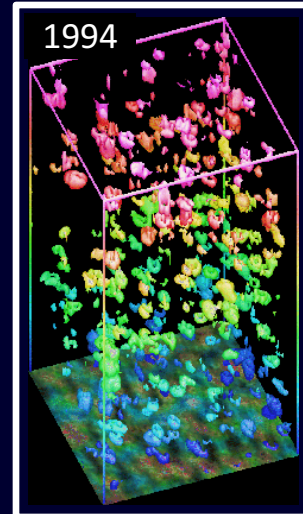
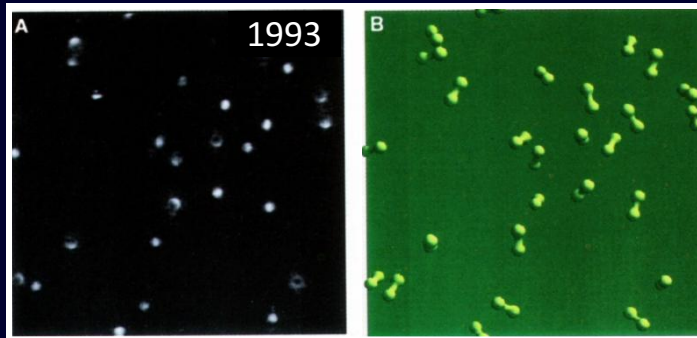
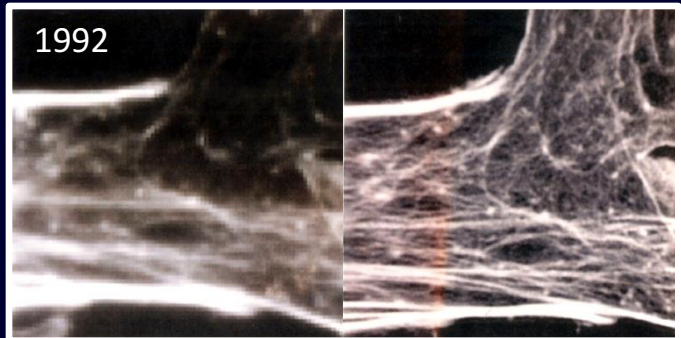
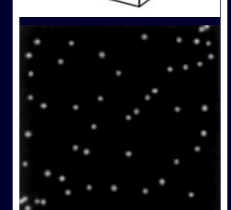
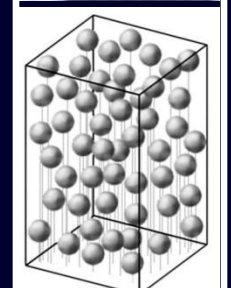
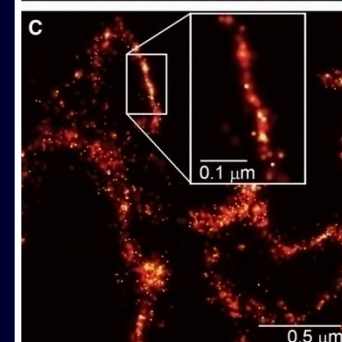
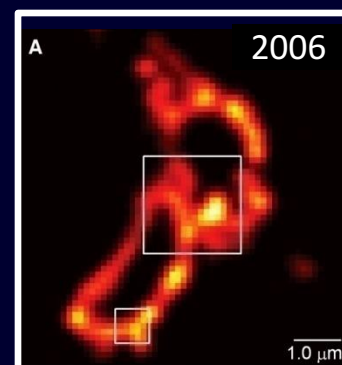
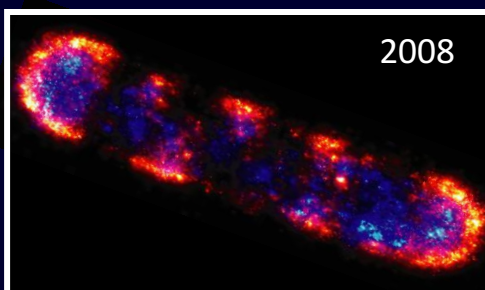
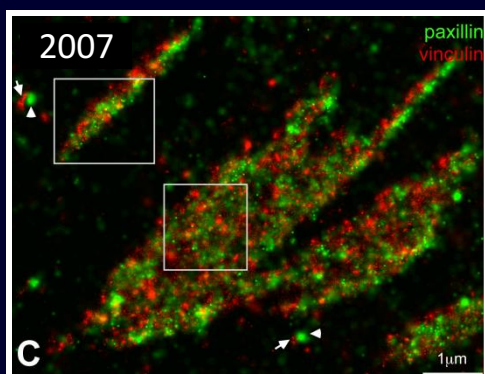
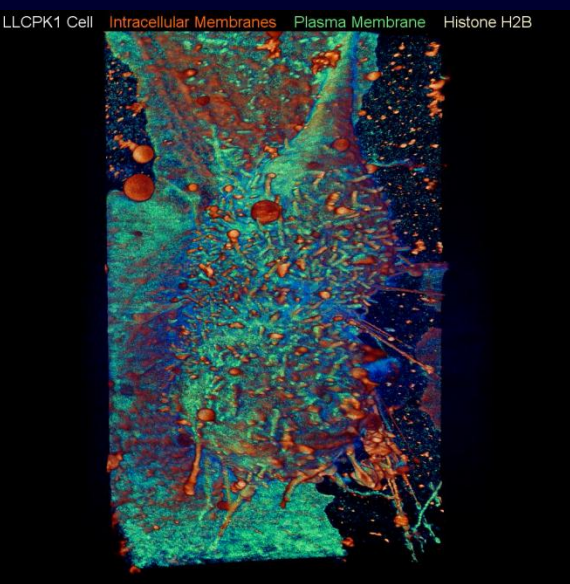


Single Molecules, Cells, and Super-Resolution Optics

Eric Betzig
Janelia Research Campus, HHMI



2014



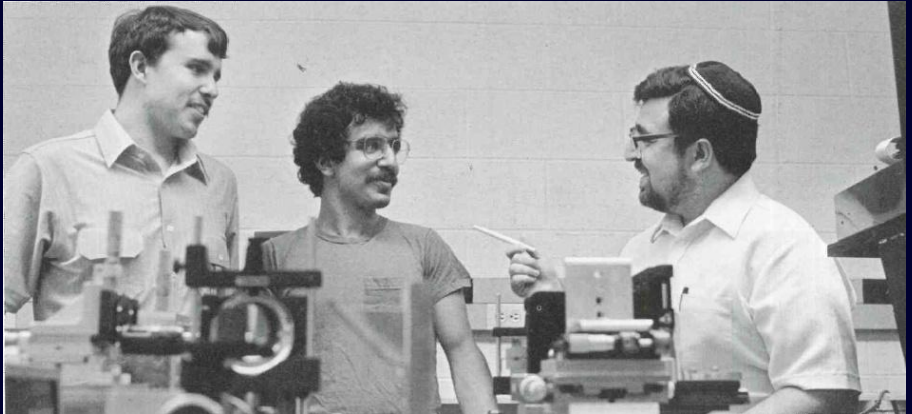
1
9
9
5

Cornell and the Beginnings of Near-Field Optical Microscopy

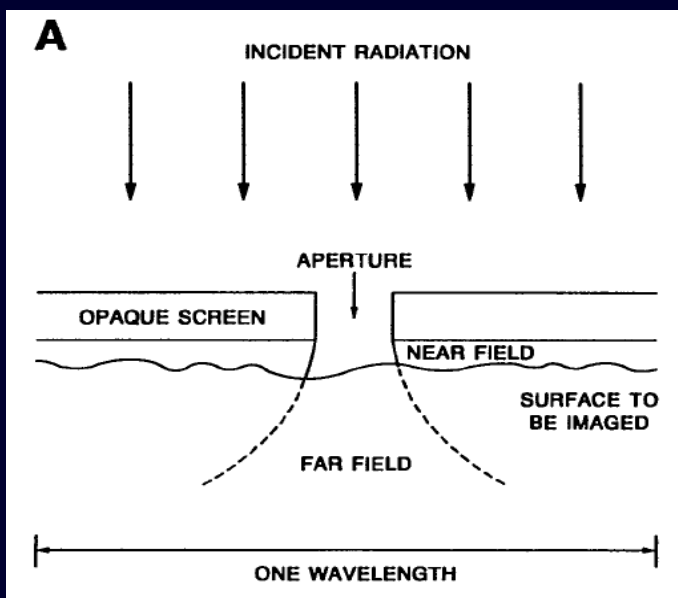
Mike Isaacson and his STEM



Me, Alec Harootunian, and Aaron Lewis, 1983



concept

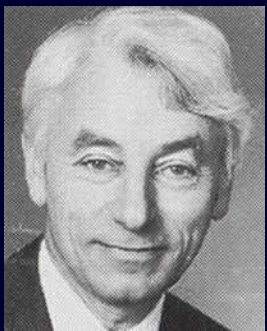


A. Lewis, et al.,
Ultramicroscopy **13**,
227 (1984)

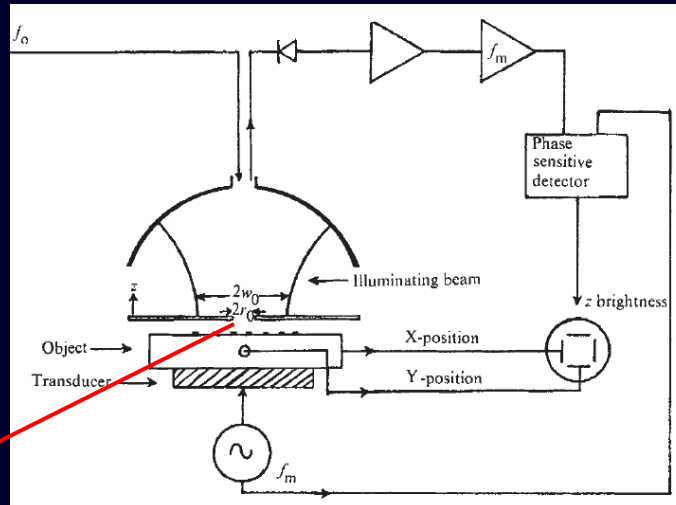
The Long History of Breaking Abbe's Law: Near-Field

near-field microwave ($\lambda = 3 \text{ cm}$) microscopy

Sir Eric Ash



sub-wavelength
aperture



object



image



Resolution of $1/60$ of the wavelength!

