Imaging at the molecular level: Measuring molecular motion and interactions

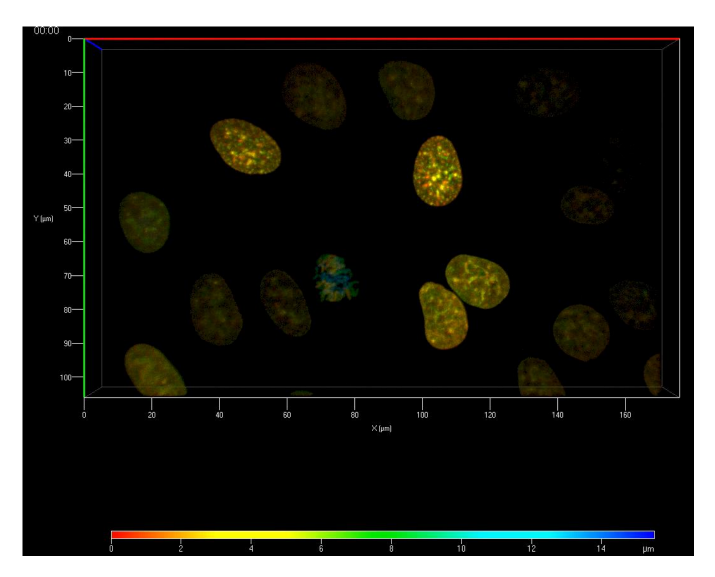
Chris Lagerholm Wolfson Imaging Centre Oxford Weatherall Institute of Molecular Medicine

Imaging at the molecular level: Measuring molecular motion and interactions

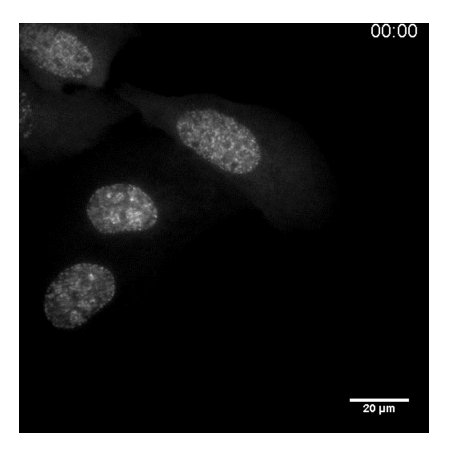
- $\cdot\,$ Brief intro to theory of molecular motion
- <u>Fluorescence techniques for measuring molecular motion</u>
- Single particle tracking (SPT)
- Fluorescence Recovery after Photobleaching (FRAP)
- Fluorescence Loss in Photobleaching (FLIP)
- Fluorescence localization after Photobleaching (FLAP)
- Fluorescence Correlation Spectroscopy (FCS)
- Raster Image Correlation Spectroscopy (RICS)
- <u>Fluorescence techniques for measuring molecular interactions</u>
- Förster Resonance Energy Transfer (FRET)
- Fluorescence Cross-Correlation Spectroscopy (FCCS)

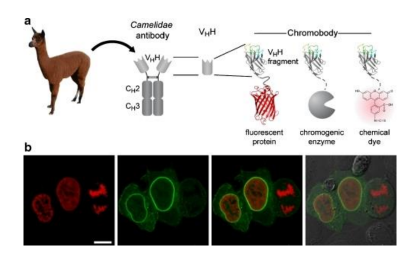
Many biological samples are dynamic at a wide range of time-scales

Example 1: Cell cycle; 3D re-construction of HeLa cells expressing H2B-GFP imaged on Spinning-disc Confocal at 6 Z-stacks/h

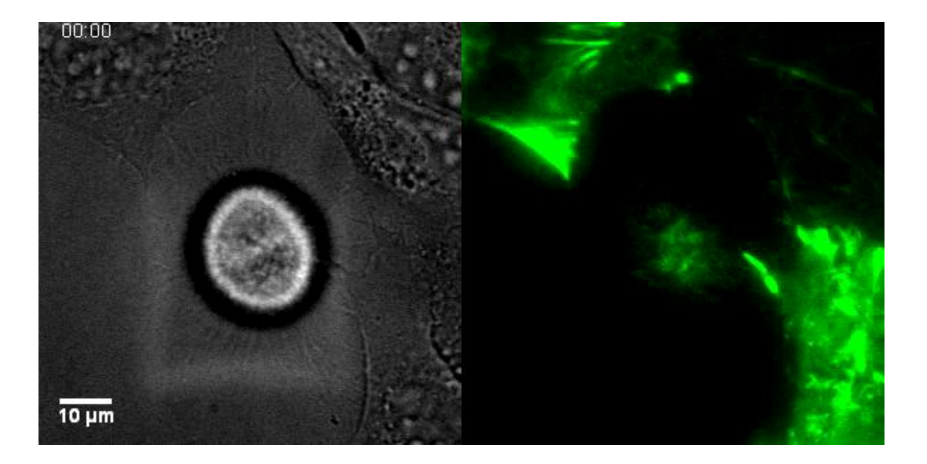


Example 2: Cell cycle; U2OS cells expressing PCNA-Chromobody-RFP imaged on Widefield/TIRF microscope at 24 z-planes/h

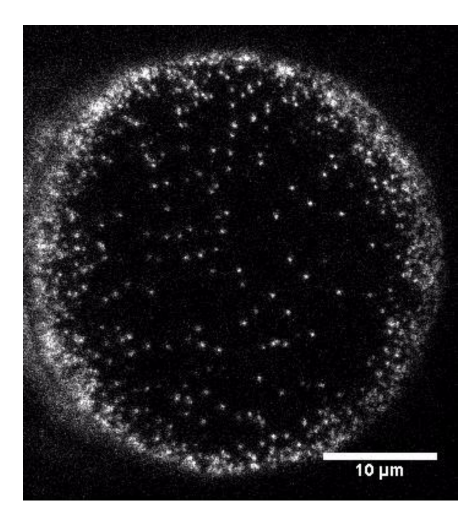




Example 3: Mitosis; HeLa cells expressing LifeAct-RFP imaged in TIRF microscope at 24 z-planes/h



Examples 4: Molecular motion of phospholipid analogue (KK114-PE) in substrate supported bilayer (SLB) imaged in TIRF mode at 65 Hz



- In but time-lapse imaging alone does not reveal quantitative information about movements of cell organelles, proteins, lipids, or DNA
- In particular, time lapse imaging is not sufficient to differentiate between different modes of motion of a molecule i.e.
 - 1. Random (Brownian) diffusion
 - 2. Directed active (motor driven) transport
 - 3. Reversible bound and moving by 1, 2, or combination thereof
 - 4. ...

Brief intro to theory of molecular motion

ON THE MOVEMENT OF SMALL PARTICLES SUSPENDED IN A STATIONARY LIQUID DEMANDED BY THE MOLECULAR-KINETIC THEORY OF HEAT

I N this paper it will be shown that according to the molecular-kinetic theory of heat, bodies of microscopically-visible size **suspended** in a liquid will perform movements of such magnitude that they can be easily observed in a microscope, on account of the molecular motions of heat. It is possible that the movements to be discussed here are identical with the so-called "Brownian molecular motion"; however, the information available to me regarding the latter is so lacking in precision, that I can form no judgment in the matter (I).

$$\left\langle (x(t) - x_0)^2 \right\rangle = \int_{-\infty}^{\infty} x^2 N[0, 2Dt] \, dx = 2Dt$$

Einstein, A. (1905). Über die von der molekularkinetischen Theorie der Wärme geforderte Bewegung von in ruhenden Flüssigkeiten suspendierten Teilchen. *Annalen der Physik, 17,* 549-560.

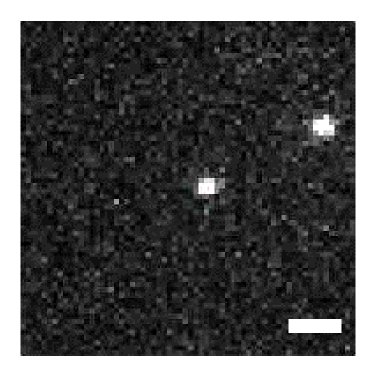
Einstein, A. (1956). *Investigations on the theory of the Brownian movement* (A. D. Cowper, Trans.). New York, NY: Dover Publications.

