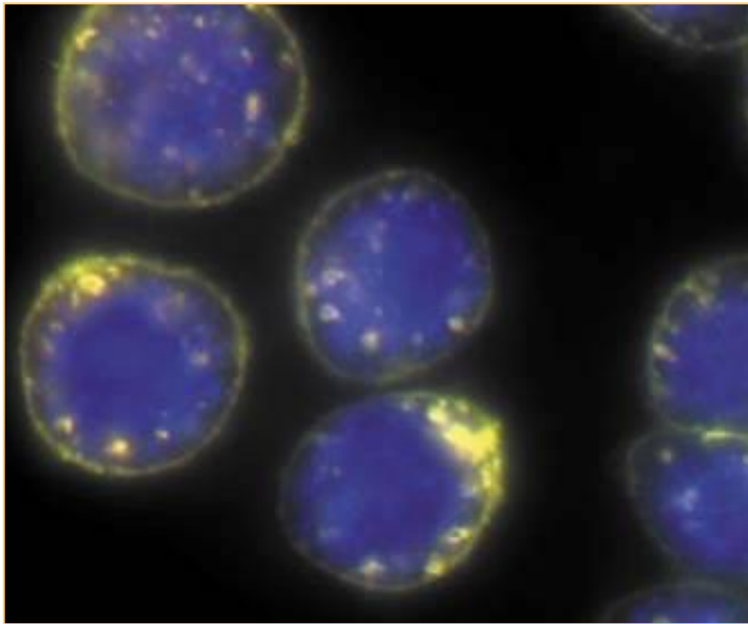
Fixed cells: anti-cytochrome oxidase subunit I Ab

Live cells: MitoTracker® Red/Green/Orange

JC-1 (red J-aggregates at high conc., red to green depends on membrane potential)

Mitochondrial targeting sequence-GFP

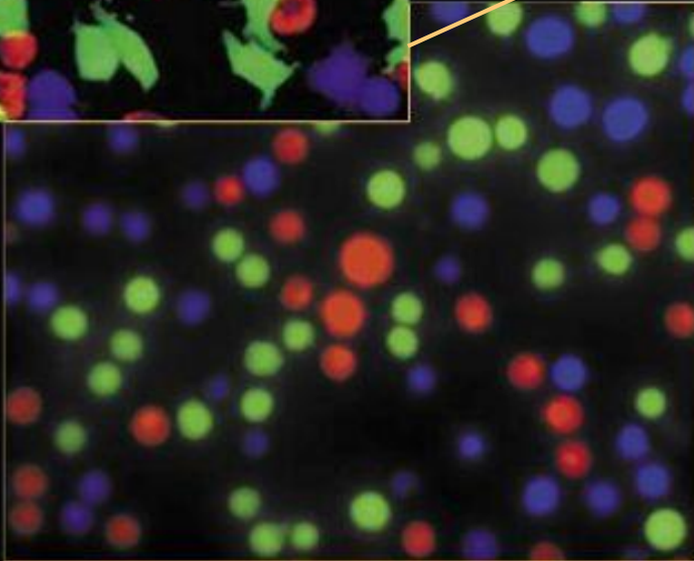
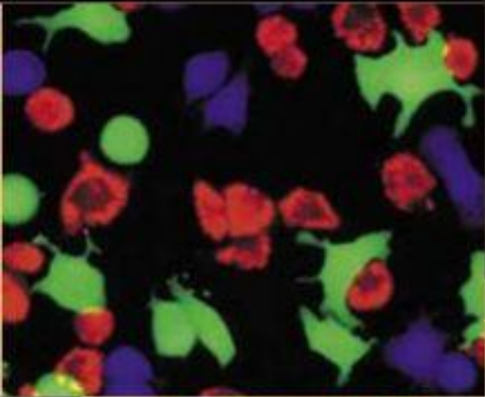
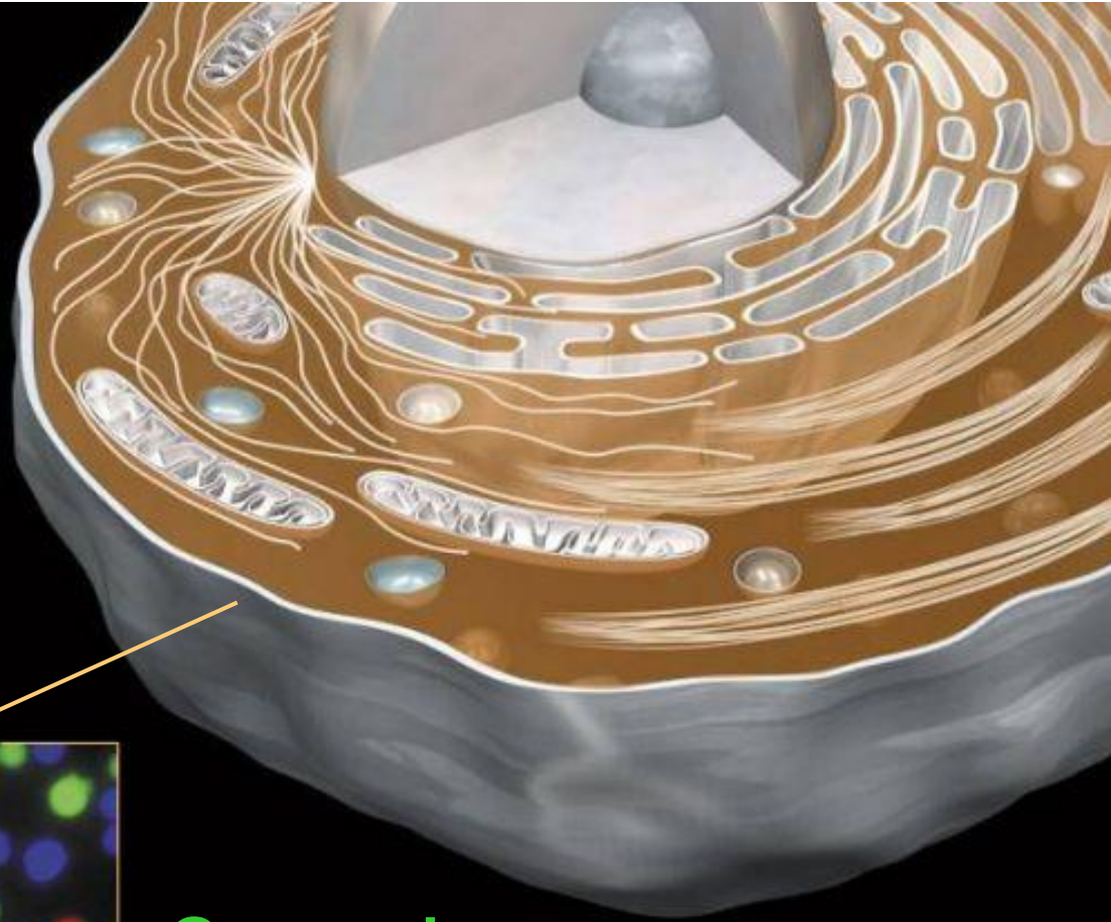




Lipid Rafts

BODIPY® FL C₅-ganglioside GM1

Fluorescent Cholera Toxin subunit B (CT-B)