XXXVIII. A Suggested Method for extending Microscopic Resolution into the Ultra-Microscopic Region. By E. H. SYNGE*.



Edward "Hutchie"
Synge, *Phil. Mag.* **6**,
356 (1928)

- J.A. O'Keefe (1956)
- A.V. Baez (acoustics, 1956)
- C.W. McCutchen (1967)
- U. Ch. Fischer (lithography, 1981)
- D.W. Pohl (1984)
- G.A. Massey (1984)
- J. Wessel (1985)

The Long History of Breaking Abbe's Law: Far-Field

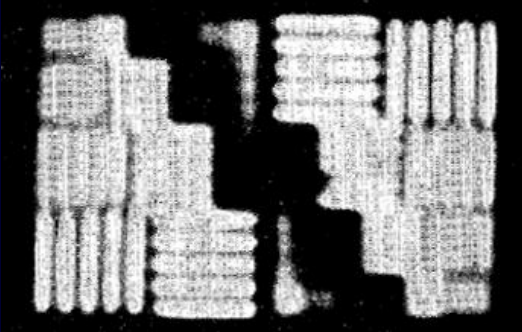
Structured Light

Optical Systems with Resolving Powers Exceeding the Classical Limit*

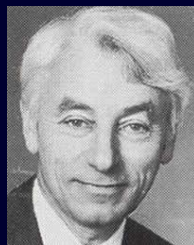
W. Lukosz†
Institut A für Physik, Technische Hochschule, 33 Braunschweig, Germany
(Received 27 April 1966)

W. Lukosz, JOSA **56**, 1463 (1966)

test pattern, conventional



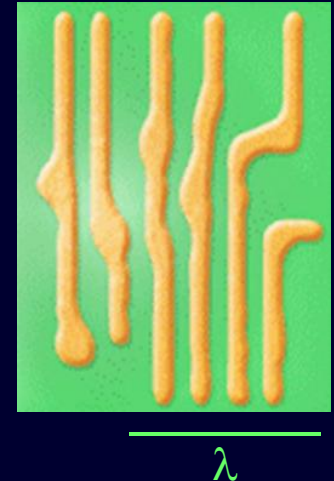
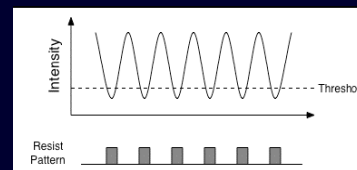
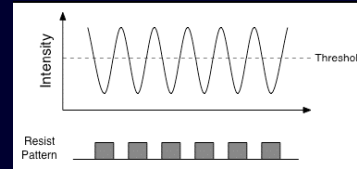
Sir Eric Ash



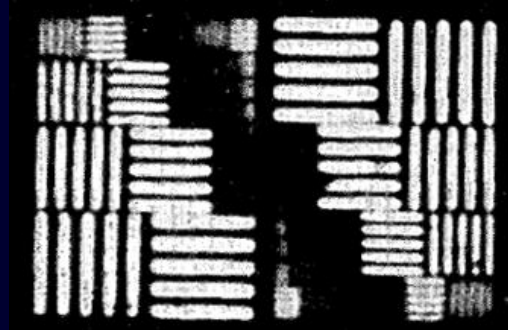
nominal exposure

Nonlinear Interaction with Sample

integrated circuit linewidth control



test pattern, super-resolved

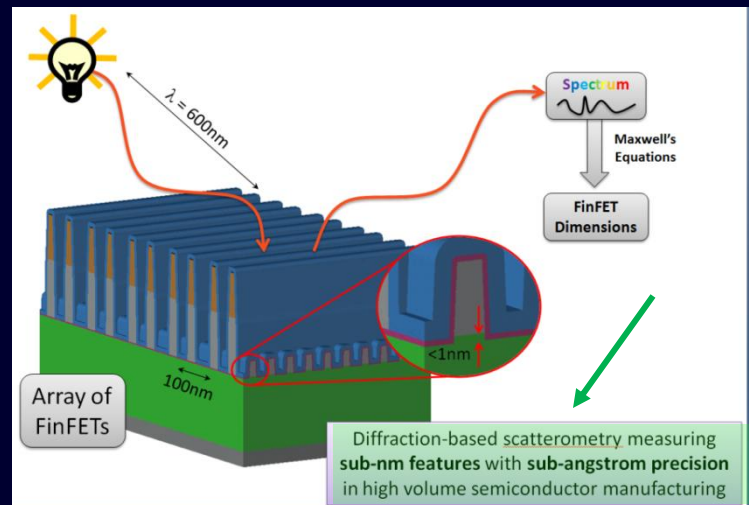


Resolution 3×
beyond Abbe's
Limit!

A. Bachl, W. Lukosz, JOSA **57**, 163 (1967)

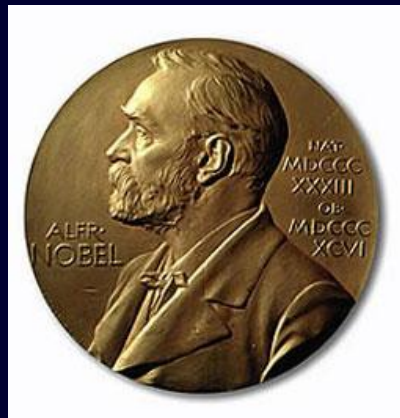
intentional
overexposure

A Priori Information: wafer inspection

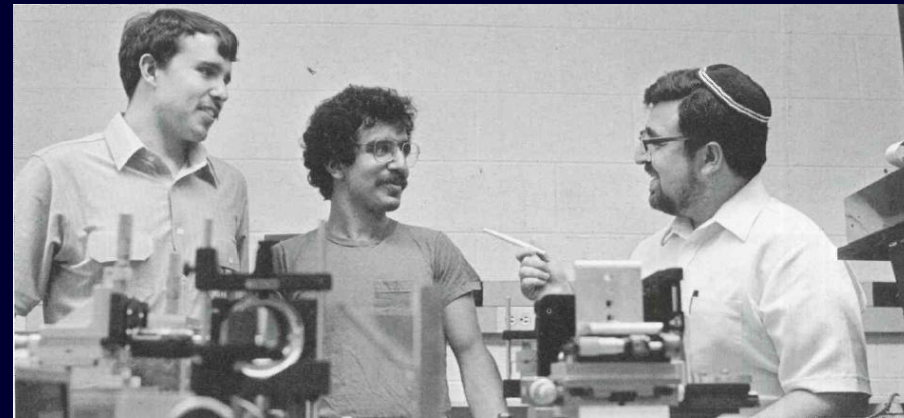


Making Near-field Optical Microscopy Work

Edwin Neher and Bert Sakmann, Nobel 1991



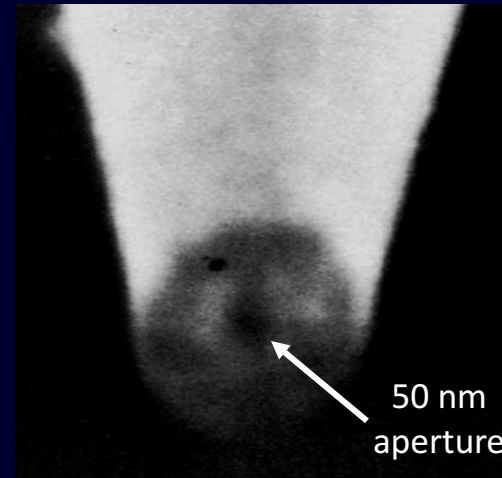
Me, Alec Harootunian, and Aaron Lewis, 1983



patch clamp: single ion channel recording



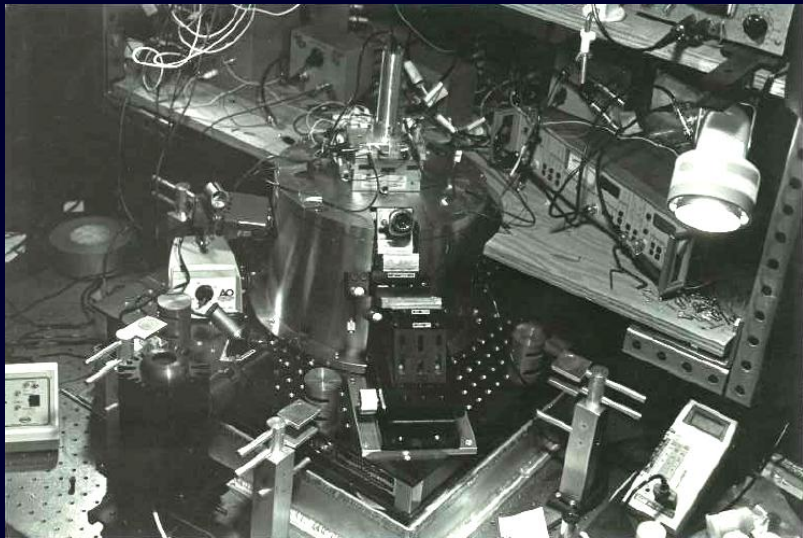
end of aluminum coated pipette



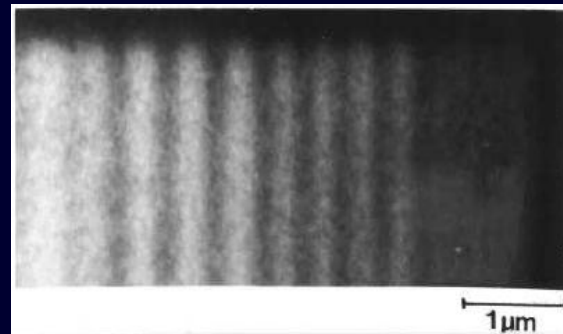
E. Betzig, et al., *Biophys. J.* **49**, 269 (1986)

Making Near-field *Optical* Microscopy Work

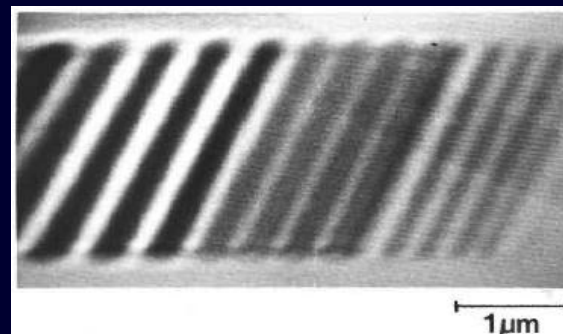
my near-field scanning optical microscope (NSOM)



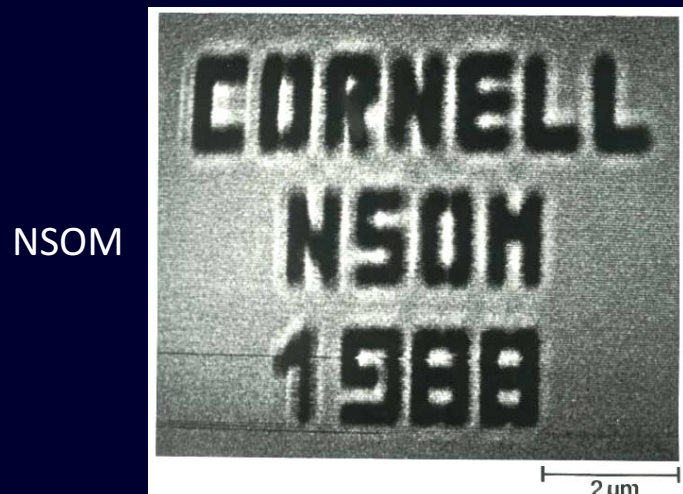
microscope control room



diffraction limited



NSOM

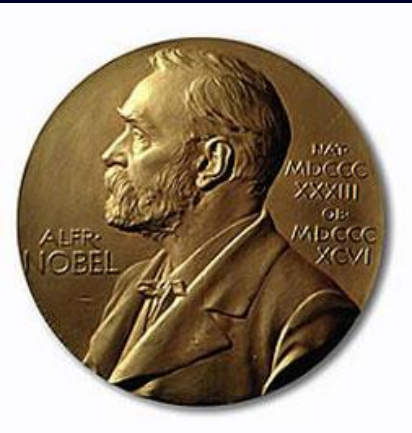
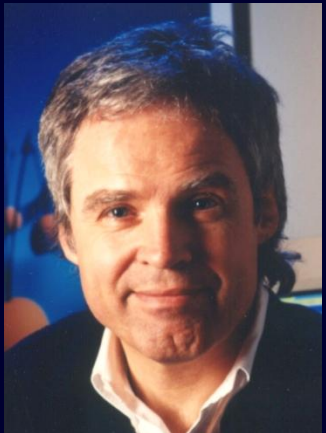


