2020/2021 - Light microscopy in Cellular Biology

Fluorescence Microscopy

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FLUORESCENCE A SMALL SUMMARY

- What is it?
- Where does it come from?
- Advantages
- Disadvantages

- Chromophores are components of molecules which absorb light
- e.g. from protein most fluorescence results from the indole ring of tryptophan residue
- They are generally aromatic rings

FLUORESCENCE

ABS - Absorbance S 0.1.2 - Singlet Electronic Energy Levels FL - Fluorescence T 1,2 - Corresponding Triplet States I.C.- Nonradiative Internal Conversion IsC - Intersystem Crossing PH - Phosphorescence

FLUORESCENCE MICROSCOPY: BASICS OF THEORY

- Absorbance spectrum limits excitation.
- Energy states limit excitation
- Molecule returns to lowest vibrational state emitting heat
- Light is emitted on return to ground state

Excited state

excitation

shorter wavelength, higher energy

Ground state

→ Stoke's shift

emission

longer wavelength, less energy

Fluorophores (Fluorochromes, chromophores) Fluorophores (Fluorochromes, chromophores)

• Special molecular • Special molecular structure structure

• Aromatic systems (Pi-• Aromatic systems (Pisystems) and metal systems) and metal complexes (with complexes (with transition metals) transition metals)

• characteristic • characteristic excitation and emission excitation and emission spectra spectra

400nm

STOKE'S SHIFT

 \checkmark The emission spectrum of an excited fluorophore is usually shifted to longer wavelengths when compared to the absorption or excitation spectrum

> Excitation 495 nm Emission: 520 nm

- The intensity of the fluorescence is very weak in comparison with the excitation light (10 -3 to 10 -5).
- The emitted light re-radiates spherically in all directions.
- Dark background is required to enhance resolution.

STOKE'S SHIFT

As Stokes' shift values increase, it becomes easier to separate excitation from emission light through the use of fluorescence filter combinations.

•produces yellow-orange emission

•to achieve maximum fluorescence intensity

•a fluorophore is usually excited at wavelengths near or at the peak of the excitation curve,

•And detected at widest possible range of emission wavelengths that include the emission peak

 (e)

Wavelength (nm)

PARAMETERS

• Extinction Coefficient

- E refers to a single wavelength (usually the absorption maximum)
- **Quantum Yield**
	- \bullet \mathbf{Q}_{f} is a measure of the integrated photon emission over the fluorophore spectral band

• At sub-saturation excitation rates, fluorescence intensity is proportional to the product of ε and Q_{f}
Mumber of emitted photons φ = Number of absorbed photons

• Lifetime $1 - 10 \times 10^{-9}$ secs (1-10 ns)

FLUORESCENCE

Stokes Shift

• is the energy difference between the lowest energy peak of absorbance and the highest energy of emission

Wavelength

Intensity

related to the **probability** of the event

Wavelength

the **energy** of the light absorbed or emitted

MULTICHANNEL FLUORESCENCE LABELLING

• Direct coupling to macromolecules

- Fluorescent dyes and substrates
- Fluorescent fusion proteins
- Fluorescent Antibodies

Ch1(Green) UBI-GFP

Ch2(Red) Texas Red anti-rabbit & Rabbit anti- **BGal**

Arterial edothelial cell Ch1(Green) FITC Tubulin Ch2(Red) mitotracker Ch3(Blue) DAPI

PHOTOBLEACHING

- Defined as the irreversible destruction of an excited fluorophore (discussed in later lecture)
- Methods for countering photobleaching
	- Scan for shorter times
	- Use high magnification, high NA objective
	- Use wide emission filters
	- Reduce excitation intensity
	- Use "antifade" reagents (not compatible with viable cells)

Not a chemical process

Dynamic quenching =- Collisional process usually controlled by mutual diffusion Typical quenchers – oxygen Aliphatic and aromatic amines (IK, NO2, CHCl3)

Static Quenching

Formation of ground state complex between the fluorophores and quencher with a non-fluorescent complex (temperature dependent – if you have higher quencher ground state complex is less likely and therefore less quenching