Cytosol

Live cells:

CellTracker™ Green CMFDA

Calcein, AM

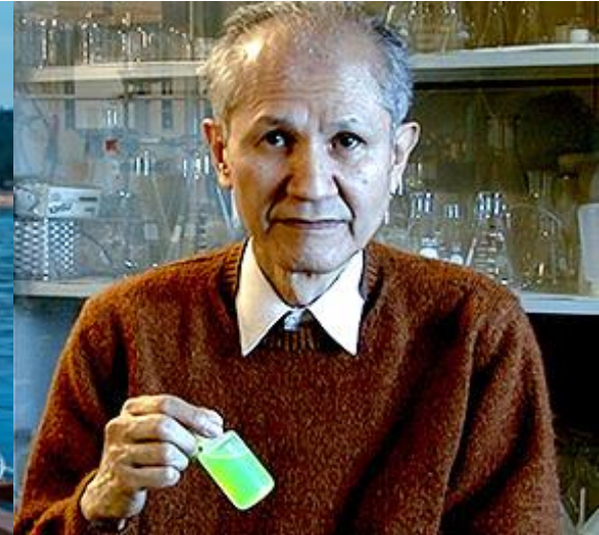
Qtracker

GFP with nuclear export sequence

The breakthrough of fluorescent proteins from jellyfish

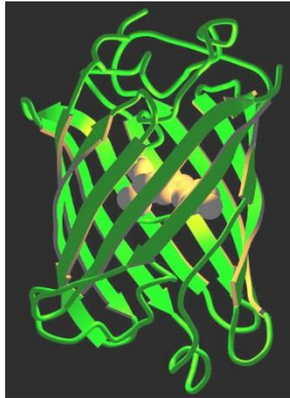


*Aequorea
victoria*

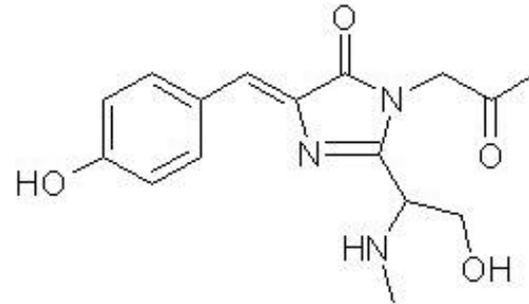


Osamu
Shimomura

The breakthrough of fluorescent proteins for live cell imaging



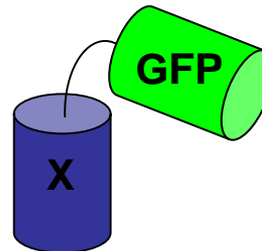
GFP fold
 β -can



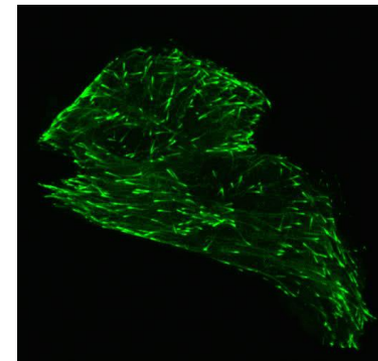
GFP chromophore
from Ser-Tyr-Gly



Link GFP sequence to gene of your favourite protein



GFP folds and becomes fluorescent



GFP lights up your favourite protein in cell

Fluorescent proteins are more than just labels

Photoactivation/Photoswitching

PA-GFP, Dronpa, Eos

Reporting on environment

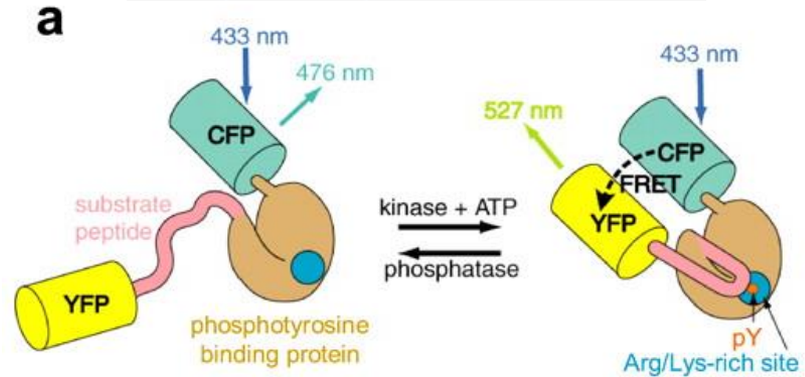
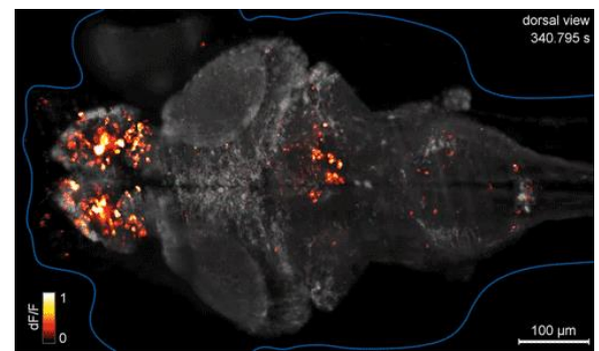
Ca²⁺, phosphorylation, cAMP, cGMP, pH, neurotransmitters, voltage, cell cycle, redox

Reporting on protein-protein interaction

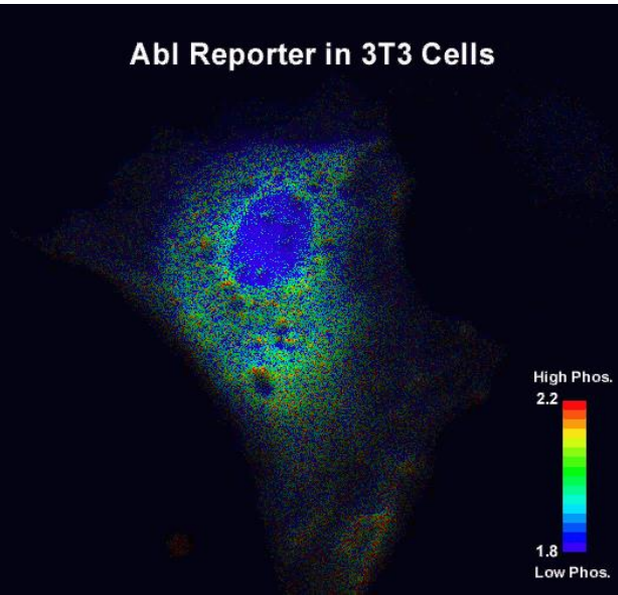
CFP/YFP FRET, split fluorescent proteins