Initial Struggles at Bell Labs

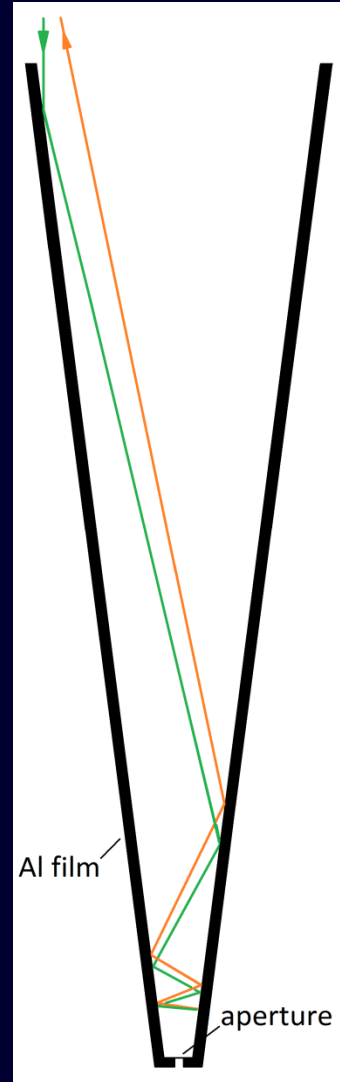
AT&T Bell Labs, Murray Hill, NJ



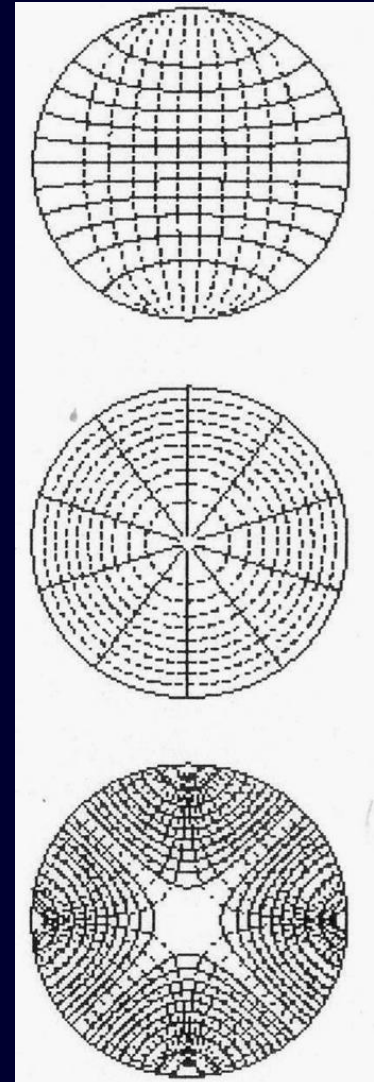
Horst Störmer, 1998 Nobel in Physics



retroreflection
in pipette



lowest order waveguide
modes at tip



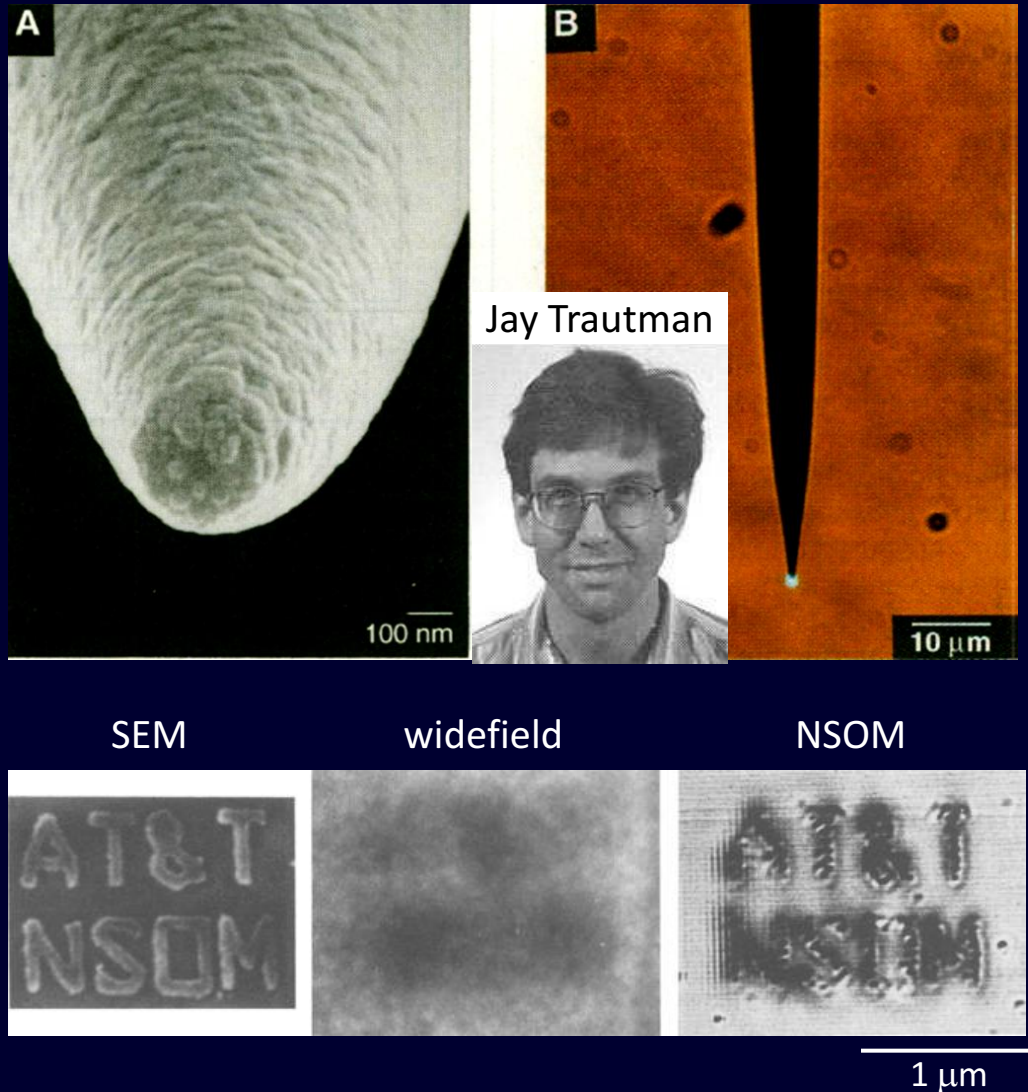
TE_{21}

TM_{01}

TE_{11}

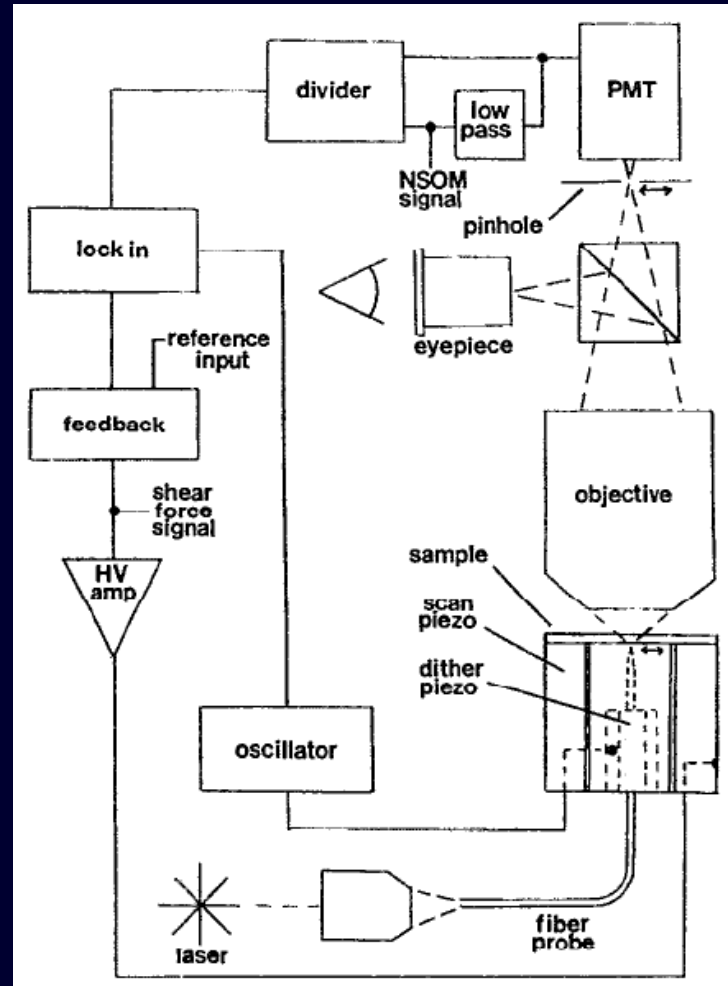
Making NSOM Routine

adiabatically tapered optical fiber probe



E. Betzig, J.K. Trautman, *et al.*, *Science* **251**, 1468 (1991)

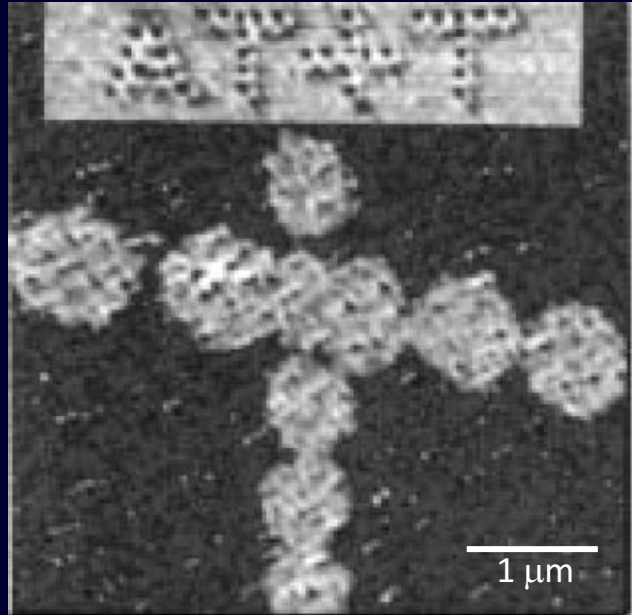
shear force distance regulation



E. Betzig, *et al.*, *Appl. Phys. Lett.*
60, 2484 (1992)

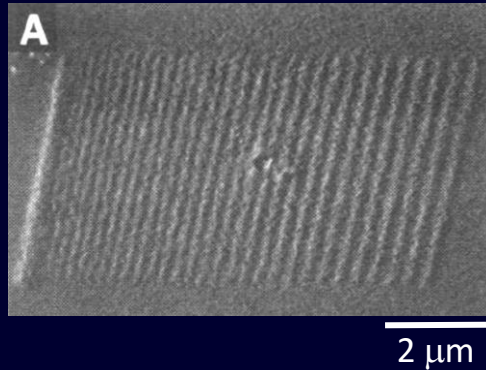
The Golden Age of NSOM

high density data storage

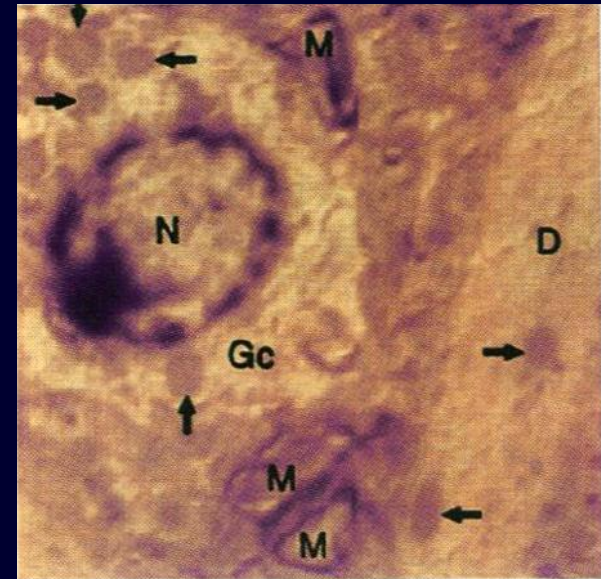


E. Betzig, et al., *Appl. Phys. Lett.* **61**, 142 (1992)

photolithography

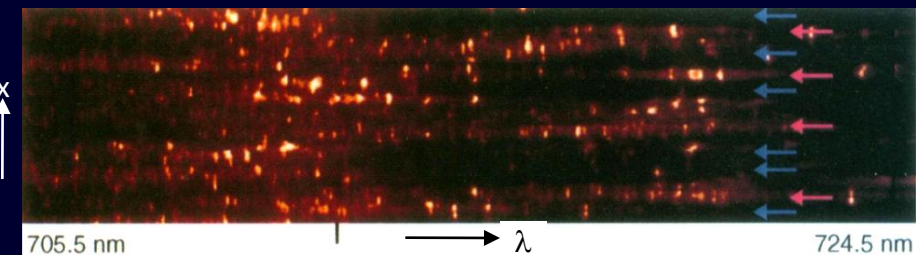


histological section,
monkey hippocampus



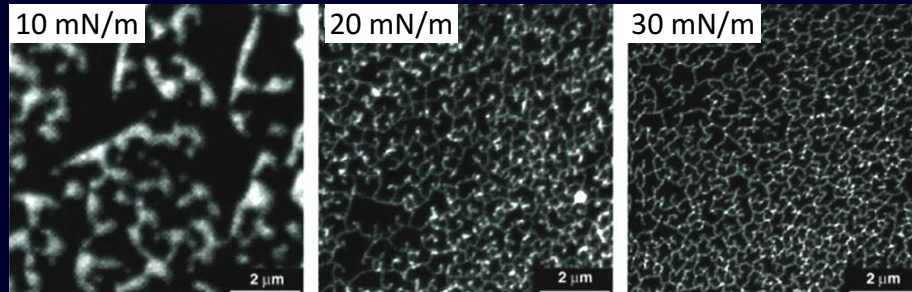
E. Betzig, J.K. Trautman, *Science* **257**, 189 (1992)

nanoscale spectroscopic imaging



H.F. Hess, et al., *Science* **61**, 142 (1994)

fluorescence: phase change in phospholipid monolayers



J. Hwang, et al., *Science* **270**, 610 (1995)

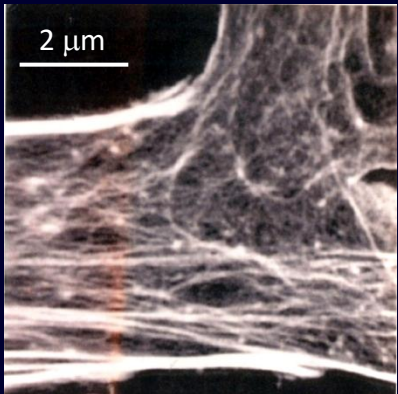