Fluorescence techniques for measuring molecular motion

- 1) Single particle tracking (SPT)
- 2) Fluorescence Recovery after Photobleaching (FRAP)
- 3) Fluorescence Loss in Photobleaching (FLIP)
- 4) Fluorescence localization after Photobleaching (FLAP)
- 5) Fluorescence Correlation Spectroscopy (FCS)
- 6) Raster Image Correlation Spectroscopy (RICS)

1) Single particle tracking (SPT)

anti-CD73 Fab'-biotin + sAv-605 Qdot



189 Hz (5.3 ms integration) with 160X magnification and Andor EMCCD

Playback 100 Hz, 2000 frames Scale bar = 1µm

Single Particle Tracking

• Time lapse imaging of spatially resolved single particles, molecules, or subcellular structures

 Technique results in time trajectories of sub-pixel positions of single objects

- Such trajectories contain information about:
 - diffusion coefficients
 - velocities
 - step sizes
 - spatial and temporal confinement

Probe Considerationsfor SPT

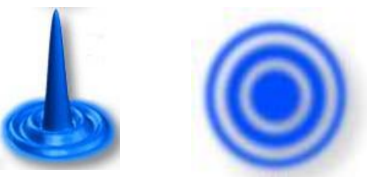
- Very bright and stable
- Small
- Monovalent (i.e. one probe per molecule of interest)
- Low non-specific binding

Typical SPT Probes

	Gold	Cy3	Quantum dots
Means of detection	Scattering	Fluorescence	Fluorescence
Size (diameter)	40 nm	~1-2 nm	~10-20 nm
Advantages	Photostable	Small Monovalent	Photobleaching resistant
Disadvantages	Immobilization Large Multivalent	Photobleaching (~5 s)	Non-specific binding Blinking

Experimental considerations for SPT

• Because of diffraction, the image of point source as focused by a microscope is an **Airy pattern**



• The center of the Airy pattern is known as the Airy disk

 $r_{Airy} = \begin{array}{l} \frac{0.61 \lambda}{NA_{objective}} \end{array} \qquad \begin{array}{l} \lambda = wavelength \ of \ light \\ NA=numerical \ aperture=n \ sin \ \alpha \\ n = index \ of \ refraction \ of \ immersion \ media \\ \alpha = half-angle \ of \ angular \ aperture \end{array}$

•The Airy disk represents 84 percent of the total luminous energy

- NA ranges from <0.95 for air, <1.2 for water, <1.4 for oil
- For 100X magnification, 1.4 NA oil immersion objective and illumination with green light (500 nm = $0.5 \mu m$)

$$r_{Airy} = \frac{(0.61) (0.5 \ \mu m)}{1.4} \approx 220 \ nm$$

Microscope Resolution

• The diffraction of light causing the Airy disk is also the limiting factor of the resolution of a microscope

•The resolution of a microscope is defined as the minimum distance two objects have to be separated by to be resolved as two separate objects



Airy patterns of two point sources

• For 100X magnification, 1.4 NA oil immersion objective and illumination with green light (500 nm = $0.5 \ \mu m$)

Resolution =
$$\frac{(0.61) (0.5 \,\mu\text{m})}{1.4} \approx 220 \,\text{nm}$$

Sampling considerations in SPT

• Single Particle (Molecule) Tracking relies on intensified video or CCD cameras both of which utilizes arrays of **square pixels** for detection

• For optimum resolution data should be sampled at about 1/3 X the resolution. This is known as **Nyquist Sampling**

Nyquist Sampling =
$$\frac{(0.61) (0.5 \,\mu\text{m})}{3 \, 1.4} \approx 70 \,\text{nm}$$

• Hence for 100X magnification, 1.4 NA oil immersion objective and illumination with green light (500 nm = $0.5 \ \mu m$)

• Proper sampling then requires $100 \times 70 \text{ nm} = 7 \mu \text{m}$ pixels

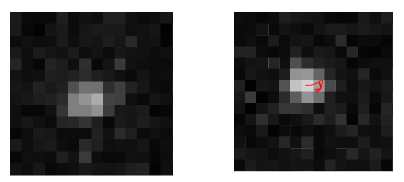
Data Analysis

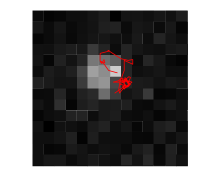
• Obtain sub-pixel resolution by curve fitting to determine centroids of single molecules

•Approximate Airy pattern with a 2D spatial Gaussian and fit each image

$$\boldsymbol{A} + \frac{\boldsymbol{B}}{2\pi \boldsymbol{w}^2} \boldsymbol{E} \boldsymbol{x} \boldsymbol{p} \left[-\frac{1}{2\pi \boldsymbol{w}^2} \left((\boldsymbol{x} - \boldsymbol{x}_{\boldsymbol{o}})^2 + (\boldsymbol{y} - \boldsymbol{y}_{\boldsymbol{o}})^2 \right) \right]$$

where ω = width of the PSF of the microscope (~220 nm)



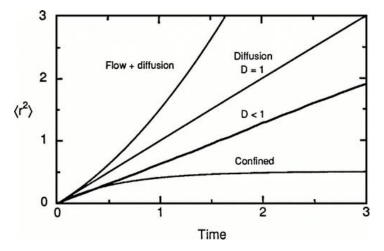


• Dynamics are analyzed with respect to that predicted by Brownian (random) motion in a 2 D (or 3D) fluid

• For 2D, the mean square displacement (MSD) is

$$< \mathbf{r}^{2} >= \frac{\prod_{k=1}^{q} \sqrt{(\mathbf{x}_{t1} - \mathbf{x}_{t2})^{2} + (\mathbf{y}_{t1} - \mathbf{y}_{t2})^{2}}}{q} = 4\mathbf{D}\Delta t \qquad \Delta t = t_{2} - t_{1}$$

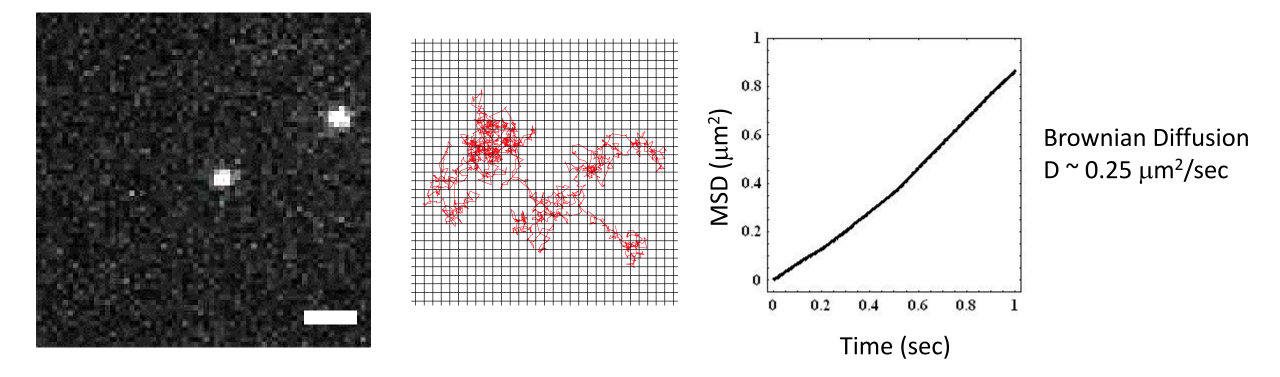
where D is the diffusion coefficient



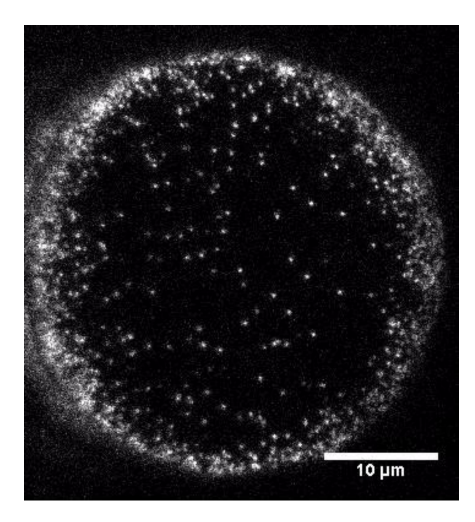
Brownian motion $\langle r^2 \rangle = 4Dt$ Anomalous diffusion $\langle r^2 \rangle = 4Dt^{\alpha}$ $\alpha < 1$ Diffusion with flow $\langle r^2 \rangle = 4Dt + (Vt)^2$ Confined diffusion $\langle r^2 \rangle \approx \langle r_c^2 \rangle [1 - A_1 \exp(-4A_2Dt / \langle r_c^2 \rangle)]$

Saxton and Jacobson (1997) Annu. Rev. Biophys. Biomol. Struct. 26: 373-99

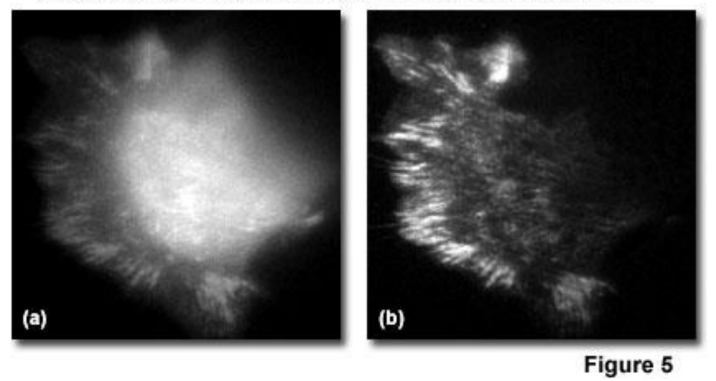
Mode of Motion & Diffusion Coefficient



Examples 4: Molecular motion of phospholipid analogue (KK114-PE) in substrate supported bilayer (SLB) imaged in TIRF mode at 65 Hz

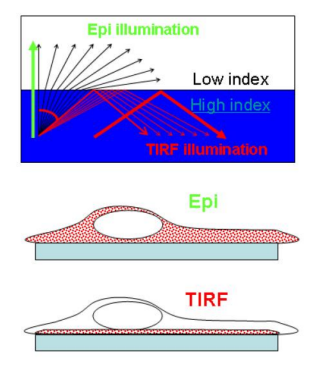


Cell Focal Adhesions in Widefield and TIR Fluorescence



http://www.microscopyu.com/articles/fluorescence/tirf/tirfintro.html

How does TIRF microscopy work?