Modifying environment

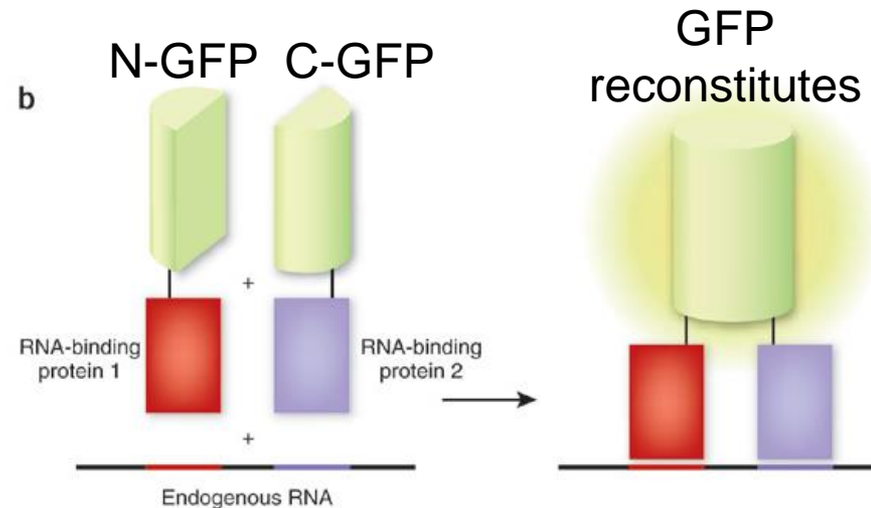
Singlet oxygen generation, Channelrhodopsin



Abl Reporter in 3T3 Cells



**Targeting advantage
to defined compartment,
cell-type,
developmental stage**

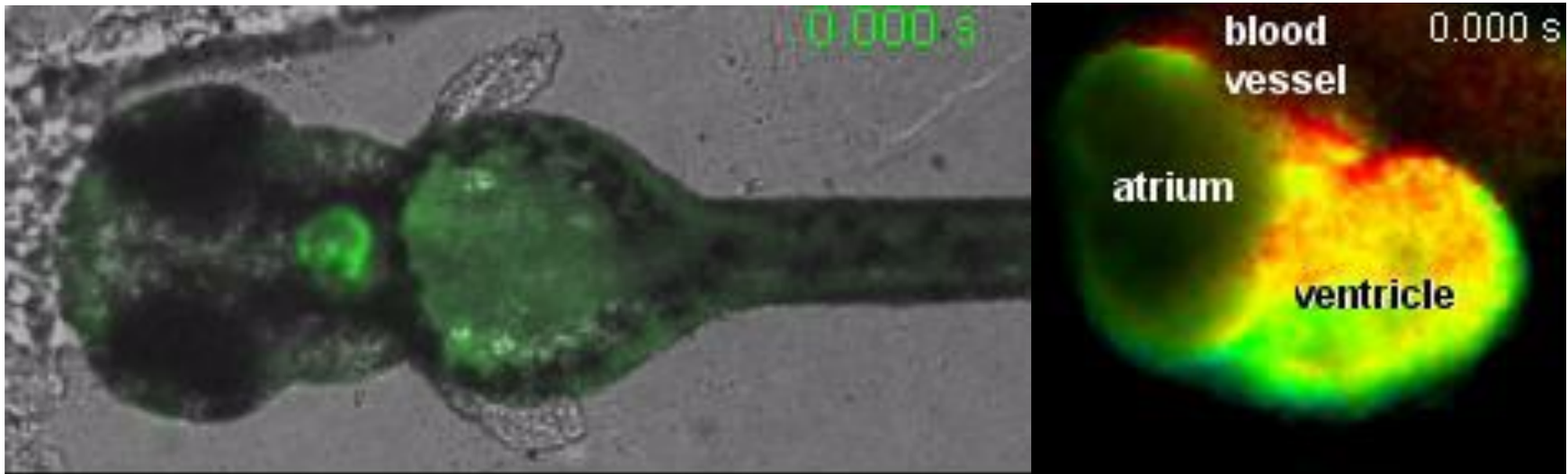


Sensing voltage with fluorescent protein

Mermaid FRET voltage-sensor
by FP fusion to voltage-sensing phosphatase

Expressed in zebrafish heart
Non-invasive testing of mutant phenotypes
and drug cardiotoxicity.

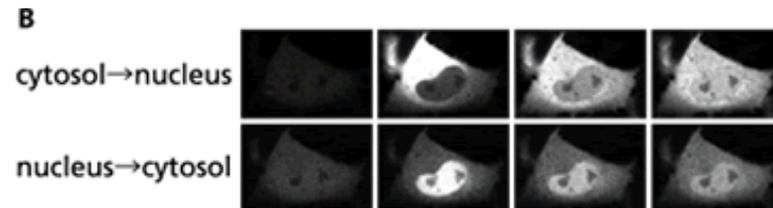
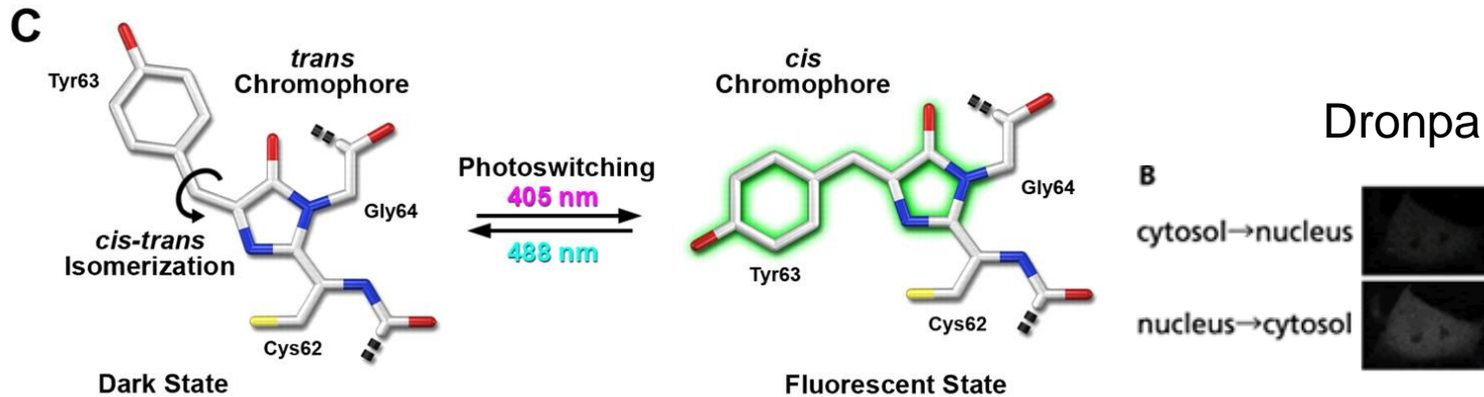
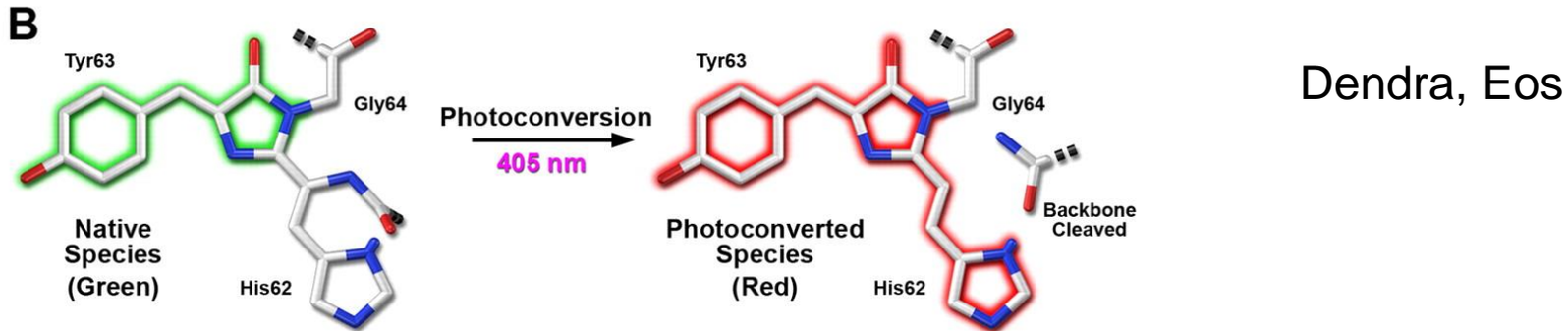
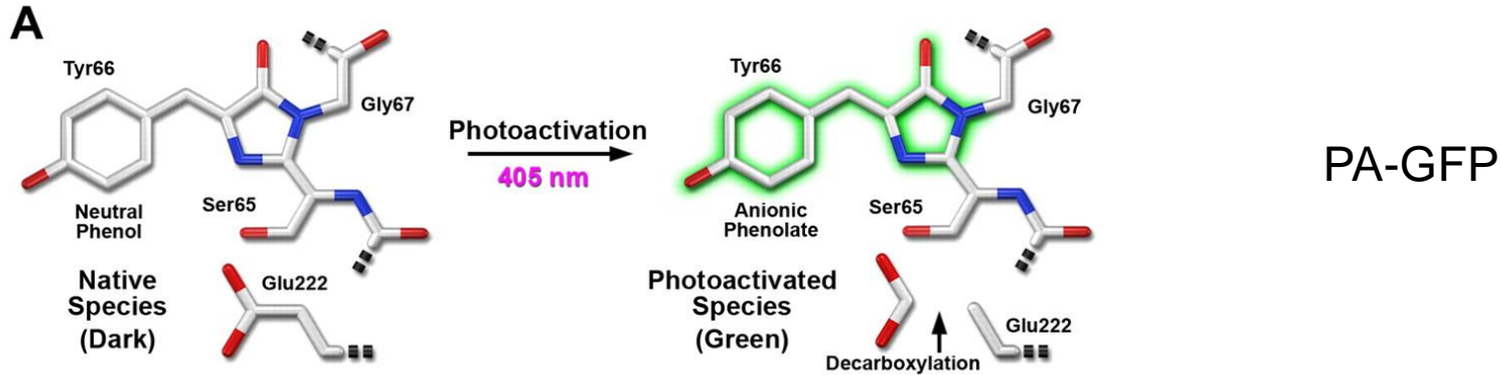
Tsutsui, Miyawaki J Physiol 2010