Single Molecule Detection (SMD)

fluorescence: actin, mouse fibroblast cell

widefield



NSOM



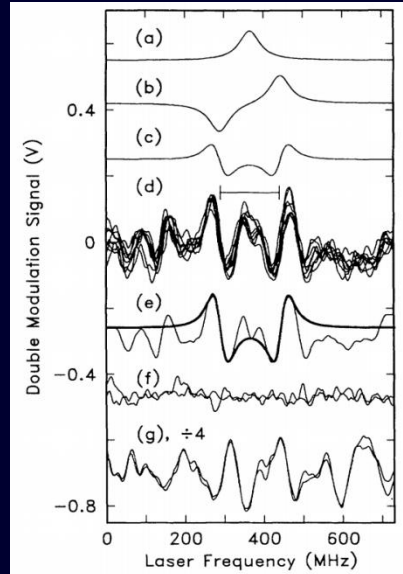
E. Betzig, et al., *Bioimaging* **1**, 129 (1993)

Nobel, 2014



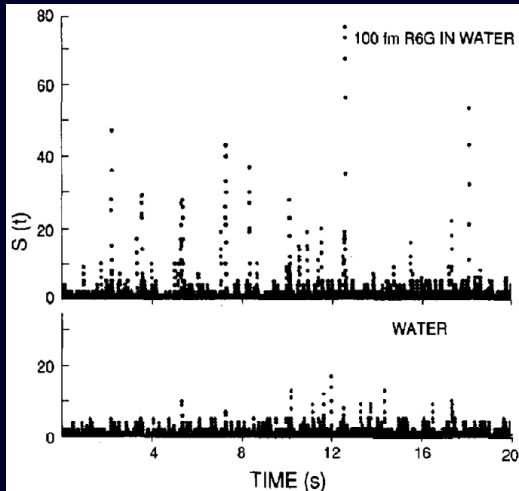
W.E. Moerner

W.E. Moerner, L. Kador,
Phys. Rev. Lett. **62**, 2535
(1989)



SM fluorescence excitation spectrum, 1.8°K

SM fluorescence bursts at room temp



Time gated:

E.B. Shera, et al., *Chem. Phys. Lett.* **174**, 553 (1990)

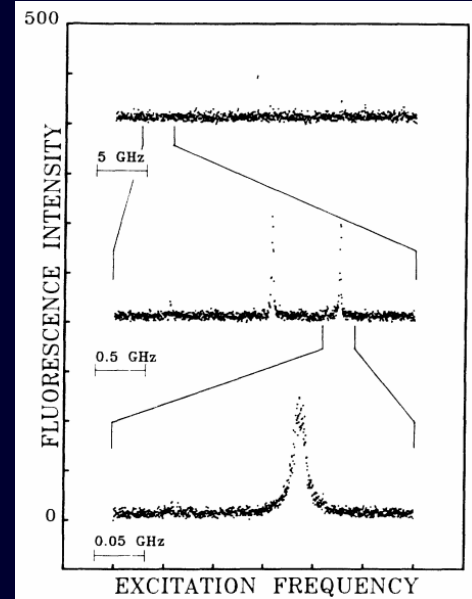
FCS:

R. Rigler, J. Widengren,
Bioscience **3**, 180
(1990)

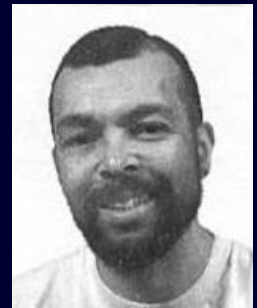


Michel Orrit

M. Orrit, J. Bernard,
Phys. Rev. Lett. **65**,
2716 (1990)

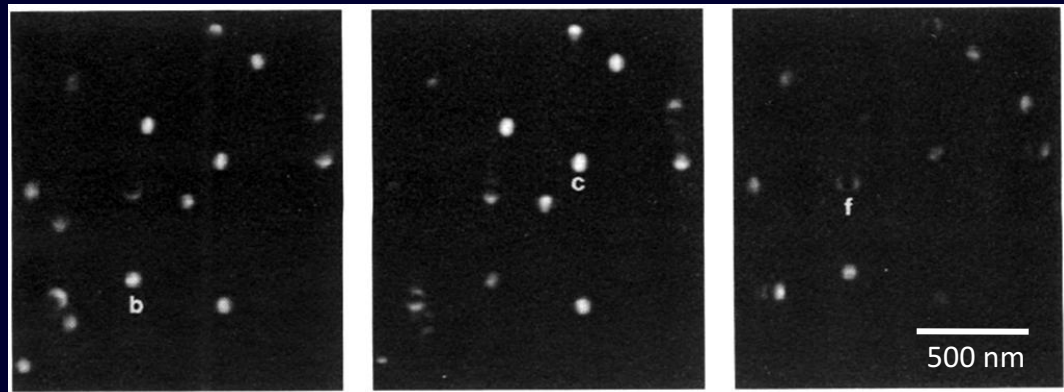


NSOM and the Birth of Single Molecule Microscopy



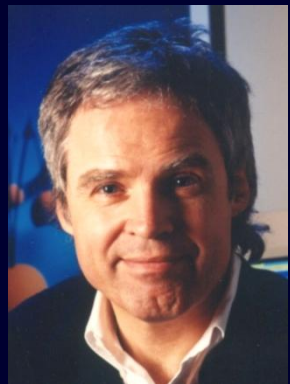
single molecule fluorescence anisotropy

random

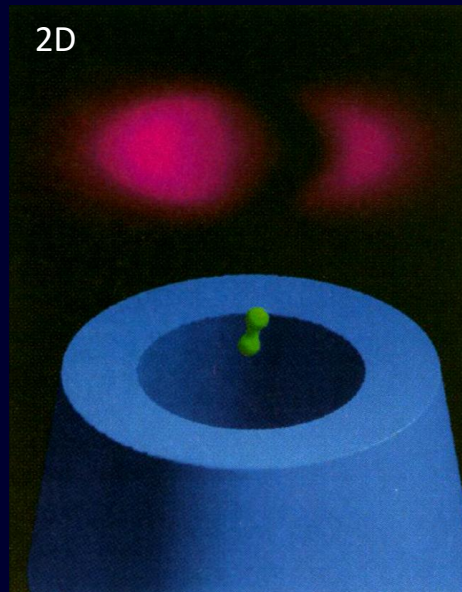
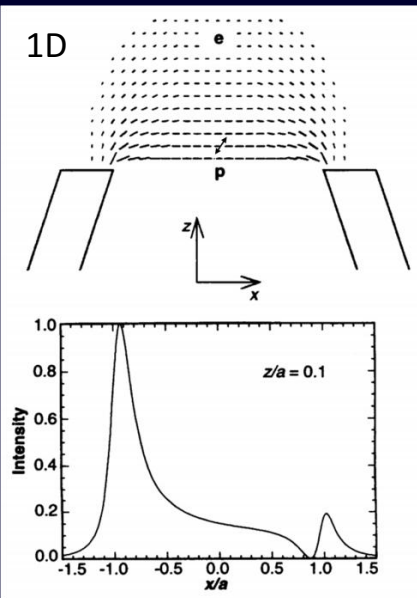


dil-C₁₈-(3)
molecules on
PMMA

E. Betzig, R.J. Chichester,
Science **262**, 1422 (1993)

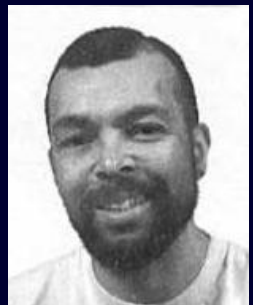


Horst Störmer



single molecule NSOM signal
 $|\mathbf{E}(\mathbf{x}) \cdot \mathbf{p}|^2$

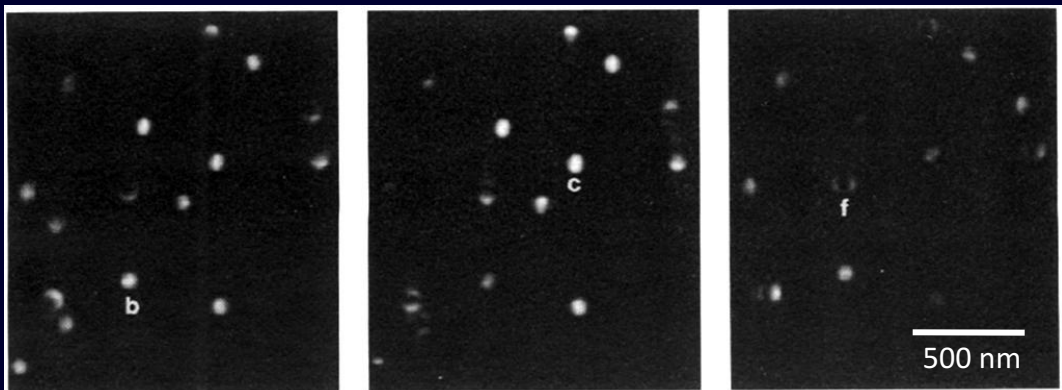
NSOM and the Birth of Single Molecule Microscopy