In TIRF, only those fluorophore molecules that are near the cover glass surface are excited When the incident angle is greater than the critical value, the laser light undergoes **total internal reflection** from the glass-water interface.

Living Cell (n = 1.33 - 1.37)

Fluorophores Fluorophores Cell

Glass Slide (n = 1.518)

Evanescent Wave ---

Laser

Excited

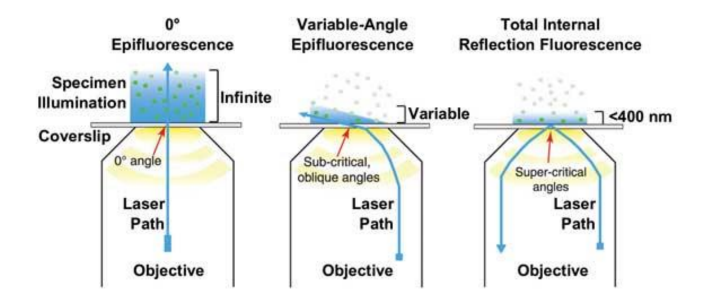
Reflection

Angle

Membrane

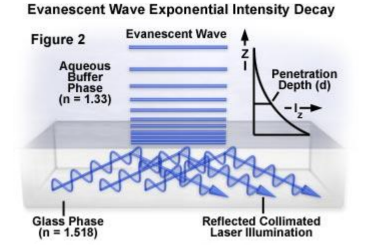
Interface

http://www.olympusmicro.com/primer/techniques/fluorescence/tirf/tirfhome.html

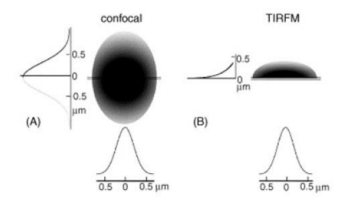




Advantages of the evanescent field



Point-spread function for confocal versus TIRF

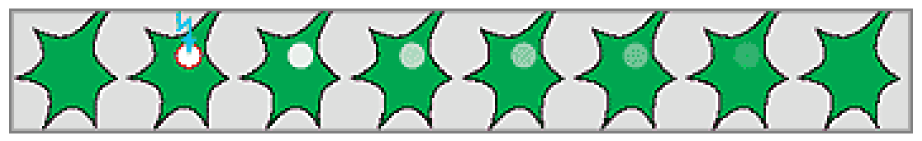


FWHM of p	point-spread function:
Confocal	300 nm X 800 nm
TIRF	250 nm X ~100 nm (length constant)

2) Fluorescence Recovery after Photobleaching (FRAP)

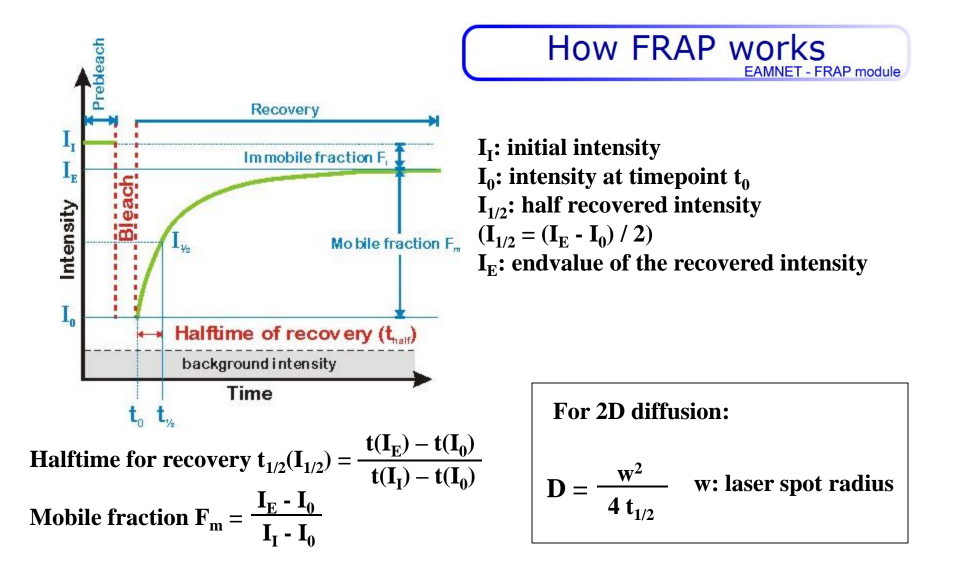
 Select photobleaching of region of interest (ROI) with very high laser intensity
Monitor the progress of fluorescence recovery in the bleached area with high temporal resolution

- Changes in intensity in the bleached region represent the sum of all movements of fluorescent molecules, whether passive (e.g., diffusion) or active (e.g., transport).
- The half-recovery time is a measure of the average speed of protein movement.



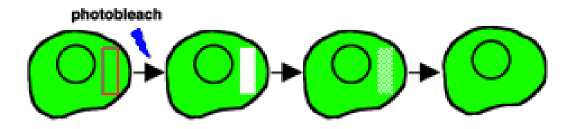
Bleached spot size ~ 1 μm

(www.zeiss.com)

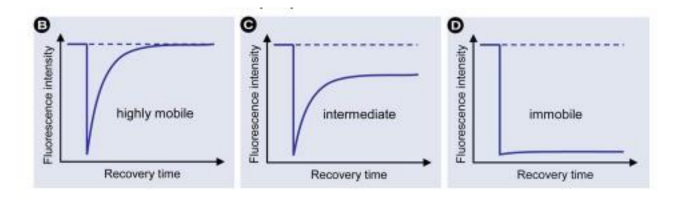


http://www.embl-heidelberg.de/eamnet/frap/html/how_frap_works.html

Can also do FRAP with most confocal microscopes



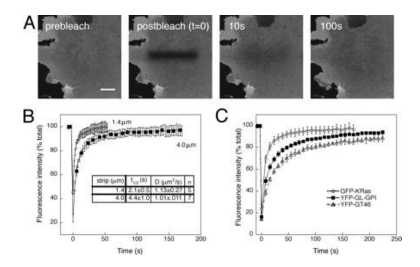
In this geometry, D is typically calculated numerically by comparing fluorescence recovery curves with simulated curves
(See i.e. http://www.embl.de/eamnet/html/frap_analysis.html)



Dynamics of putative raft-associated proteins at the cell surface

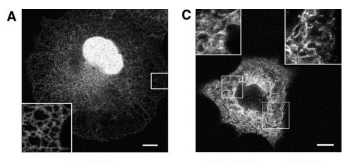
Anne K. Kenworthy,^{1,2,3} Benjamin J. Nichols,^{1,4} Catha L. Remmert,² Glenn M. Hendrix,² Mukesh Kumar,⁵ Joshua Zimmerberg,⁵ and Jennifer Lippincott-Schwartz¹

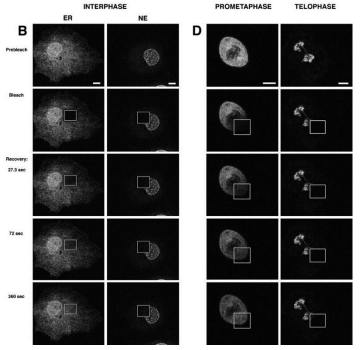
The Journal of Cell Biology, Volume 165, Number 5, June 7, 2004 735-746



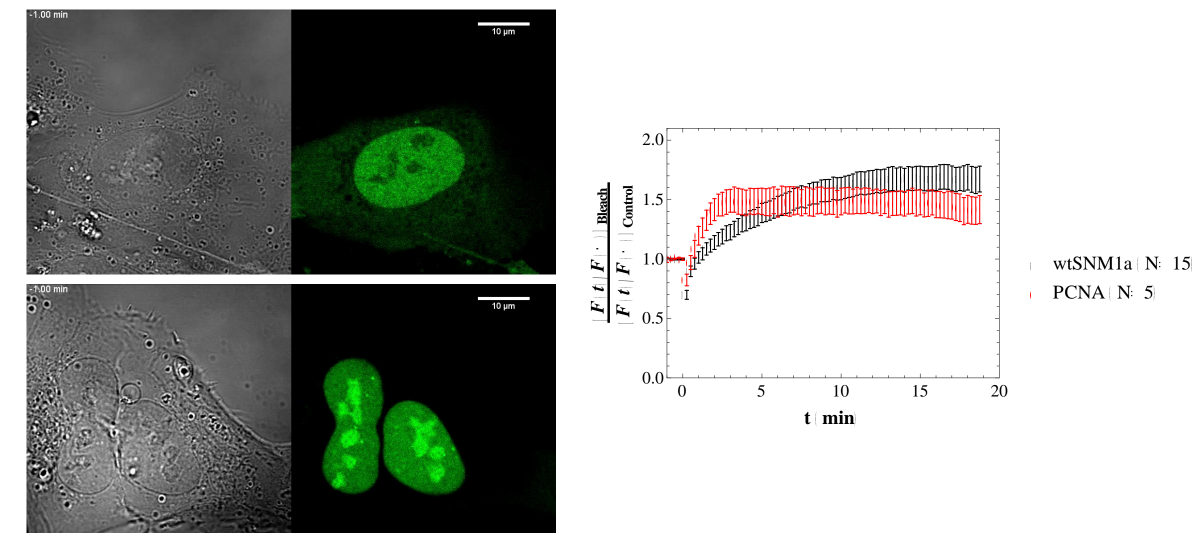
Large-scale lateral diffusion measurements by confocal microscopy. (A) Selected images from a confocal FRAP experiment at 37 °C of GFP-KRas expressed in COS-7 cells. Bleach box, 4 μ m wide. Bar, 10 μ m. (B) Kinetics of recovery for 1.4- (circles) versus 4- μ m-wide (squares) bleach box. Calculated D and t1/2 values are indicated. Data shown are for GFP-KRas expressed in COS-7 cells at 37 °C. (C) Kinetics of recovery for YFP-GT46 (triangles), YFP-GL-GPI (squares), and GFP-KRas (circles) in COS-7 cells at 37 °C using a 4- μ m-wide bleach box. Each curve shows the mean ± SD from seven to nine cells from a single experiment. The calculated Ds were as follows: GFP-KRas, 1.01 ± 0.11 μ m2/s; YFP-GL-GPI, 0.47 ± 0.07 μ m2/s; YFP-GT46, 0.23 ± 0.02 μ m2/s.