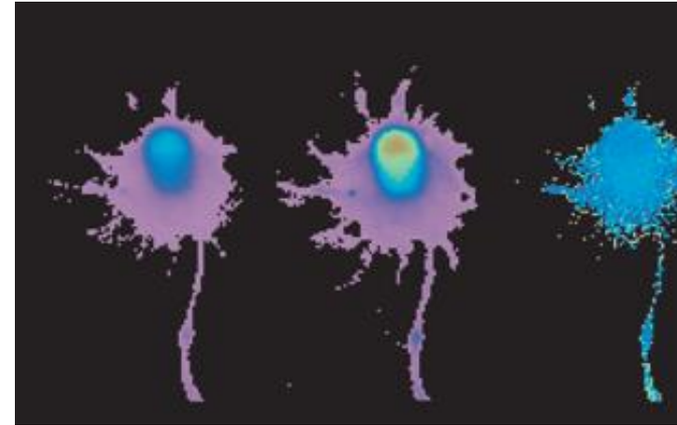
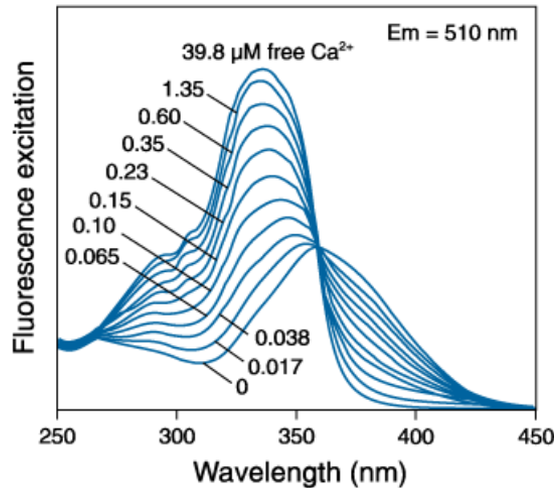
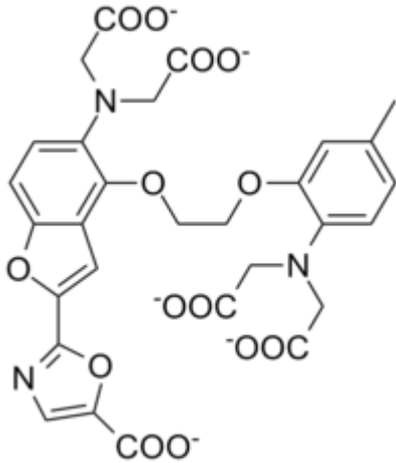
FRET sensor ratio crucial

best is YC2.60 cameleon: 600%,
if <20% then lost in cellular noise

Chromophores in switching



Small molecule fluorescent sensors



Fura-2 sensing calcium

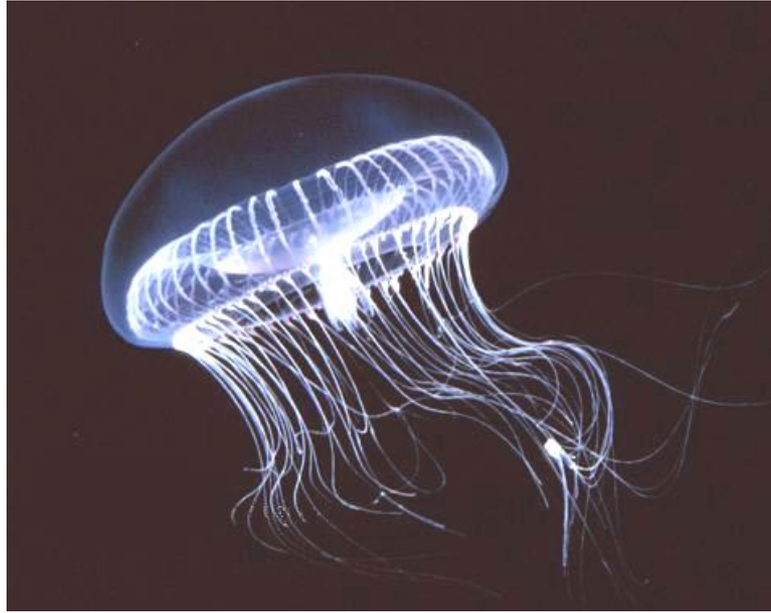
Metal ions: calcium, magnesium, zinc, sodium, potassium, chloride, mercury

pH (also dyes to conjugate to proteins, CyPher from GE, SNARF from Invitrogen)

Reactive oxygen species, nitric oxide

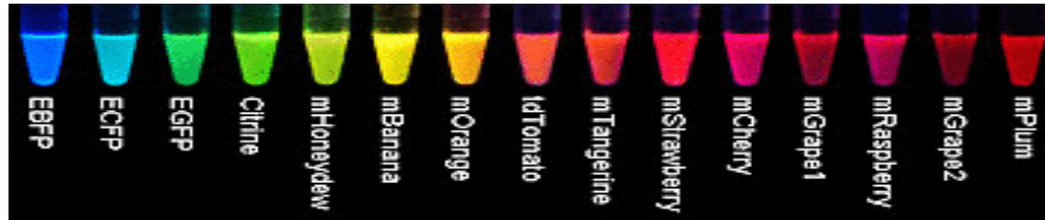
Transmembrane potential

How good is a fluorescent protein?