Rob Chichester

single molecule fluorescence anisotropy

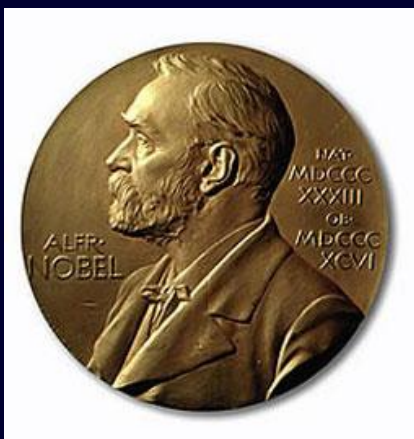
random



dil-C₁₈-(3)
molecules on
PMMA

E. Betzig, R.J. Chichester,
Science **262**, 1422 (1993)

Hans Bethe, 1967 Nobel in Physics



H.A. Bethe, *Phys. Rev.* **66**, 163 (1944)

E fields at aperture: theory vs. experiment

$z/a = 0.1$

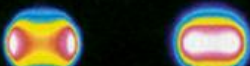
$z/a = 0.2$

data

$z/a = 0.4$

$z/a = 0.8$

$$|E_x|^2$$



$$|E_y|^2$$



$$|E_z|^2$$



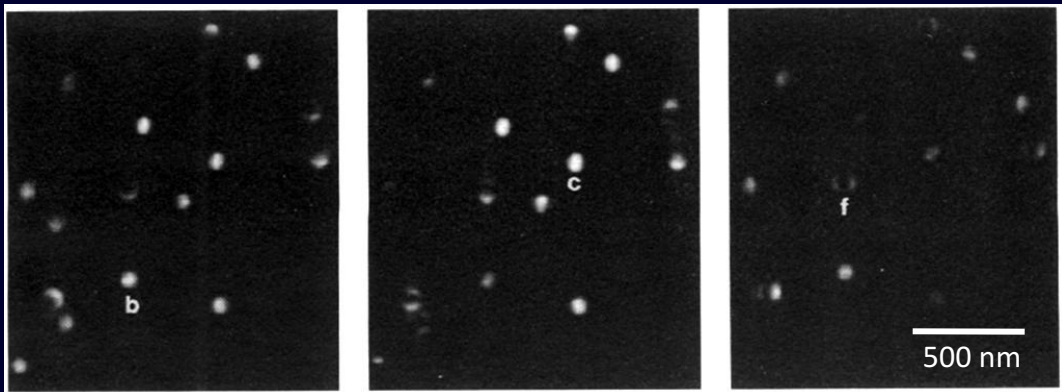
200 nm

NSOM and the Birth of Single Molecule Microscopy



single molecule fluorescence anisotropy

random

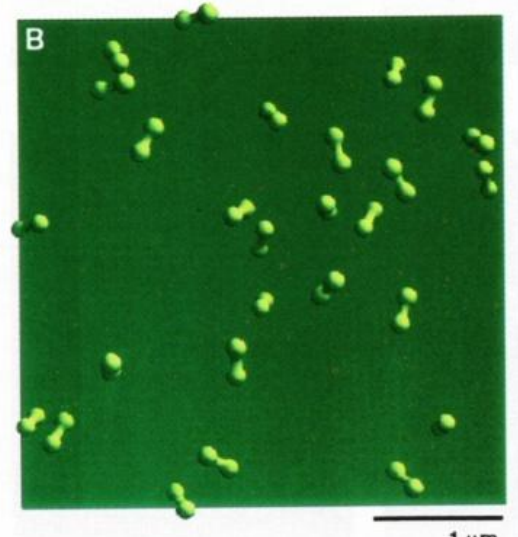
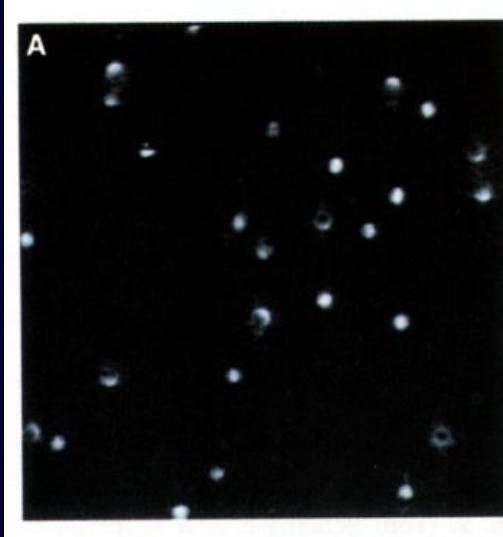


dil-C₁₈-(3)
molecules on
PMMA

E. Betzig, R.J. Chichester,
Science **262**, 1422 (1993)

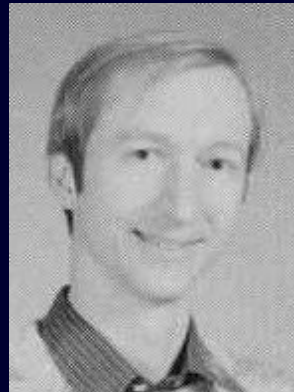
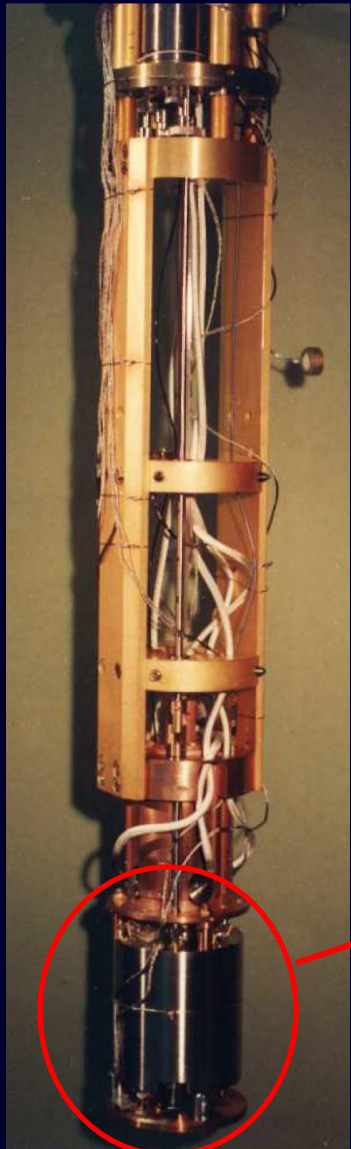
- first imaging of single molecules at room temp
- first super-resolution imaging of single molecules
- first measurement of single molecule dipole orientations
- first localization of single molecules to fraction of PSF width (12 nm xy, 6 nm z)

single molecule dipole orientations



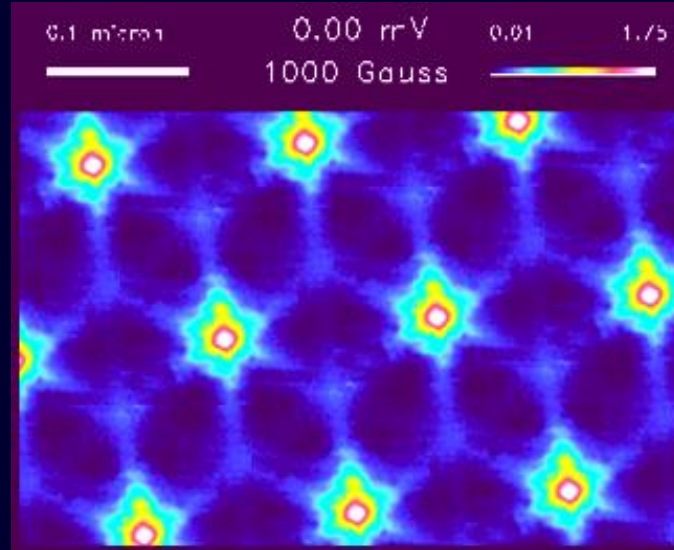
Cryogenic Near-field Spectroscopy

scanning tunnel spectroscopy of
Abrikosov flux lattice in NbSe_2



Harald Hess

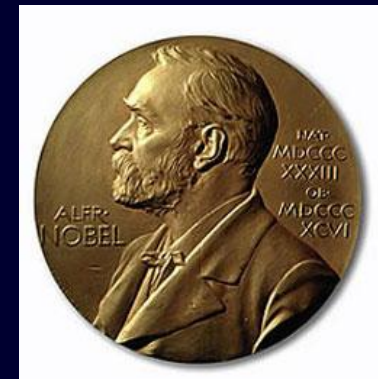
Harald's low temp STM



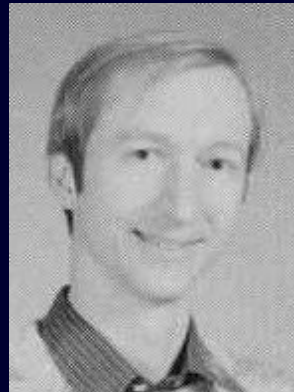
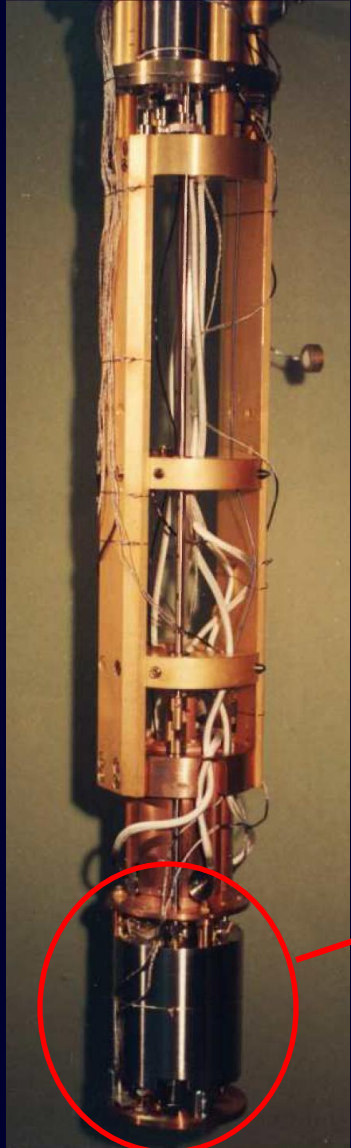
H.F. Hess, et al., *Phys. Rev. Lett.* **62**, 1691 (1989)