J. Ellenberg, E. D. Siggia, J. E. Moreira, C. L. Smith, J. F. Presley, H. J. Worman, and J. Lippincott-Schwartz (1997) Nuclear Membrane Dynamics and Reassembly in Living Cells: Targeting of an Inner Nuclear Membrane Protein in Interphase and Mitosis. J. Cell Biol. 138: 1193-1206





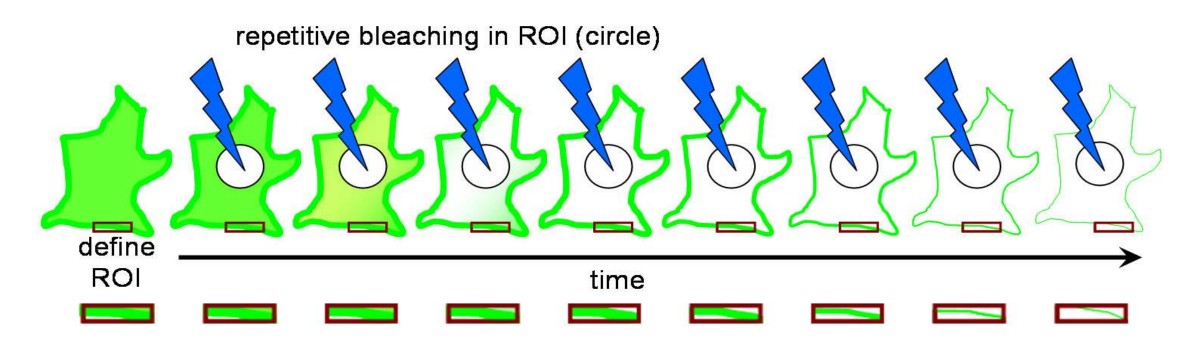
Distribution and mobilities of LBR-GFP in interphase and mitotic membranes. (A) Confocal section close to the lower cell surface showing steady-state expression of LBR-GFP in an interphase cell. (*Inset*) Boxed region at higher magnification showing LBR- GFP distribution within the ER network. (*B*) Qualitative FRAP experiments in ER and NE membranes in interphase cells expressing LBR-GFP. (*Left*) Photobleach recovery in ER membranes. (*Right*) Photobleach recovery in NE membranes. Note the complete recovery of fluorescence in the ER and the lack of recovery in the NE. (*C*) Thin confocal section through the mitotic apparatus showing the steady-state expression pattern of LBR-GFP in metaphase cells. (*Insets*) Boxed regions at higher magnification showing the tubular membrane network within which LBR-GFP redistributed. Note its resemblance to the interphase ER shown in *A*. (*D*) Qualitative FRAP experiments in mitotic membranes of cells expressing LBR-GFP. (*Left*) Photobleach recovery in telophase membranes. (*Right*) Photobleach recovery in telophase membranes. Example 3: DNA damage; U2OS cells expressing PCNA-GFP or wtSNM1A-GFP imaged on Confocal microscope at 240 z-planes/h



Characterization of role of *SNM1A*, a 5'–3' exonuclease, in DNA damage response (Lonnie Swift, Ghadir Almuhaini, Christoffer Lagerholm, Peter McHugh)

3) FLIP – Fluorescence Loss in Photobleaching

- a small region is repeatedly bleached and the loss of fluorescence in another region is measured.
- useful to show connectivity of compartments or for measuring turnover of molecules between different compartments



J. Ellenberg, E. D. Siggia, J. E. Moreira, C. L. Smith, J. F. Presley, H. J. Worman, and J. Lippincott-Schwartz (1997) Nuclear Membrane Dynamics and Reassembly in Living Cells: Targeting of an Inner Nuclear Membrane Protein in Interphase and Mitosis. J. Cell Biol. 138: 1193-1206

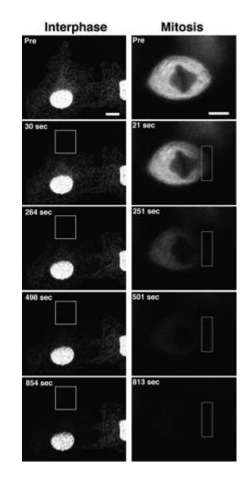
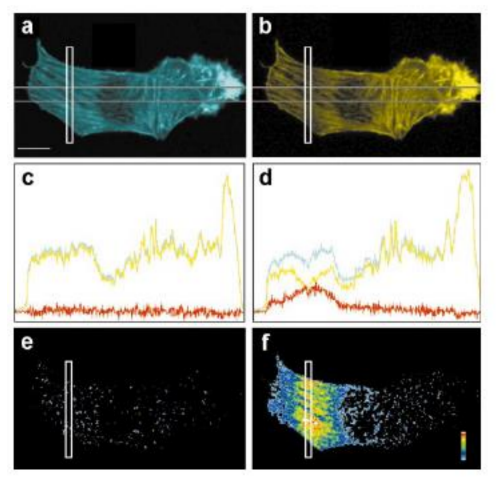


Fig. 6. FLIP to probe the continuity of interphase and mitotic membranes containing LBR-GFP. FLIP experiments were performed on interphase membranes (*left*) and metaphase membranes (*right*). Note the complete loss of fluorescence from both interphase ER membranes and mitotic membranes over a similar time course, but not from NE membranes in interphase. ER fluorescence that remained in interphase is from an adjacent cell whose membranes were not connected to those within the photobleached box. Bars, $10 \,\mu\text{m}$.

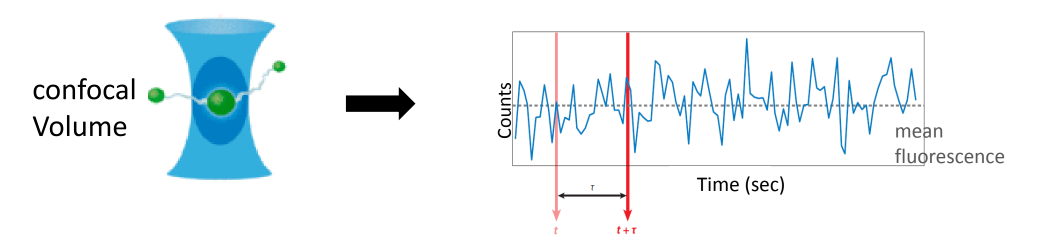
4) FLAP – Fluorescence Localization after Photobleaching

- In FLAP a protein is tagged with two fluorescent labels: one is photobleached and the other acts as a reference
- The use of a reference fluorochrome allows the tracking of the distribution of the labelled molecules by simple image differencing (I) and thus enables measurement of fast relocation dynamics.



Dunn, G. A., Dobbie, I. M., Monypenny, J., Holt, M. R., & Zicha, D. (2002). Fluorescence localization after photobleaching (FLAP): a new method for studying protein dynamics in living cells. *J Microsc, 205* (Pt 1), 109-112.

FCS – Autocorrelation of Fluorescence Fluctuations

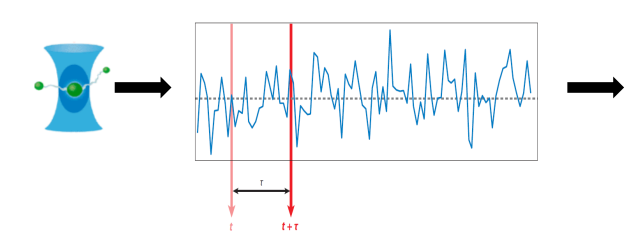


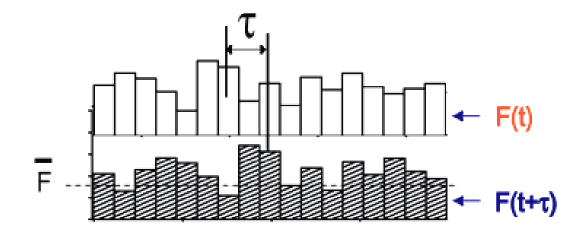
What causes fluctuations in intensity? a) Poison noise, b) diffusion of molecules



Bursts of photons are seen as single fluorophores diffuse in and out of the laser beam