- Many quenchers act by reducing oxygen concentration to prevent formation of singlet oxygen
- Satisfactory for fixed samples but not live cells!
- Antioxidents such as propyl gallate, hydroquinone, pphenylenediamine are used
- \bullet Reduce O_2 concentration or use singlet oxygen quenchers such as carotenoids (50 mM crocetin or etretinate in cell cultures); ascorbate, imidazole, histidine, cysteamine, reduced glutathione, uric acid, trolox (vitamin E analogue)

PROBES FOR PROTEINS

PROBES FOR NUCLEIC ACIDS

DNA PROBES

• AO

- Metachromatic dye
	- concentration dependent emission
	- double stranded NA Green
	- single stranded NA Red
- AT/GC binding dyes
	- AT rich: DAPI, Hoechst, quinacrine
	- GC rich: antibiotics bleomycin, chromamycin olivomycin, rhodamine 800

INDO-1: 1H-Indole-6-carboxylic acid, 2-[4-[bis[2-[(acetyloxy)methoxy]-2- oxoethyl]amino]-3- [2-[2-[bis[2- [(acetyloxy)methoxy]-2-oxoetyl]amino]-5- methylphenoxy]ethoxy]phenyl]-, (acetyloxy)methyl ester $[C_{47}H_{51}N_3O_{22}]$ (just in case you want to know....!!)

FLUO-3: Glycine, N-[4-[6-[(acetyloxy)methoxy]-2,7- dichloro-3-oxo-3H-xanthen-9-yl]-2-[2-[2- [bis[2-[(acetyloxy)methoxy]-2- oxyethyl]amino]-5- methylphenoxy]ethoxy]phenyl]-N-[2- [(acetyloxy)methoxy]-2-oxyethyl]-, (acetyloxy)methyl ester

PH SENSITIVE INDICATORS

SNARF-1: Benzenedicarboxylic acid, 2(or 4)-[10-(dimethylamino)-3-oxo-3H- benzo[c]xanthene-7-yl]- **BCECF:** Spiro(isobenzofuran-1(3H),9'-(9H) xanthene)-2',7'-dipropanoic acid, ar-carboxy-3',6'-dihydroxy-3-oxo-

PROBES FOR OXIDATION STATES

DCFH-DA: 2',7'-dichlorodihydrofluorescein diacetate (2',7'-dichlorofluorescin diacetate; H2DCFDA)

DCFH-DA - dichlorofluorescin diacetate HE - hydroethidine 3,8-Phenanthridinediamine, 5-ethyl-5,6-dihydro-6-phenyl-DHR-123 - dihydrorhodamine 123 Benzoic acid, 2-(3,6-diamino-9H-xanthene-9-yl)-, methyl ester $C_{24}H_{16}C_{12}O_7$ $C_{21}H_{21}N_3$ $C_{21}H_{18}N_2O_3$

SPECIFIC ORGANELLE PROBES

BODIPY - borate-dipyrromethene complexes NBD - nitrobenzoxadiazole DPH – diphenylhexatriene TMA - trimethylammonium

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OTHER PROBES OF INTEREST

• GFP - *Green Fluorescent Protein*

- GFP is from the chemiluminescent jellyfish *Aequorea victoria*
- excitation maxima at 395 and 470 nm (quantum efficiency is 0.8) Peak emission at 509 nm
- contains a p-hydroxybenzylidene-imidazolone chromophore generated by oxidation of the Ser-Tyr-Gly at positions 65- 67 of the primary sequence
- Major application is as a reporter gene for assay of promoter activity
- requires no added substrates

MULTIPLE EMISSIONS

- Many possibilities for using multiple probes with a single excitation
- Multiple excitation lines are possible
- Combination of multiple excitation lines or probes that have same excitation and quite different emissions
	- e.g. Calcein AM and Ethidium (**ex 488 nm**)
	- emissions **530 nm** and **617 nm**

FILTER COMBINATIONS

• The band width of the filter will change the intensity of the measurement

FLUORESCENCE OVERLAP

Fluorescence microscopy

– Principle and practical consideration

Fluorescence microscopy

Excites and observe fluorescent molecules

The most commonly used microscopy

High resolution, sensitive with low background, multi-channel…

comes with variations (fancy names).

deconvolution, OMX, deltavision confocal, spinning disc, two photon TIRF, FRAP, FRET, FLIM, iFRAP, FCS …

PALM, STED, STORM, SIM, (super-resolution)

still in development

What can you do with a fluorescence microscope?