Alexei Abrikosov, 2003 Nobel in Physics

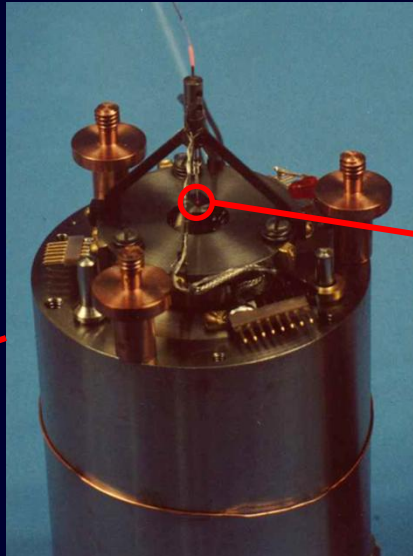


Cryogenic Near-field Spectroscopy

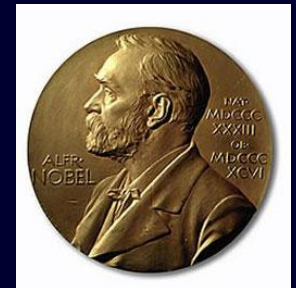
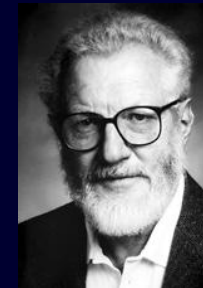


Harald Hess

Harald's low temp STM

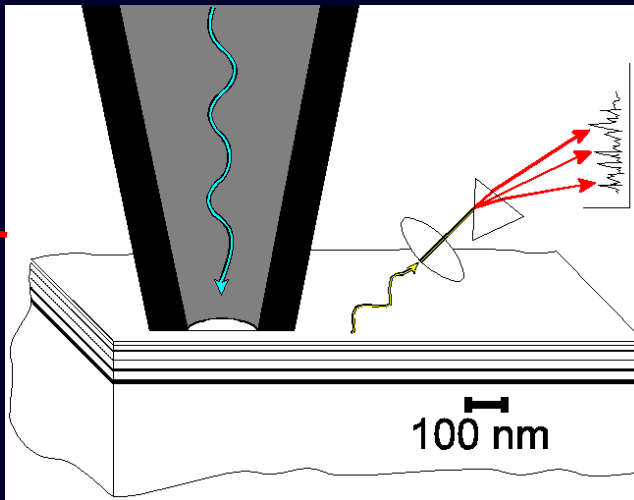


Alferov & Kroemer, 2000 Nobel in Physics



semiconductor
laser diode

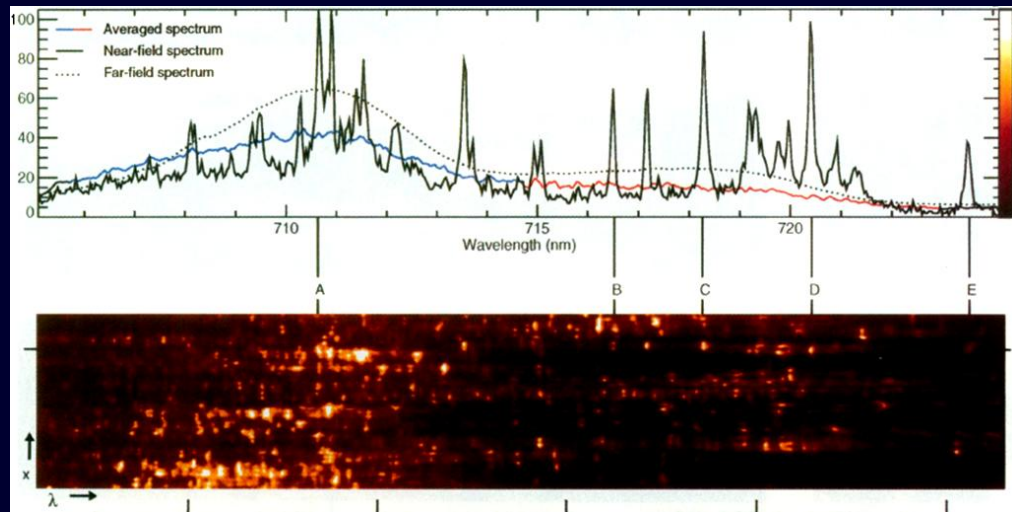
NSOM fiber probe



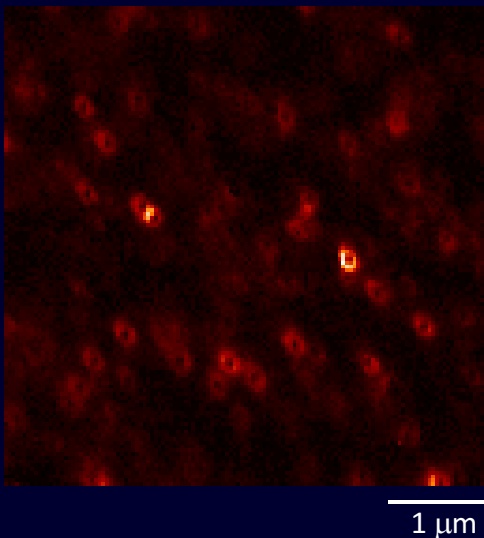
GaAs / AlGaAs
multiple
quantum well

Cryogenic Near-field Spectroscopy

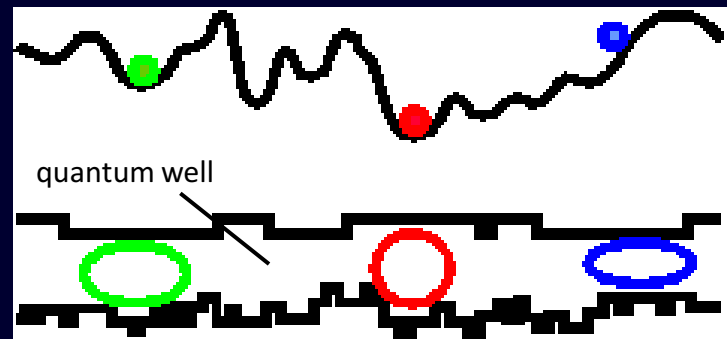
single exciton transitions, 23Å quantum well, 2°K



exciton recombination sites
scrolling from $\lambda = 700$ to $\lambda = 730$ nm



exciton energy variations due to
interface roughness



isolation of discrete
sites in x,y,λ space



H.F. Hess, E. Betzig, *et al.*,
Science **264**, 1740 (1994)

My First Mid-Life Crisis

NSOM engineering limitations:

- poor yield during manufacture
- fragile probes
- topographical artifacts
- weak signals
- probe tips get hot
- large probe tip ($0.25 \mu\text{m}$)

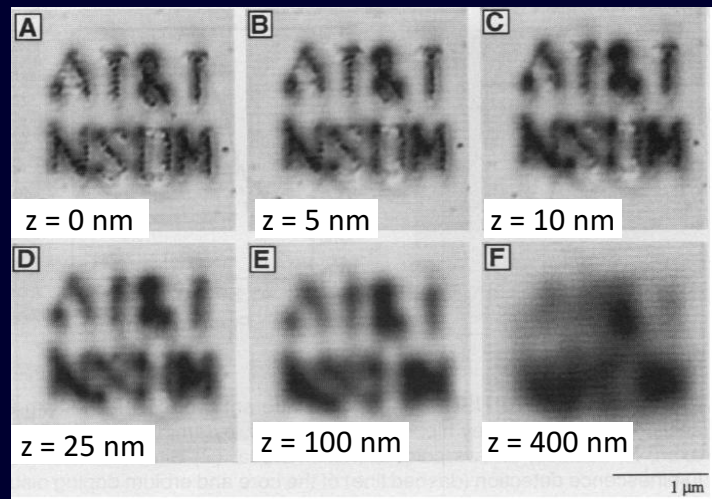
Cells aren't flat!



3D lattice light sheet microscopy,
D. Mullins, T. Ferrin, E. Betzig, *et al.*

NSOM fundamental limitations:

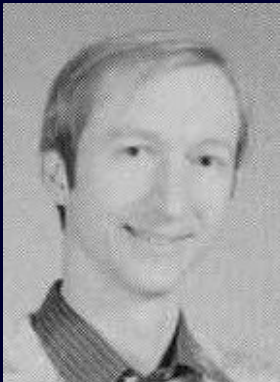
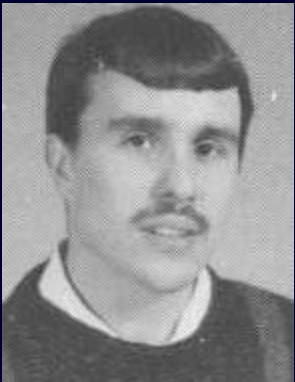
- probe perturbs fields at sample
- complex contrast mechanisms
- nonlinear image formation - artifacts
- the near-field is VERY, VERY short



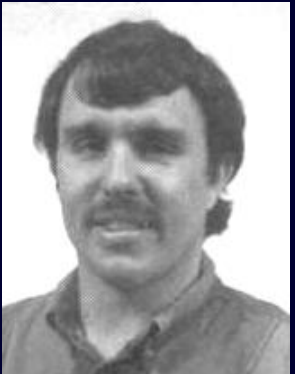
E. Betzig, J.K. Trautman, *Science* **257**, 189 (1992)

My First Mid-Life Crisis

me and Harald, 1989

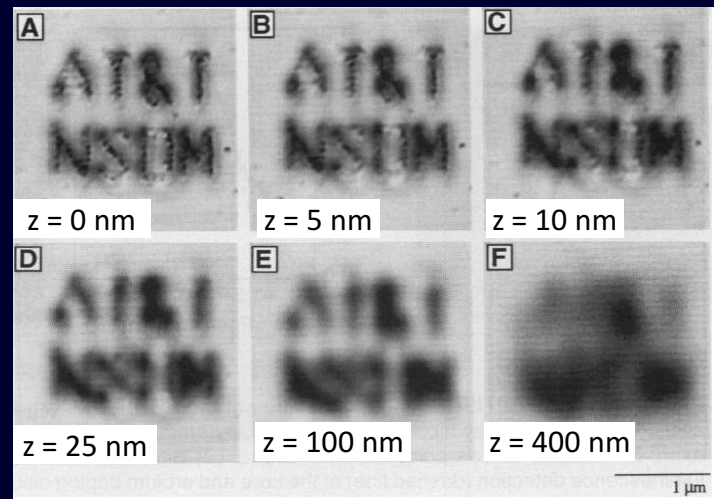


me and Harald, 1994



NSOM fundamental limitations:

- probe perturbs fields at sample
- complex contrast mechanisms
- nonlinear image formation - artifacts
- the near-field is very, very short



E. Betzig, J.K. Trautman, *Science* **257**, 189 (1992)

Multidimensional Localization Microscopy

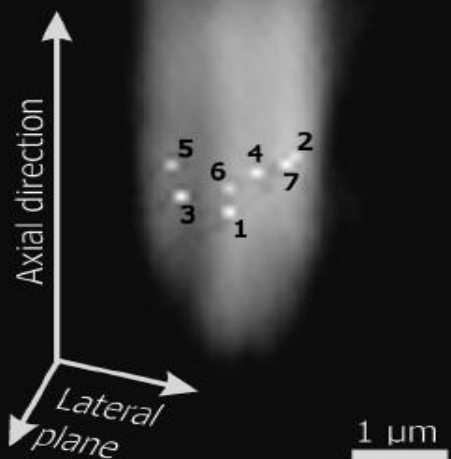
February 1, 1995 / Vol. 20, No. 3 / OPTICS LETTERS 237