A. victoria GFP is good for jellyfish,
but not great for cell biologists!

How good is a fluorescent protein?



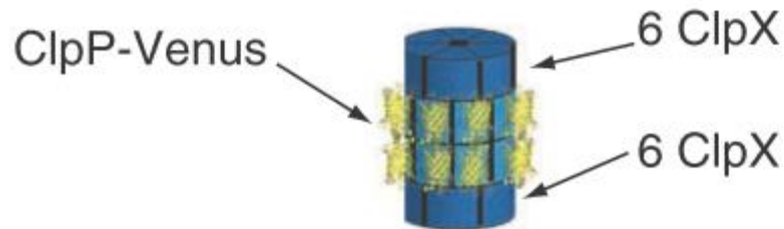
A. *victoria* GFP is terrible!

EGFP is OK, but there are now better...

1. Excitation and emission λ good match to filters on your microscope
look at other fluorophores at same time
2. Bright $\epsilon \times QY$ Clover, YPet 2.5 x EGFP
mRuby2 3x mCherry
3. Stable to photobleaching EBFP bad, mCherry and YPet good
4. Non-toxic attach on right part of your protein
all make H_2O_2 , FPs can transfer electrons
5. Environment-insensitive especially to pH, chloride
CyPet does not fold at $37^\circ C$, all need O_2
Photoactivatable FP did not work in ER
6. Little non-specific binding fully monomeric, A206K non-dimerising
7. Fast Maturation Venus 2 min. Red FPs can start off green
half-time ~ 15 min mCherry, 100 min TagRFP

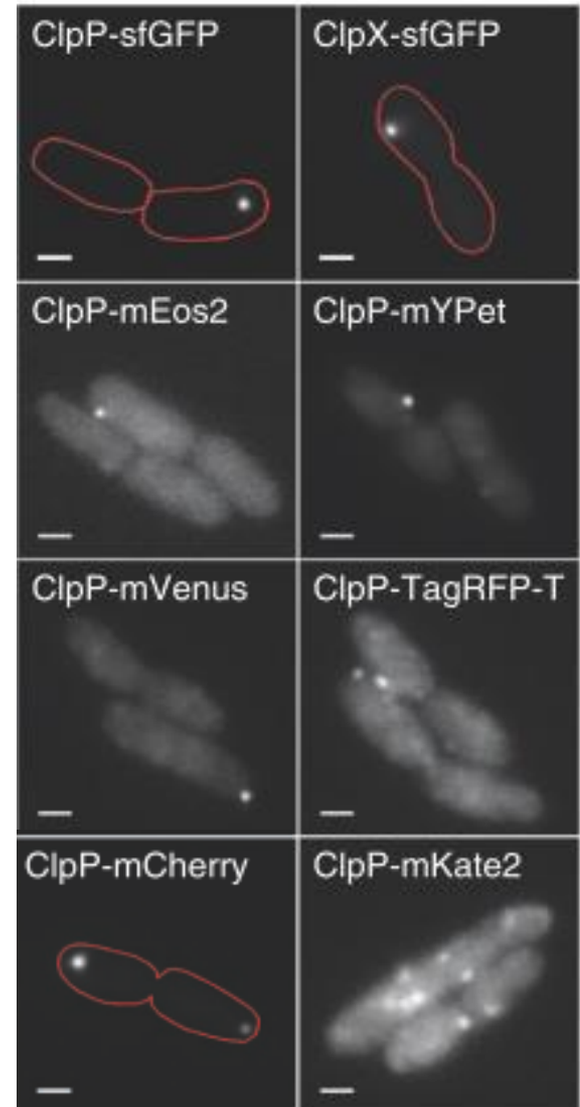
You MUST worry about FP multimerization!

Tag multimerizing protein with FP and
sometimes see foci-
are these real or caused by the tag?



With hexameric barrel involved in
E. coli protein degradation,
many commonly used FPs induce
artificial foci

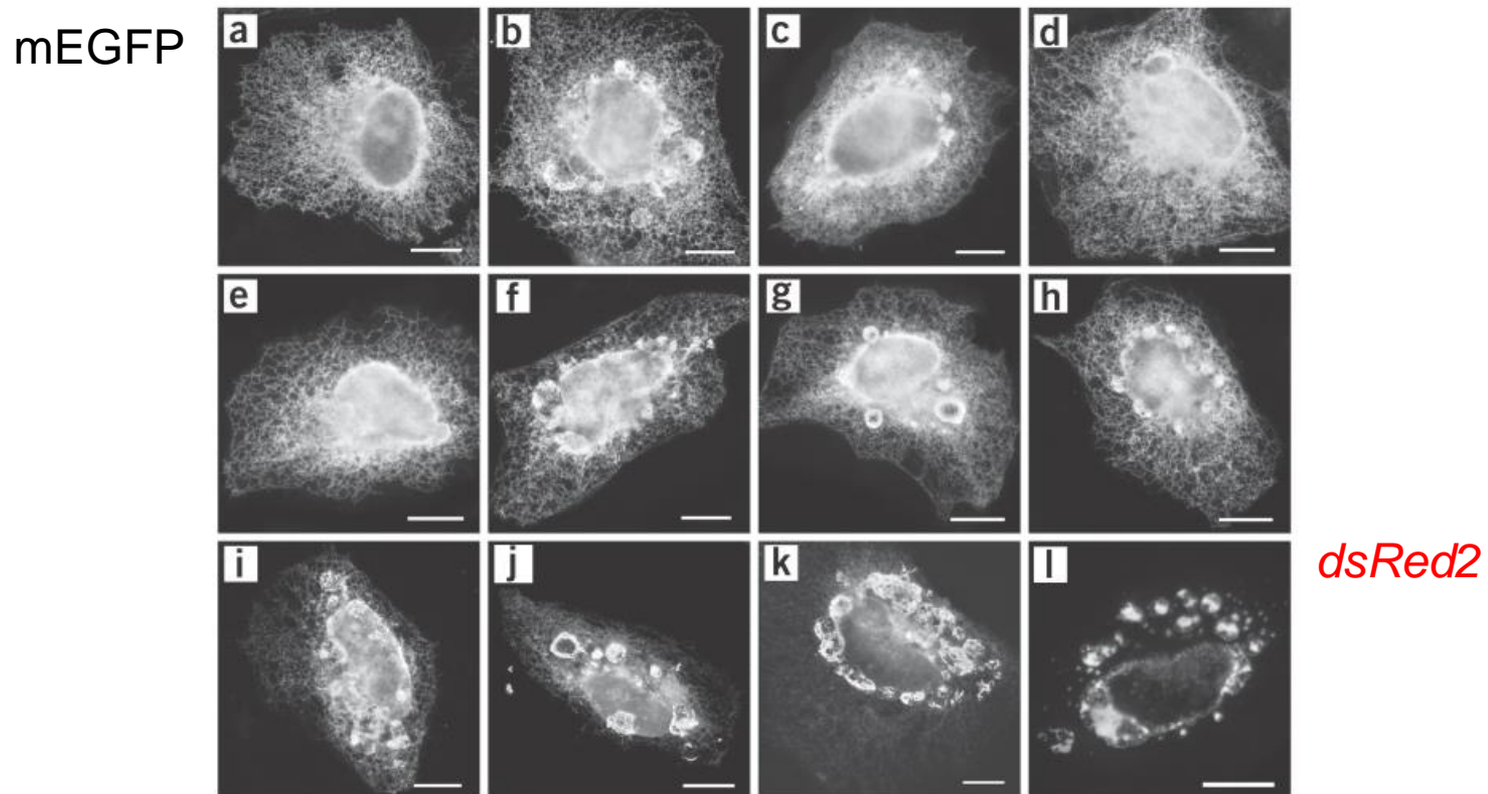
(no cluster with Ab or SNAP-Tag)
as well as affecting daughter cell
inheritance of proteolysis ability
mCherry, sfGFP, mYPet poor!
mGFPmut3, Dronpa OK
D. Landgraf et al. Nature Meth 2012