FCS – Autocorrelation of Fluorescence Fluctuations



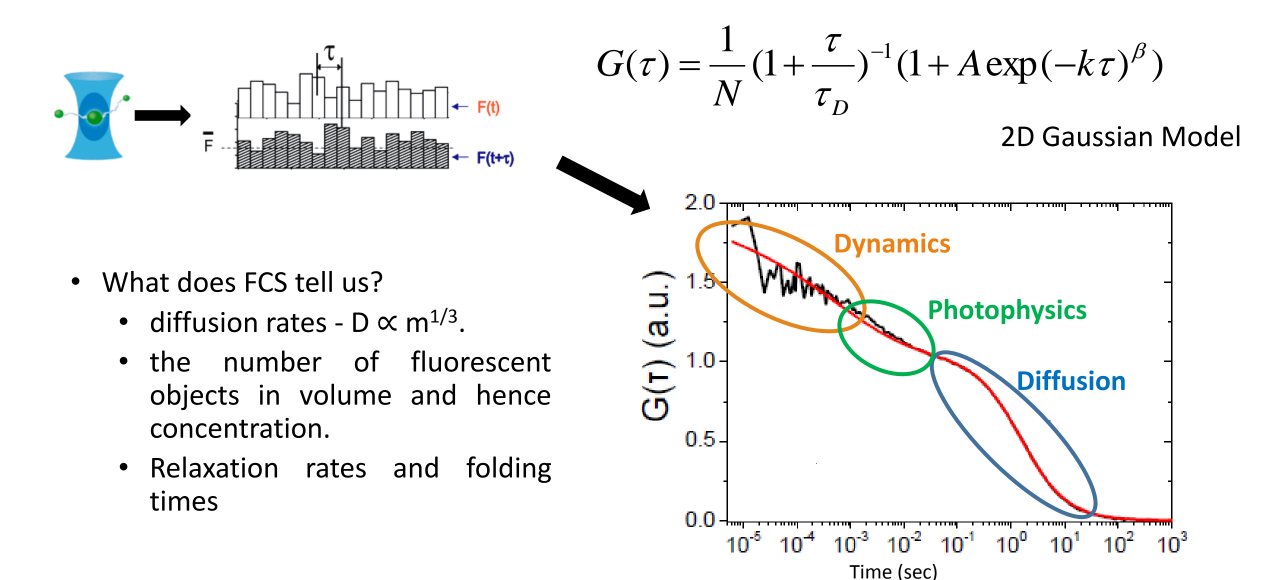


autocorrelation of fluorescence alterations

Autocorrelation describes the probability of finding the molecule still in the confocal volume at later times T

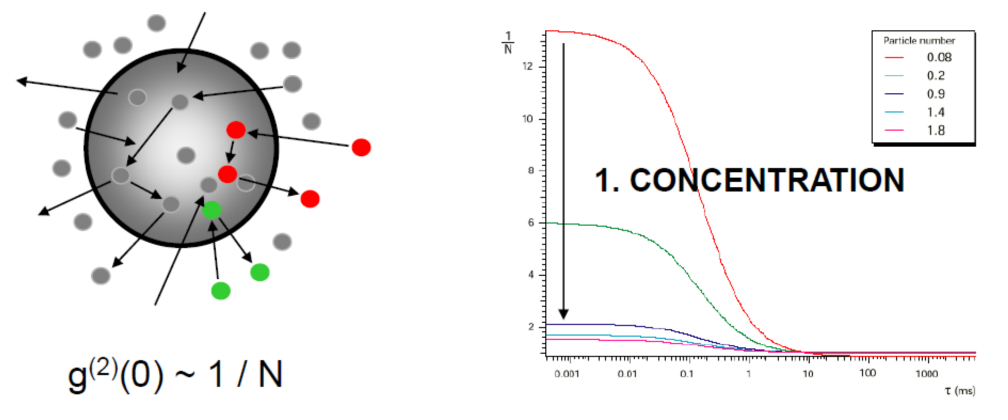
$$G(\tau) = \frac{\left\langle \delta F(t) \cdot \delta F(t+\tau) \right\rangle}{\left\langle \delta F(t) \right\rangle^2}$$
$$\delta F(t) = F(t) - \left\langle F(t) \right\rangle$$

FCS – Autocorrelation of Fluorescence Fluctuations



Fluorescence Correlation Spectroscopy

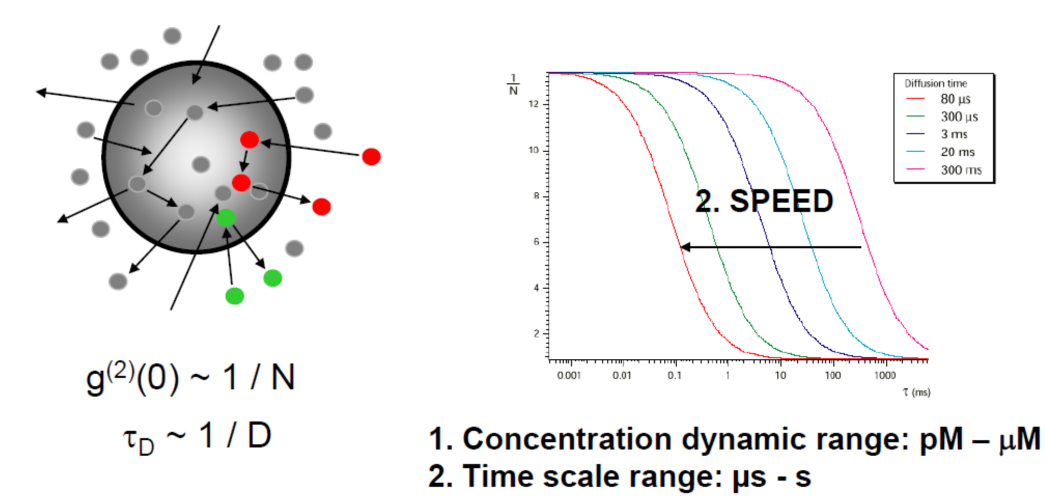
Wrapping it up: How many



1. Concentration dynamic range: pM - µM

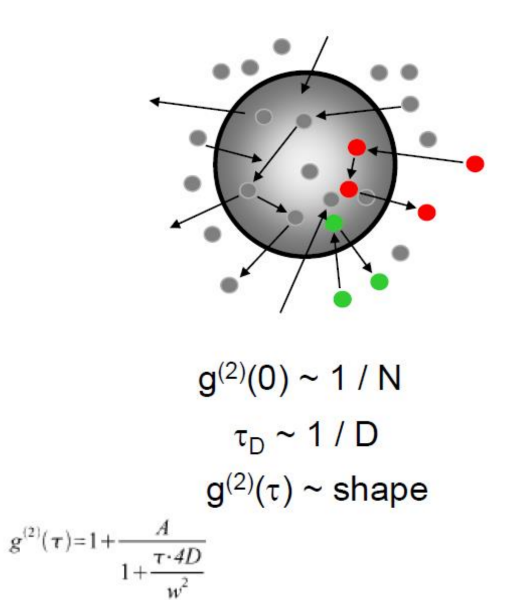
Fluorescence Correlation Spectroscopy

Wrapping it up: How many, how fast

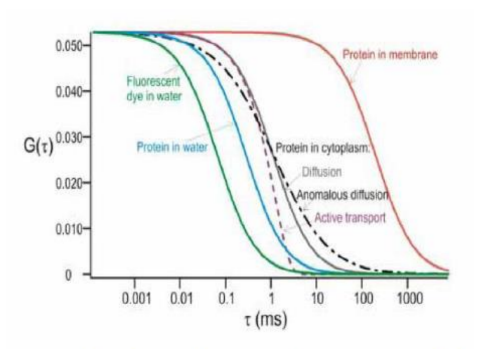


Fluorescence Correlation Spectroscopy

Wrapping it up: How many, how fast, what mechanism



3. TYPE OF MOVEMENT



- 1. Concentration dynamic range: $pM \mu M$
- 2. Time scale range: µs s
- 3. Type of dynamics: rafts, flow, free, ...

FCS – Conformational Dynamics

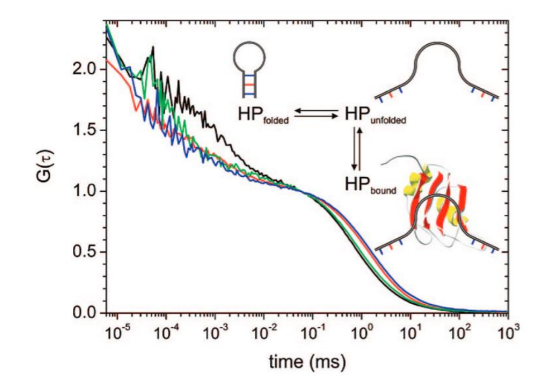
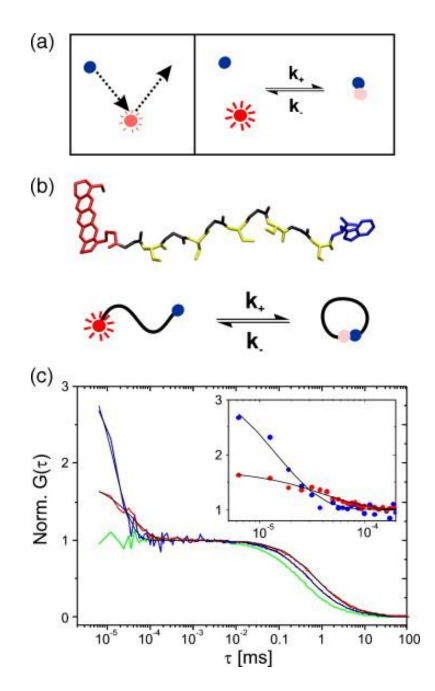


Figure 3. Normalized FCS curves measured for 10^{-9} M solutions of the RNA hairpin HP and the native ORN NT in the absence and presence of *At*GRP7 protein. The fluorescence correlation functions of free HP and NT are shown in black and green, respectively. The curve for HP bound to *At*GRP7 is shown in red and that for NT bound to *At*GRP7 in blue. The proposed model of a two-state hairpin-folding equilibrium whose unfolded state can be bound by the protein³³ is illustrated in the inset.