Proposed method for molecular optical imaging

E. Betzig

NSOM Enterprises, 17 Webster Drive, Berkeley Heights, New Jersey 07922

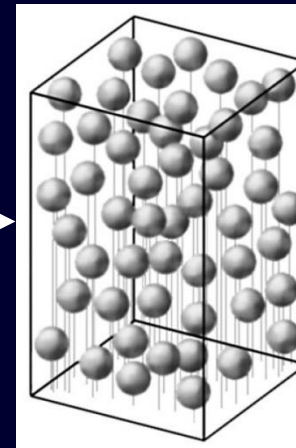
spectral isolation



A.M. van Oijen, et al., JOSA A16, 909 (1999)

higher
dimensional
isolation

original image



localization



Photobleaching: X. Qu, et al., PNAS 101, 11298 (2004)
M.P. Gordo, et al., PNAS 101, 6462 (2004)

Lifetime: M. Heilemann, et al., Anal. Chem. 74, 3511 (2002)

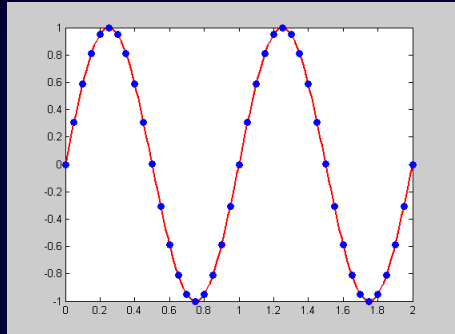
Blinking: K.A. Lidke, et al., Opt. Express 13, 7052 (2005)

Spatial Resolution and the Nyquist Criterion

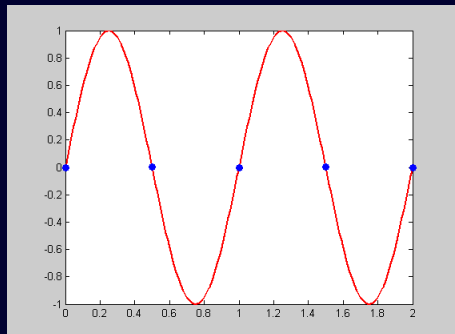
Nyquist criterion:

Sampling interval must be at least twice as fine as the desired resolution

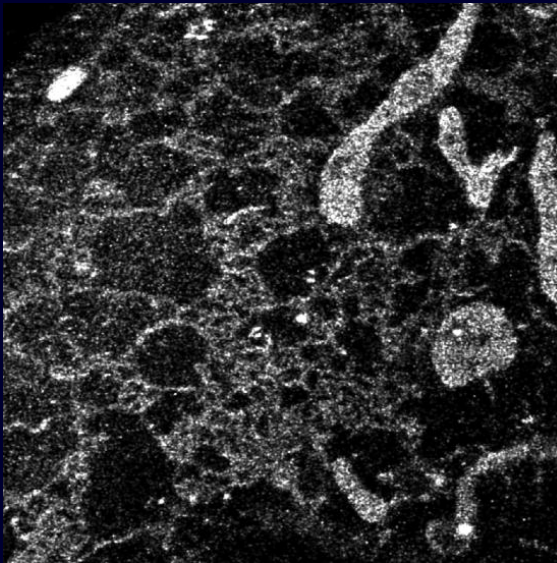
20 samples / period



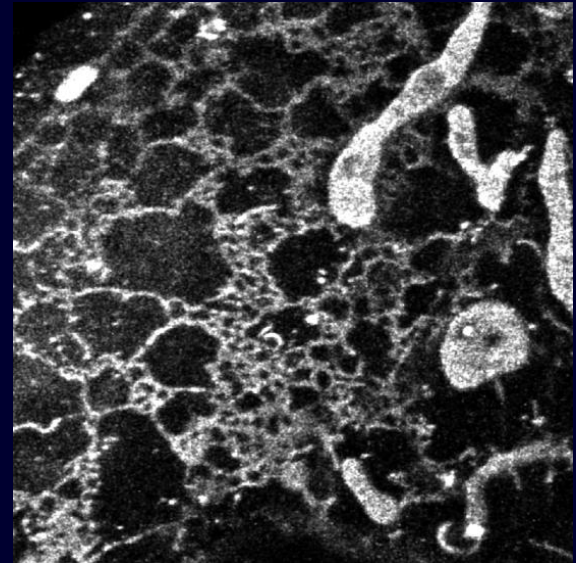
2 samples / period



initial molecular density



4x greater molecular density



2 μm

Image Dimensionality	Molecules Required per Diffraction Limited Region for 20 nm Resolution
----------------------	--

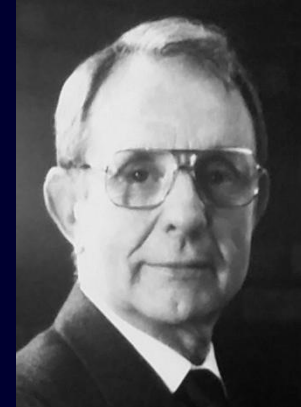
1D	25
2D	500
3D	2.9×10^4

Diffraction Limited Region:
0.25 μm dia, 0.6 μm long

And Now for Something Completely Different



Flexible Adaptive Servohydraulic Technology (FAST)



Robert Betzig

- moves 4000 kg load at 8g acceleration
- positioning precision to 5 μm



My Second Mid-Life Crisis

Searching for a New Direction

me in Joshua Tree National Park



Harald in Sedona, Arizona



me in Oahu, Hawaii

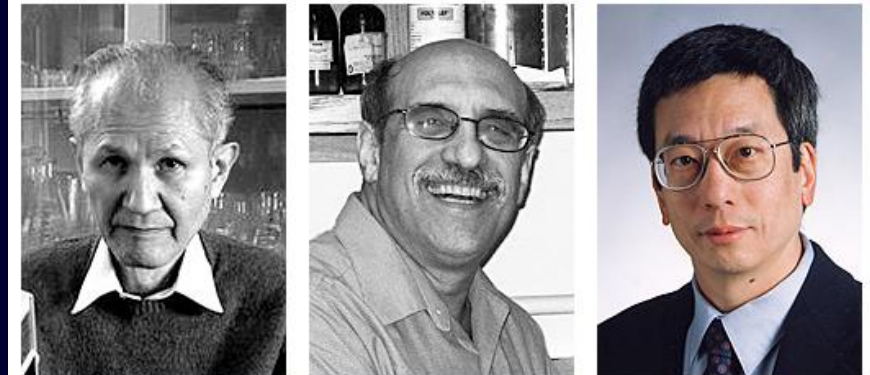


Harald in Yosemite National Park

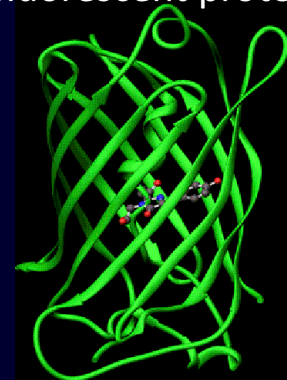


Fluorescent Proteins Revolutionize Biological Imaging

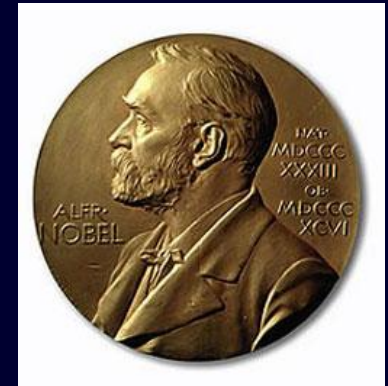
Shimomura, Chalfie, & Tsien



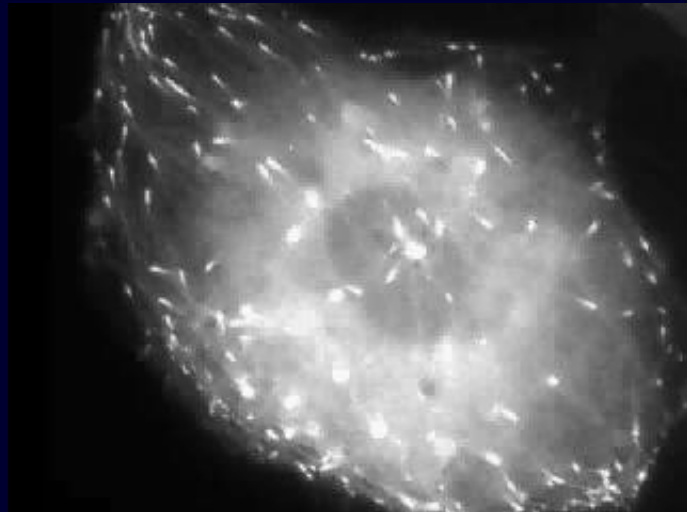
1994: green
fluorescent protein



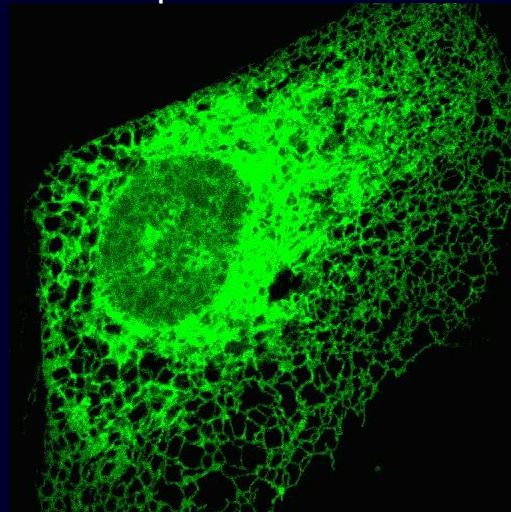
2008: Chemistry Nobel



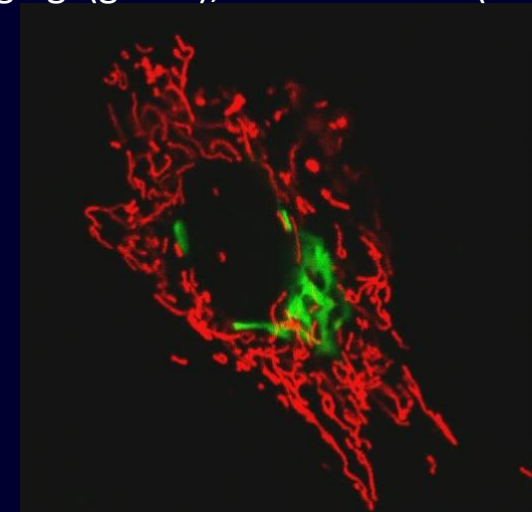
microtubule ends



endoplasmic reticulum

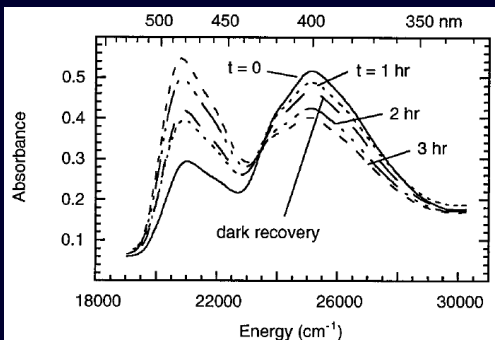


golgi (green), mitochondria (red)