For example:

Determine the localisation of specific (multiple) proteins

Determine the shape of organs, cells, intracellular structures

Examine the dynamics of proteins

Study protein interactions or protein conformation

Examine the ion concetration etc.

can observe in live cells

FLUORESCENCE TECHNIQUES

SPECIAL APPLICATIONS:

•FRET and FLIM •FRAP and photoactivation •TIRF

FRET (FLUORESCENCE RESONANCE ENERGY TRANSFER)

- method to investigate molecular interactions
- **Principle:** a close acceptor molecule can take the excitation energy from the donor (distance ca 1-10 nm)

FRET situation: Excitation of the donor (GFP) but **emission** comes from the acceptor (RFP)

No FRET

FRET

ways to measure:

• Acceptor emission

Detect the emission of the acceptor after excitation of the donor, e.g. excite GFP with 488 but detect RFP at 610 (GFP emission at 520)

• Donor emission after acceptor bleaching take image of donor, then bleach acceptor (with acceptor excitation wavelength - RFP:580nm), take another image of donor \rightarrow should be brighter!

FRET

• a suitable FRET pair (with overlapping excitation/emission curves)

- Bleed through (because of overlapping spectra)
- **Hytellimitation of techniques (filters etc)**
- Photobleaching only with fixed samples
- Intensity depends on concentrations etc

FLIM (FLUORESCENCE LIFETIME IMAGING MICROSCOPY)

- measures the lifetime of the excited state (delay between excitation and emission)
- every fluorophore has a unique natural lifetime
- lifetime can be changed by the environment, such as:
	- \checkmark lon concentration Oxygen concentration $\mathbf{v}_{\rm pH}$ **Protein-protein interactions**

FLIM

Excitation of many electrons at the same time \rightarrow count the different times when they are falling back down (i.e. photons are emitted)

lifetime $= \frac{1}{2}$ of all electrons are fallen

EXAMPLE OF FLIM-FRET MEASUREMENT

GFP expressed in COS 1 cell: average lifetime of 2523 ps

fused GFP-RFP expressed in COS 1 cell: average lifetime of 2108 ps

Joan Grindlay, R7

You still need: a suitable FRET-pair with the right orientation of the π-orbitals

 \rightarrow Interaction of proteins is not enough, because fluorophores have to be close enough and in the right orientation!

Use of FLIM: measurements of concentration changes (Ca2+), pH change etc, Protein interactions

→ FRET: Leica confocal 2 or Olympus FV 1000

FLIM: Leica confocal 1 and soon LIFA system from Lambert Instruments

SPECIAL APPLICATIONS:

• FRET and FLIM •**FRAP and photoactivation** • TIRF

FRAP (FLUORESCENCE RECOVERY AFTER PHOTOBLEACHING)

• Intense illumination with 405 laser bleaches the sample within the selected region Δ chequipation of the recovery

to measure the mobility/dynamics of proteins under different conditions

Olympus FV 1000

PHOTOACTIVATION

• Fluorophore only becomes active $(=$ fluorescent) if excited (e.g. with 405 laser) due to structural change

Pictures taken from a activation movie: activation of a line trough the lamellipodia of the cell, activated GFP_F diffuses quickly

→ Olympus FV 1000

SPECIAL APPLICATIONS:

- FRET and FLIM
- FRAP and photoactivation
- •**TIRF**

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TIRF (TOTAL INTERNAL REFLECTION **FLUORESCENCE)**

You need:

- TIRF objectives with high NA
- TIRF condensor, where you are able to change the angle of illumination
- Glass coverslips

very thin section at the bottom of the sample \rightarrow 150-200nm to study membrane dynamics (endocytosis, focal adhesions, receptor binding)

 \rightarrow Nikon TE 2000

TIRF

TIRF VS EPI

TIRF VS EPI

SUMMARY/COMPARISON