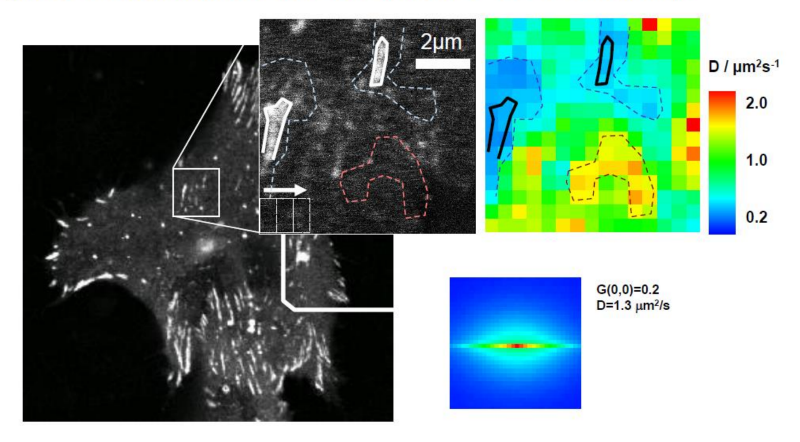


JACS., 2008, 130 (29)

6) RICS – Raster Image Correlation Spectroscopy

- RICS is a point confocal imaging form of FCS
- Technique is used to obtain quantitative diffusion maps of heterogeneous samples

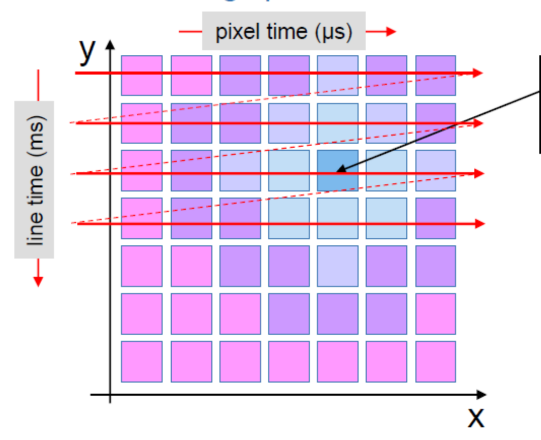
Mobility of Eos-Pax between cell adhesion structures – like a weather map.



Sample: Hari Shroff and Eric Betzig, Janelia Farm Research Campus, Ashburn, VA, USA

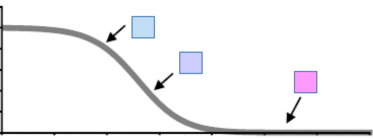
RICS = Raster Image Correlation Spectroscopy

In an image pixels are small time bins separated in space (ICS) In a raster scanned image pixels are time bins separated in space and time (RICS)



If something is happening in pixel A then, what is the probability of it happening in pixel B?

The answer is: It depends...



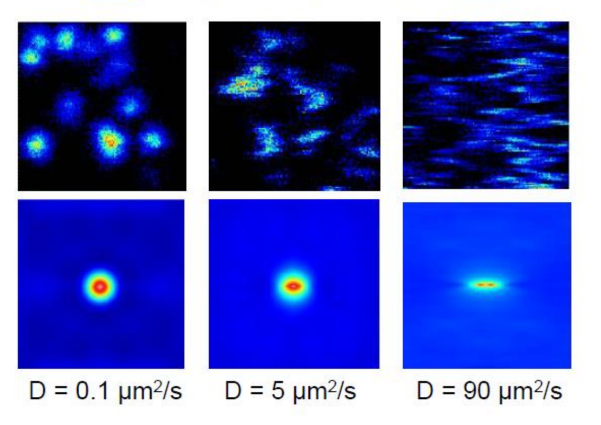
distance from pixel A and time delay from pixel A

What is shown here for one individual pixel is now repeated for all pixels...

The recipe is <u>exactly</u> the same!

$$g^{(2)}(\xi,\psi) = \frac{\langle I_r(x,y) \cdot I_r(x+\xi,y+\psi) \rangle}{\langle I_r \rangle^2}$$

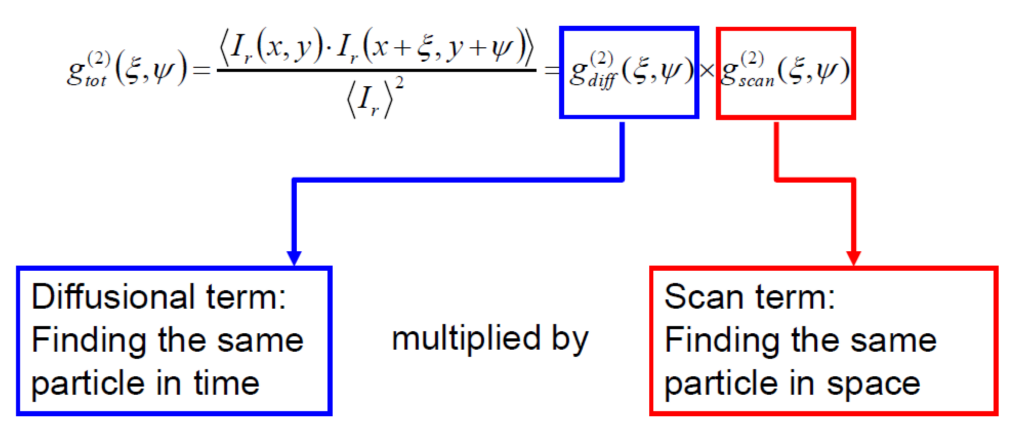
What can change is the pattern, but only if the objects move (if not: see ICS)



Raster scanned images of moving (sub-resolution) beads

This is the ICS applied to a raster scanned image of moving (sub-resolution) beads.

RICS analysis



Both are connected via the diffusion coefficient...

Summary - Fluorescence techniques for measuring molecular motion

•SPT can yield information about "heterogeneties" of single molecule behavior that is not apparent in ensemble average techniques and at a spatial resolution of ~ tens of nanometers

...but only for single molecules that are separated by a distance greater than ${\bf r}_{\rm Airy}$

• FRAP, FLIP, FLAP, FCS, and RICS are ensemble average techniques

• FCS is best choice for very fast dynamics including 3D diffusion in solution of small molecules

•Techniques can also be used to investigate binding constants and equilibria

Fluorescence techniques for measuring molecular interactions

- 1) Förster Resonance Energy Transfer (FRET)
- · 2) Fluorescence Cross-Correlation Spectroscopy (FCCS)

Why can't we image molecular interactions directly?

Microscope Resolution

• The diffraction of light causing the Airy disk is also the limiting factor of the resolution of a microscope

•The resolution of a microscope is defined as the minimum distance two objects have to be separated by to be resolved as two separate objects



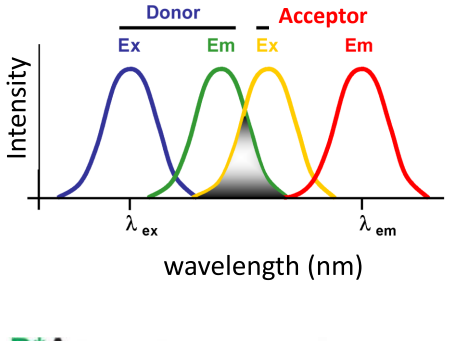
Airy patterns of two point sources

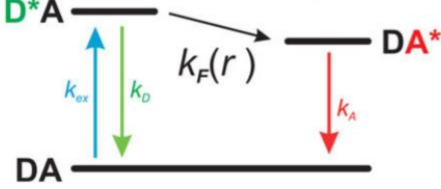
• For 100X magnification, 1.4 NA oil immersion objective and illumination with green light (500 nm = $0.5 \ \mu m$)

Resolution =
$$\frac{(0.61) (0.5 \,\mu\text{m})}{1.4} \approx 220 \,\text{nm}$$

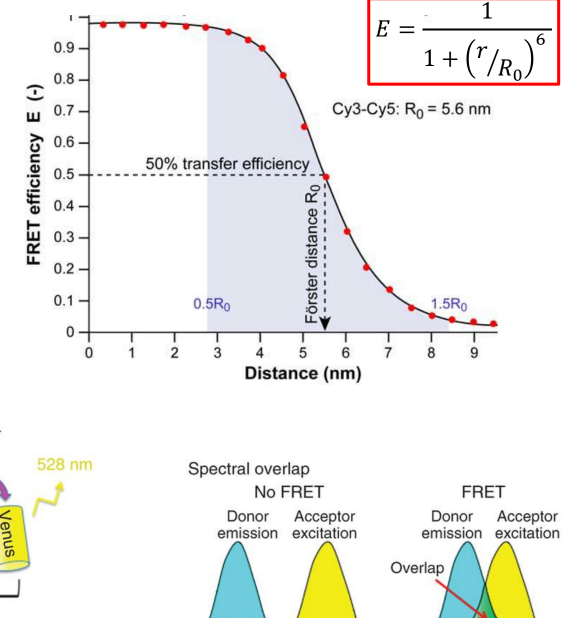
1) FRET – Förster Resonance Energy Transfer