You MUST worry about FP multimerization!

In mammalian cells,
linked to ER membrane protein,
some FPs cause whorls

PJ Cranfill et al. Nature Meth 2016



Problems with GFP in cells

- **GFP with light can donate electrons to different acceptors**
(FMN, FAD, NAD⁺, cyt. c)
GFP reddens after transfer:
photobleaching and phototoxicity
use DMEM lacking e⁻ acceptors
(riboflavin or all vitamins) for less bleaching
Lukyanov Nat Meth 2009
HAM F-12 medium 6x better EGFP stability in cells than DMEM, RPMI!
Lukyanov Biotechniques 2015
- **EGFP not good in secretory pathway**
mixed disulfide oligomers in ER and non-fluorescent in *E. coli* periplasm
(superfolder GFP behaves fine)
Erik Snapp, Traffic 2011

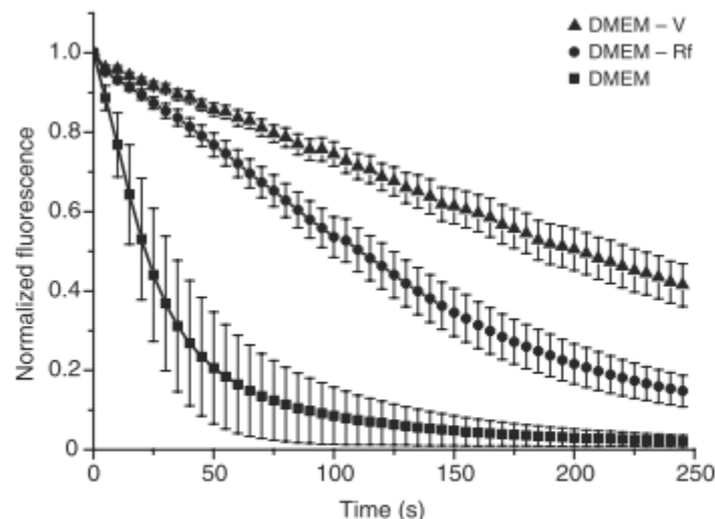
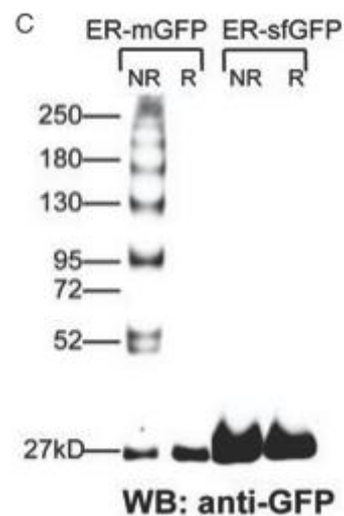


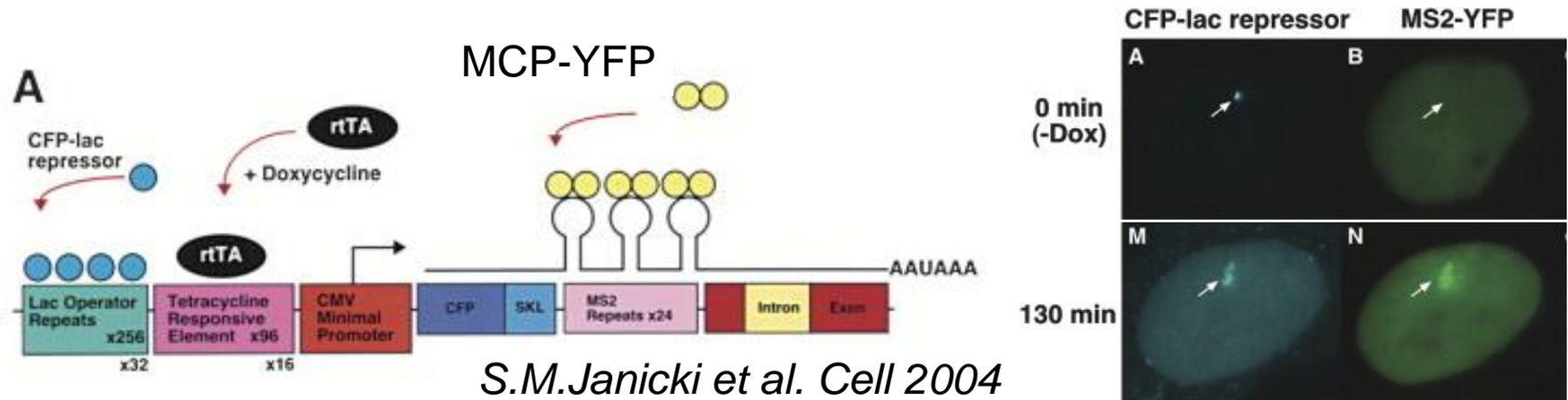
Figure 1 | Influence of cell medium on fluorescent protein photostability. Normalized bleaching curves for EGFP in live HEK293T cells maintained in DMEM, DMEM - Rf or DMEM - V. Error bars, s.d. ($n = 20$ cells).