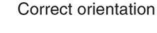
- energy is transferred non-radiatively (via long-range dipole-dipole coupling)
- relies on the close physical interaction of the two fluorophores (0.5 – 10nm)
- used to determine molecular interactions beyond the resolution limits

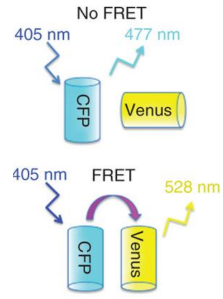


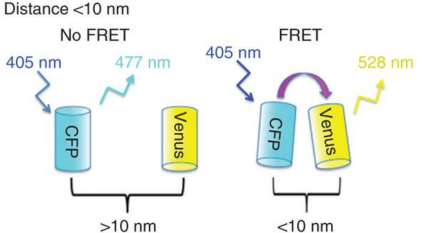


- distance between two molecules
- spectral overlap
- the quantum yield of the donor
- the relative orientation of the transition dipoles







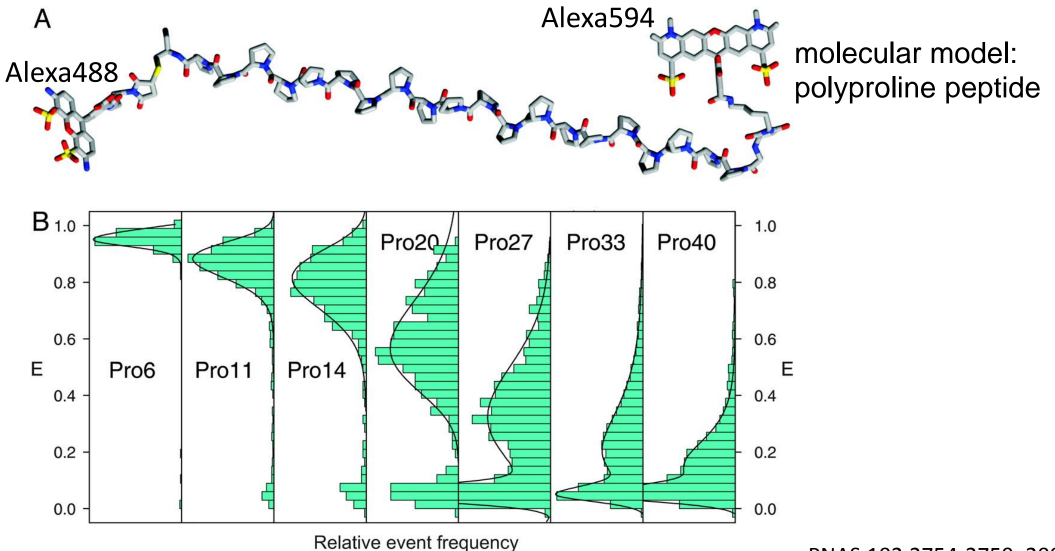


NatProt, 8, 265-281, 2013

FRET Pairs

		Donor	Acceptor	Donor	Acceptor	Förster	Cy5	Cy5.5	649	694	> 0.28	250,000	>8.0
Donor	Acceptor	Excitation	Emission	QY	3,	Distance	Fluorescein	Tetramethylrhodamine	487	574	0.93 ²	~ 87,000	4.9-5.5
<u> </u>		λ_{max} (nm)	λ_{max} (nm)	(-)	(M ⁻¹ cm ⁻¹)	(nm)	FITC	TRITC	494	572	0.92	100,000	5.4
Fluorescent protein FRET pairs							Phycoerythrin	APC ³	(546), 565	660	0.98	700,000	4.0-11.0
BFP	DsRFP	380	586	0.18	72,500	3.1–3.3	Europium	APC	340	660		700,000	9.0
EBFP2	mEGFP	383	507	0.56	57,500	4.8	Tryptophan	Dansyl	280	525	0.01-0.354	4050	2.1
CFP	GFP	433	509	0.40	21,000	4.7-4.9	Dansyl	FITC	335	519	< 0.035 ⁵	77,000	3.3-4.1
CFP	YFP	433	526	0.40	77,000	~ 5.0	Dansyl	Octadecylrhodamine	335	625	< 0.035	106,000	4.3
Cerulean	YFP	440	526	0.62	77,000		Europium	Cy5	340	670		250,000	7.0
ECFP	EYFP	440	527	0.40	83,400	4.9	Atto 488	Atto 647N	501	670	0.8	150,000	5.1
Cerulean	Venus	440	528	0.62	92,200	5.4	Atto 488	Atto 590	501	621	0.8	120,000	6.0
MiCy	mKO	472	559	0.90	51,600	5.3	Atto 550	Atto 647N	554	670	0.8	150,000	6.5
GFP	YFP	475	526	0.77	77,000	5.5-5.7	Atto 550	Atto 655	554	684	0.8	125,000	6.4
GFP	mRFP	475	579	0.77	50,000	~ 4.7	Atto 590	Atto 655	594	684	0.8	125,000	7.3
CyPet	YPet	477	530	0.51	104,000	5.1	Alexa 405	Alexa 430	401	541		16,000	
TFP1	mVenus	492	528	0.85	92,200	5.1	Alexa 488	Alexa 514	495	542	0.92	80,000	-
EGFP	mCherry	507	510	0.60	72,000	5.1	Alexa 488	Alexa 532	495	554	0.92	81,000	-
Venus	mCherry	528	610	0.57	72,000	5.7	Alexa 488	Alexa 546	495	573	0.92	104,000	6.4
Venus	tdTomato	528	581	0.57	138,000	5.9	Alexa 488	Alexa 610	495	628	0.92	138,000	
Venus	mPlum	528	649	0.57	41,000	5.2	Alexa 647	Alexa 680	650	702	0.33	184,000	-
Fluorescent protein-dye FRET pairs							Alexa 647	Alexa 700	650	723	0.33	192,000	-
EGFP	Alexa Fluor 555	484	568	0.6	155,000	6.3	Alexa 647	Alexa 750	650	780	0.33	240,000	-
EGFP	Alexa Fluor 546	484	573	0.6	112,000	5.7	Non-fluoresc	ent acceptor pairs					
EGFP	Alexa Fluor 594	484	618	0.6	92,000	5.3	Rhodamine 6G	Malachite Green	526	NF	0.95	76,000	6.1
EGFP	Alexa Fluor 568	484	603	0.6	88,000	5.4		(Abs 628 nm)					1000 B
Dye and dye-biofluorochrome FRET pairs							Alexa 488	QSY 35 (Abs 475 nm)	495	NF	0.92	23,000	4.4
Cy2	СуЗ	489	570	> 0.12	150,000	5.0-6.0	Alexa 488	Dabcyl (Abs 453 nm)	495	NF	0.92	32,000	4.9
Cy3	Cv5	550	670	> 0.15	250,000	>5.0	Alexa 647	QSY 21 (Abs 661 nm)	650	NF	0.33	90,000	6.9

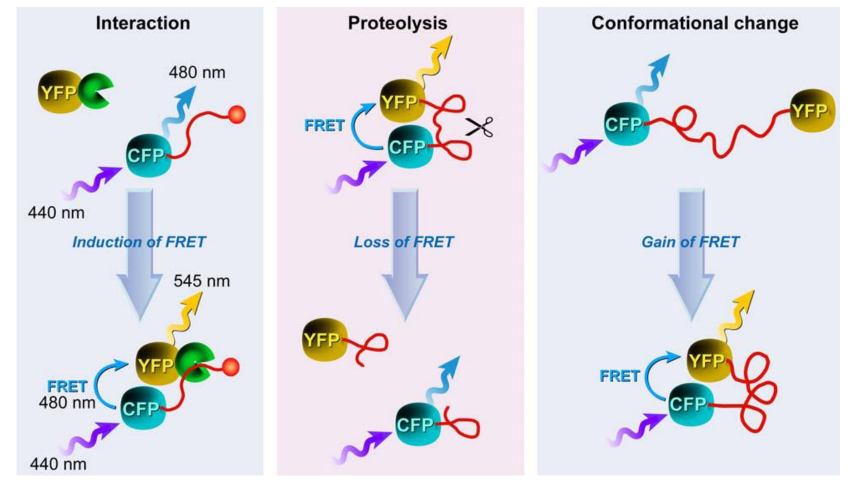
FRET – traditionally used as Molecular Ruler



PNAS,102,2754-2759, 2005

Fundamental cellular FRET-Approaches

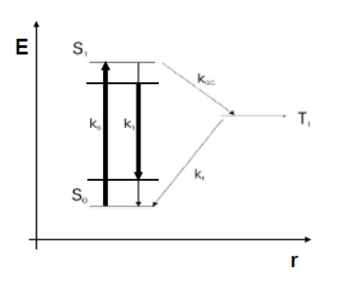
- FRET-biosensors: fusion proteins or other appropriate pairs linked by a sensory domain
- monitoring of donor/ acceptor channels and detection of changes in the FRET signal as a result of biological activity