Fluorescent RNA imaging

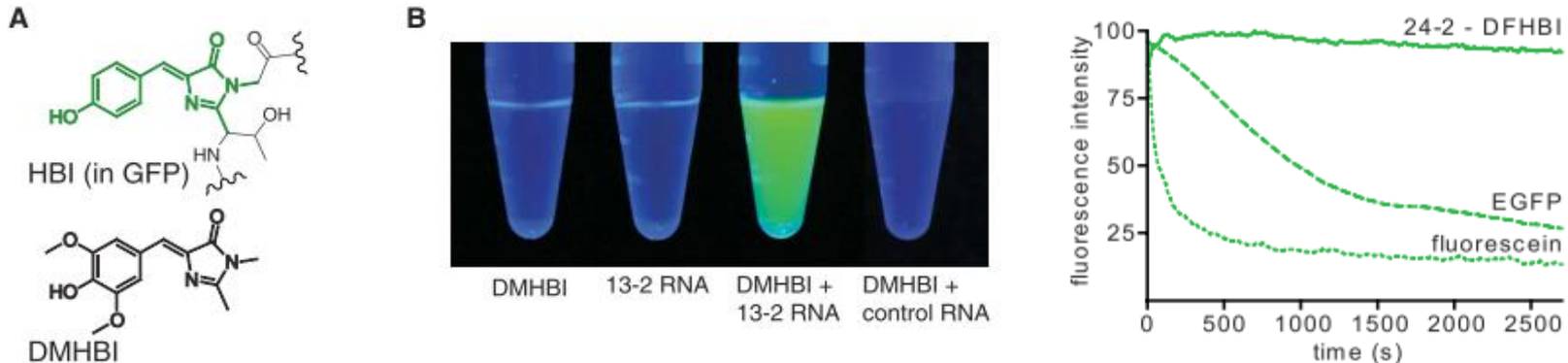
See single mRNA: MS2 mRNA stem-loops bound by MCP-YFP

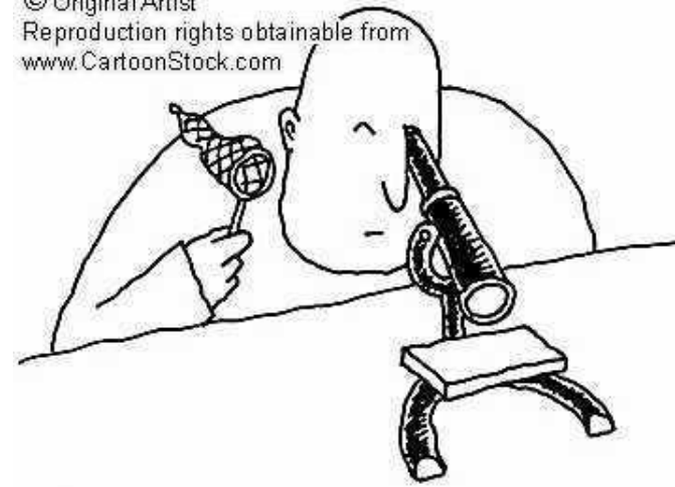
See product of translation: mRNA encodes CFP-SKL which goes to peroxisomes



Spinach RNA 60 nt aptamer binds cell-permeable fluorogenic dye

Photostable. Used to label 5S RNA in HEK cells. *Samie Jaffrey Science 2011*





Overview

1. What kind of structures are fluorescent
2. How to make and target fluorescent probes
3. Fluorescent probes for cellular structure and function
4. Using light to control cells

Why use light to control biology?

Light control allows extreme temporal and spatial control.

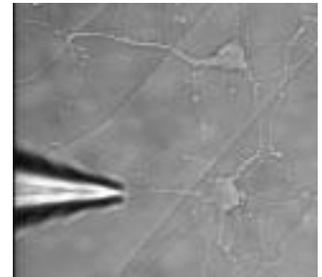


Temporal control

genes < chemicals < light
min-hr *s-min* $\mu\text{s-s}$

Spatial control

chemicals / genes < light
one or many cells *1 μm part of cell*



(note micropipettes for precise small molecule delivery)

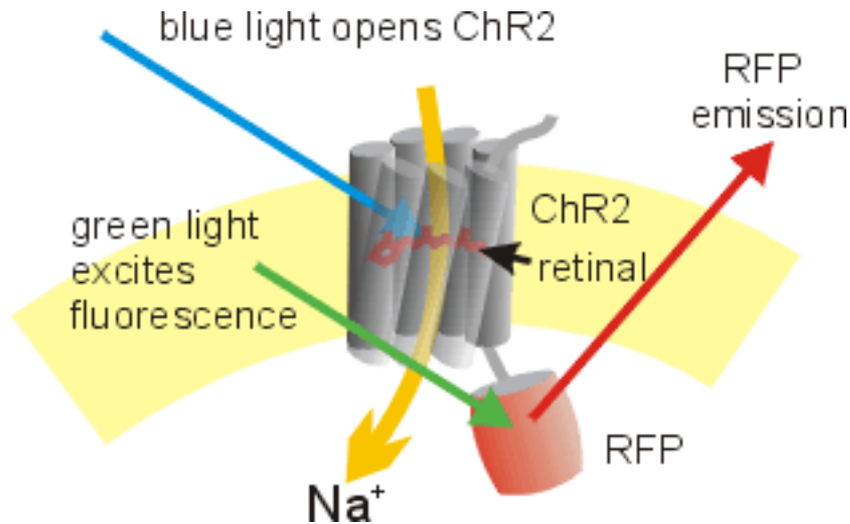
(often combine chemical/light control or gene/light control)
optogenetics/chemogenetics

Limitations of light? \$\$\$\$\$

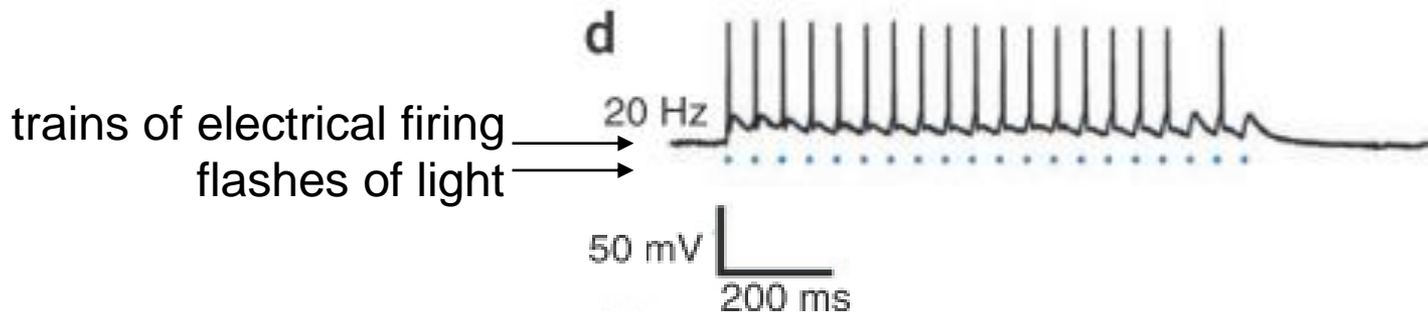
and usually data on one cell at a time

Controlling biology with light: light-gated ion channels

Channelrhodopsin from an alga, like rhodopsin, undergoes retinal isomerisation in response to light, and changes conformation, but opens a Na^+ channel. This allows light to control membrane voltage and trigger neuron firing.