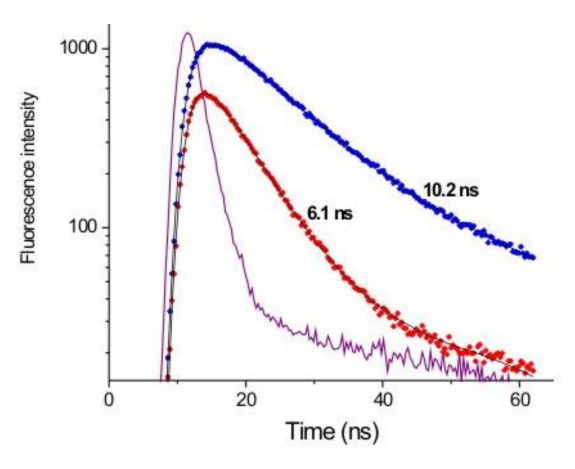


A variation of FRET: FRET-FLIM

FLIM – Fluorescence Lifetime Imaging

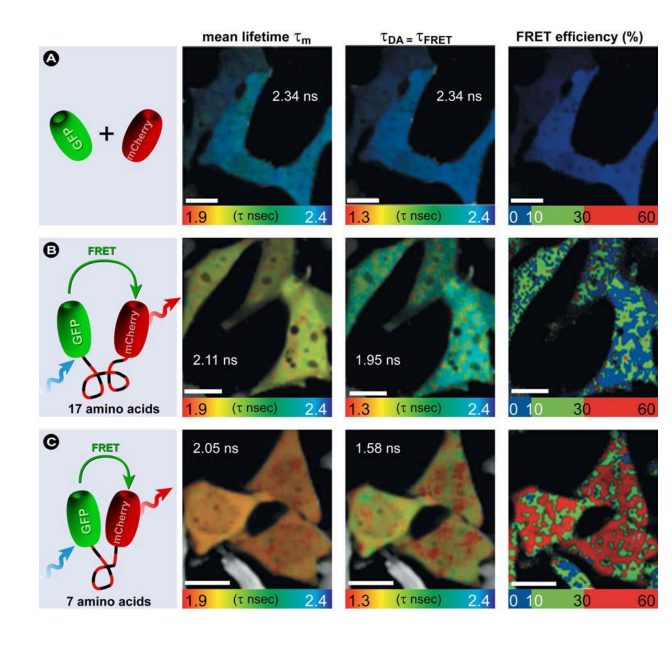
- ...is an imaging technique for producing an image based on the differences in the exponential decay rate of fluorophores.
- the lifetime of the fluorophore signal, not the intensity, is used to create the image in FLIM.





FRET-FLIM

- measurement of lifetime dynamics pixelby-pixel
- mapping of spatial distributions to measure biomolecule concentrations, interactions between biomolecules, and conformational changes with a much higher accuracy than conventional FRET methods



FLIM with TCSPC in practice

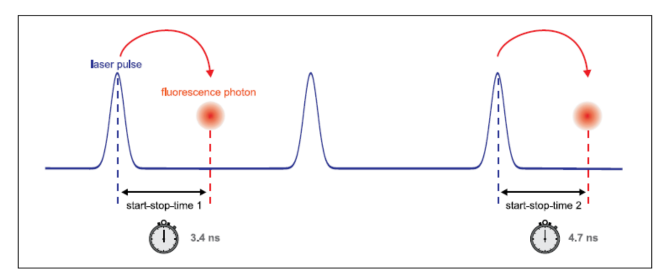


Figure 1: Measurement of start-stop times in time-resolved fluorescence measurement with TCSPC.

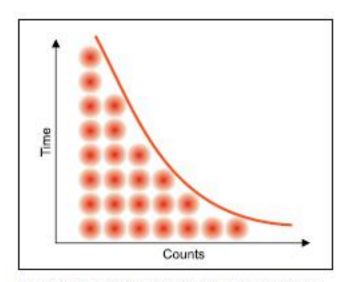
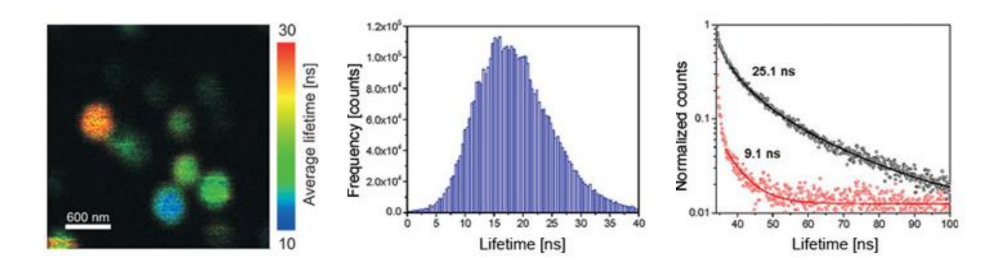


Figure 2: Histogram of start-stop times in time-resolved fluorescence measurement with TCSPC.

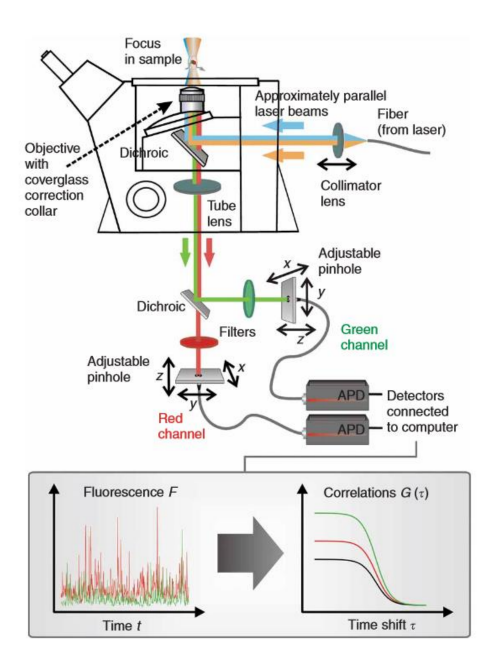
FLIM Measurements

- essential components of a FLIM set-up:
 - pulsed laser source
 - detector (APD or PMT)
 - TCSPC unit to measure the time between excitation and fluorescence emission (timedomain FLIM)
- the delay times are sorted into a histogram

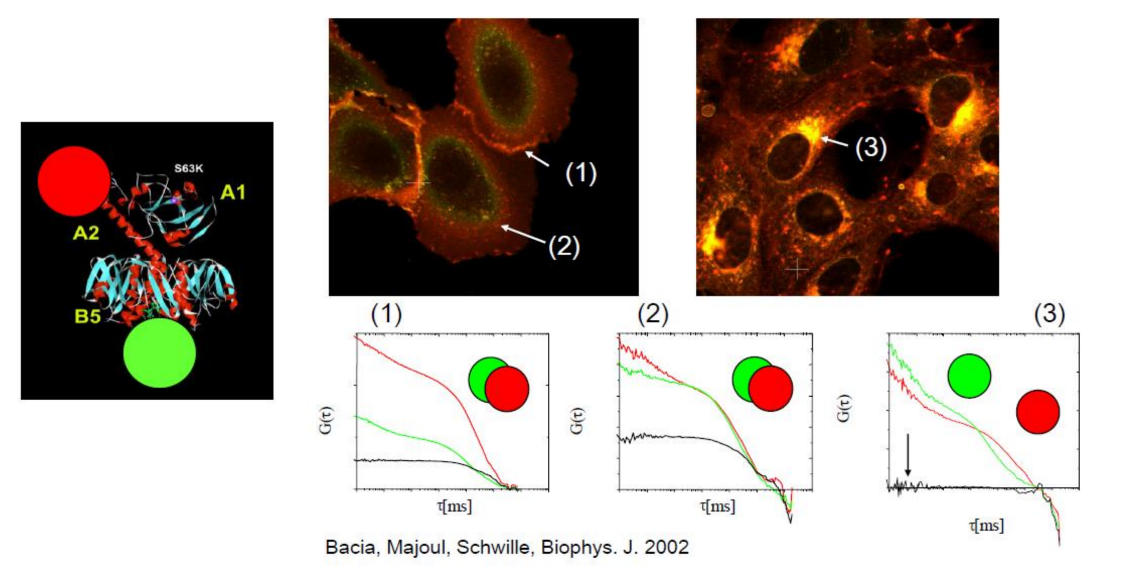


2) FCCS – Dual-Colour Extension of FCS

- interaction of two differently labeled molecular species with higher precision than single-color FCS
- high specificity: FCCS curve is only formed if the differently labeled molecules are bound and moving together (co-diffuse).
- From amplitude and decay time: binding constants, mobility of the bound complex, concentrations of all of the species



Toxicity of Choleratoxin unfolds as the active A-units separated from B₅ in Golgi



Summary - Fluorescence techniques for measuring molecular interactions

•FRET is a direct method for measuring molecular interactions beyond the resolution limits (~0.5-10 nm)

•FRET-FLIM is particularly powerful method because it minimizes spectral crosstalk problems

•FCCS is extension of FCS that enables the direct measurement of molecular interactions provided that the interacting molecules are mobile

Jablonski-Diagram

- molecules have electronic, vibrational and rotational energy levels
- absorption = electron moves to a higher excited state
- vibrational relaxation (Stokes-Shift), internal conversion, intersystem crossing
- return to the ground state: fluorescence, phosphorescence