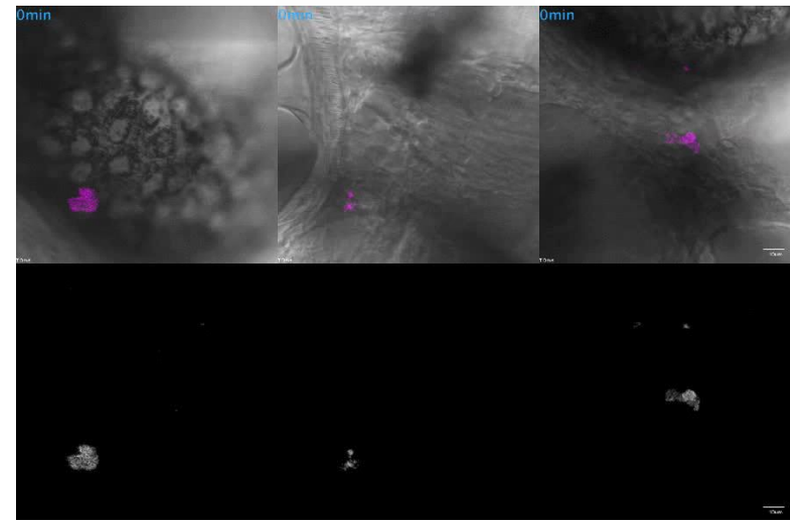
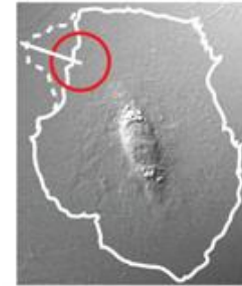
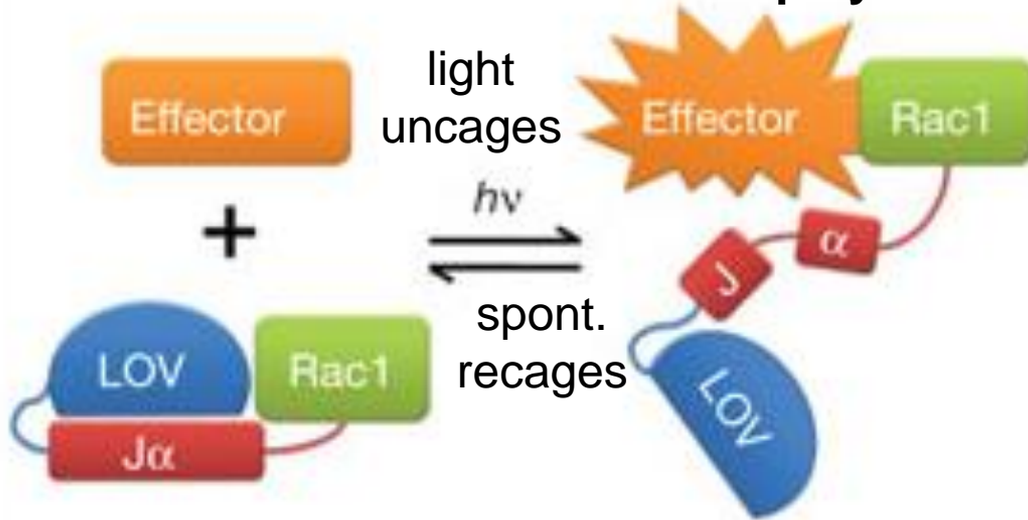
to understand neuronal firing patterns
to control secretion in diabetes
potentially in fixing neural diseases?
e.g. damping down overactivity in
epilepsy



Genetically-encoded photoactivation

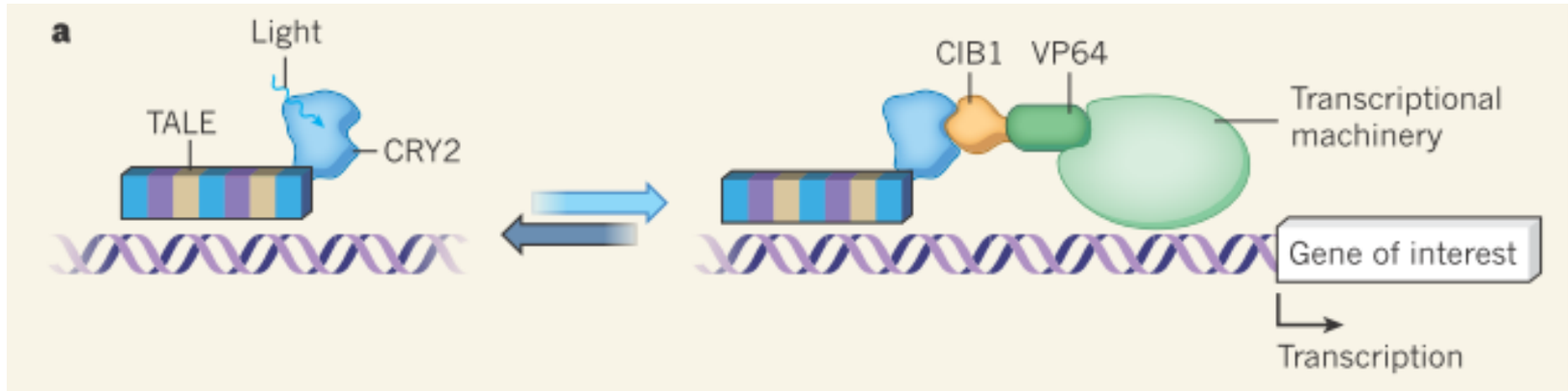
a

local actin polymerisation



1. Constitutively active Rac mutant
 2. Optimise LOV-Rac junction,
 3. knockout GTP hydrolysis and GAP/GNDI/GEF interactions
- K_d for PAK 2 μ M in dark, 200nM in light 10-fold ratio
Interaction of Rac with PAK stimulates cell protrusion and migration.

Photoactivation of transcription



1. Transcription-activator-like effector (TALE) for targeting arbitrary DNA sequence
 2. 10-fold induction
 3. Induction in minutes in animals
- Modular design (can also recruit nuclease or repressive domain)

A. Moglich et al. Nature Aug 2013

Conclusions

Choosing the right dye or fluorescent protein can make a big difference for:

- sensitivity
- signal stability
- modification to molecule/cell function
by size or multimerization

Fluorescent probes allow more than just following location:

- reporting cellular events
- uncaging biomolecule function
- controlling interactions and ion flux



References

Fluorescence probes

Molecular Probes Handbook, from Life Technologies
Principles of Fluorescence Spectroscopy 2nd edition,
by Joseph R. Lakowicz.

Protein modification

Bioconjugate Techniques, 2nd Edition
by Greg T. Hermanson.

Chemical labeling strategies for cell biology, Marks
KM, Nolan GP. Nat Methods. 2006 Aug;3(8):591-6.

Fluorescent proteins

(i) See table at

<http://nic.ucsf.edu/FPvisualization/>

(ii) Quantitative assessment of fluorescent proteins
PJ Cranfill, DW Piston et al. Nature Methods 2016

(iii) as sensors: Designs and applications of
fluorescent protein-based biosensors.

Ibraheem A, Campbell RE.

Curr Opin Chem Biol 2010;14:30-6