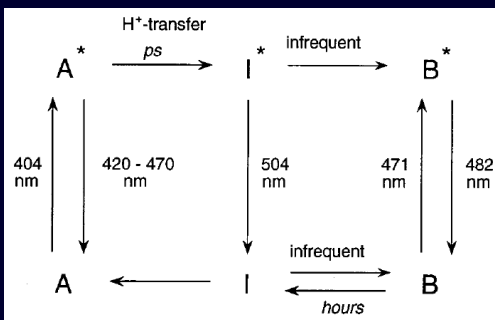


Switching Behavior in Green Fluorescent Protein

488 nm absorption increase under 398 nm illumination

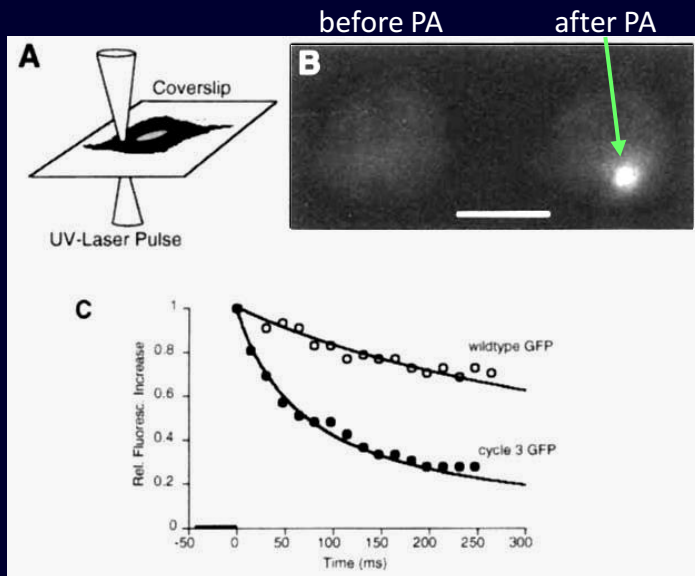


proposed mechanism



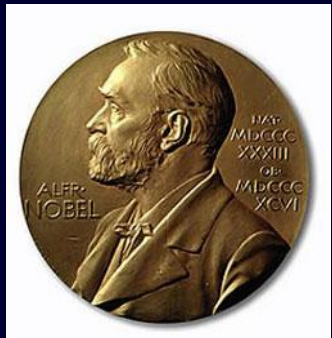
M. Chattoraj, et al., PNAS
93, 8362 (1996)

in vivo UV photoactivation (PA) of wtGFP



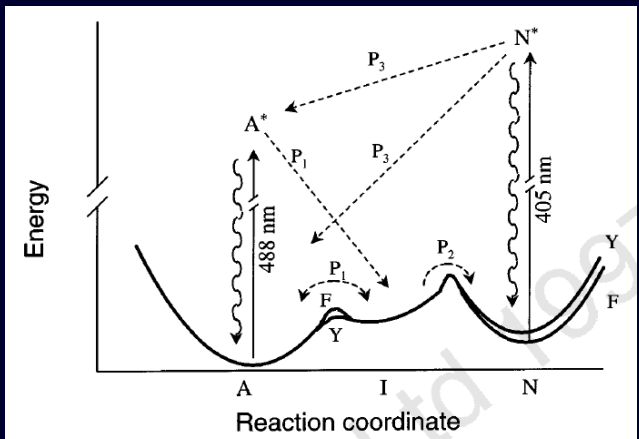
H. Yokoe, T. Meyer, Nat. Biotech. 14, 909 (1996)

W.E. Moerner, 2014 Nobel in Chemistry



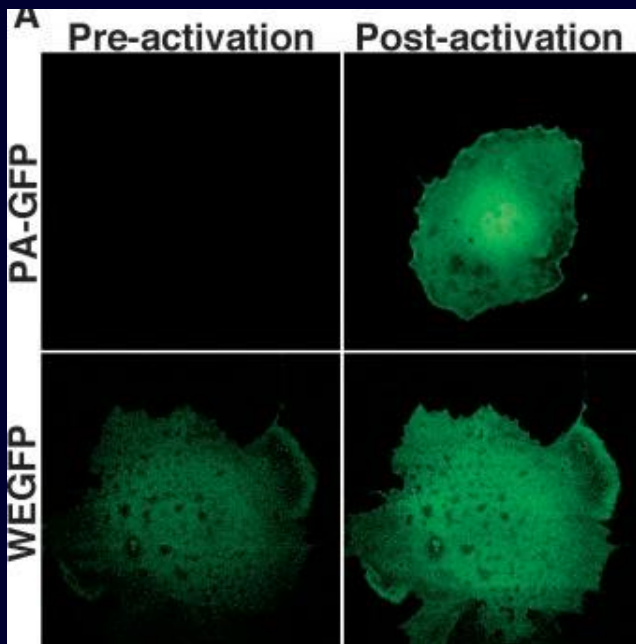
photoactivation energy diagram

R.M. Dickson, et al.,
Nature 388, 355 (1997)



Directed Mutagenesis of Photoactivated Fluorescent Proteins (PA-FPs)

increased on/off contrast of PA-GFP

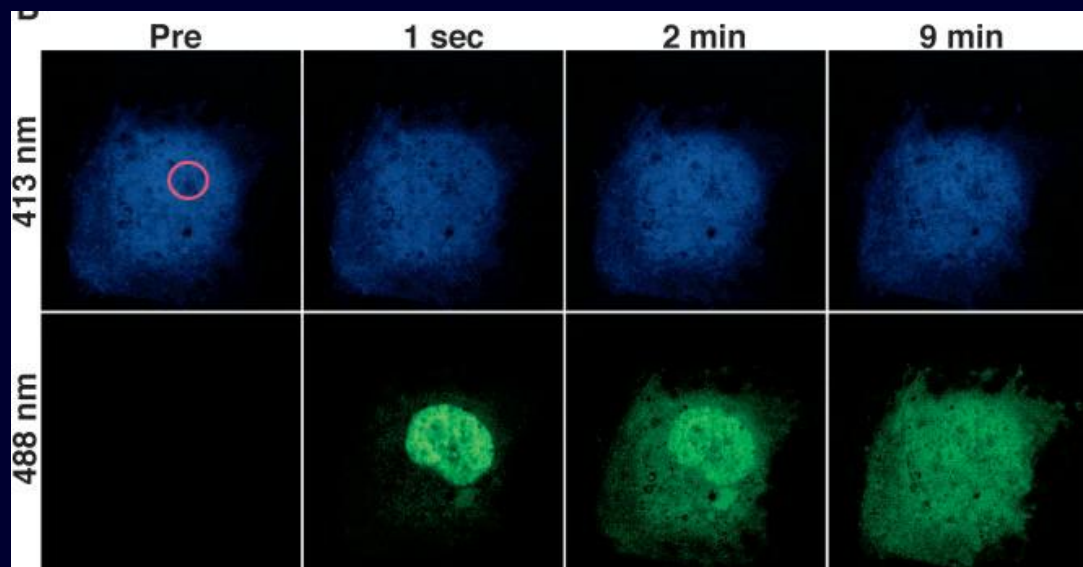


Jennifer
Lippincott-
Schwartz



George
Patterson

pulse chase: nuclear vs cytosolic diffusion



A Fateful Trip

Greg Boebinger



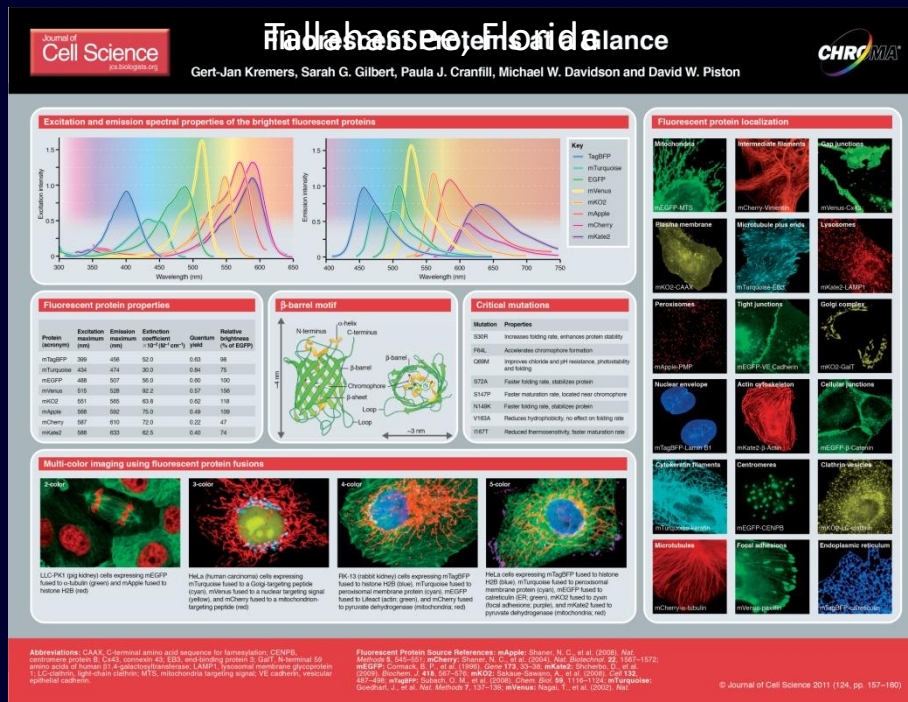
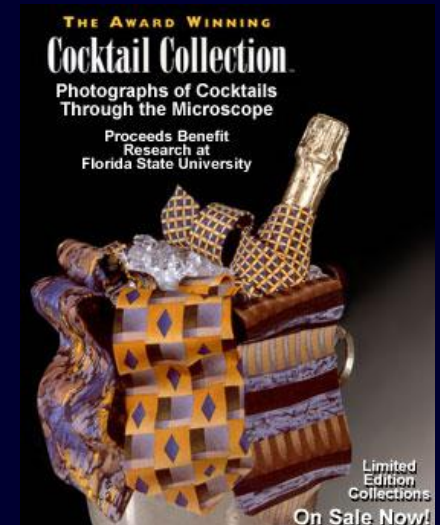
National High Magnetic Field Lab



Mike Davidson



Neckties®



website tutorials



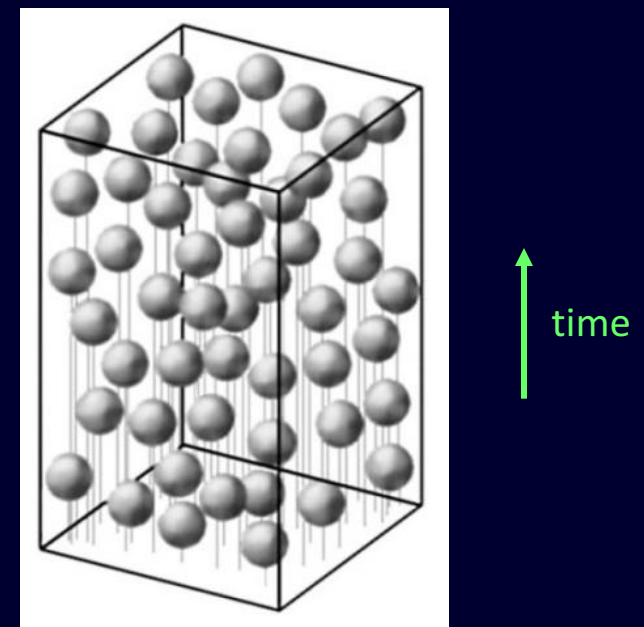
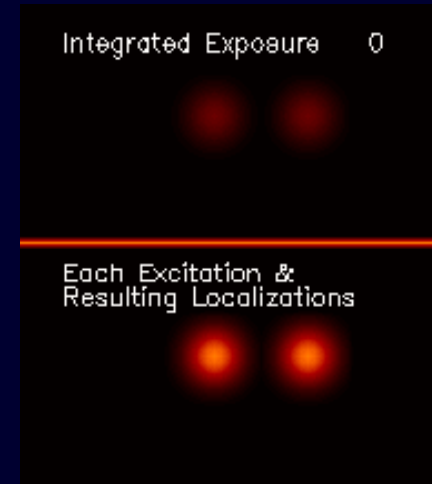
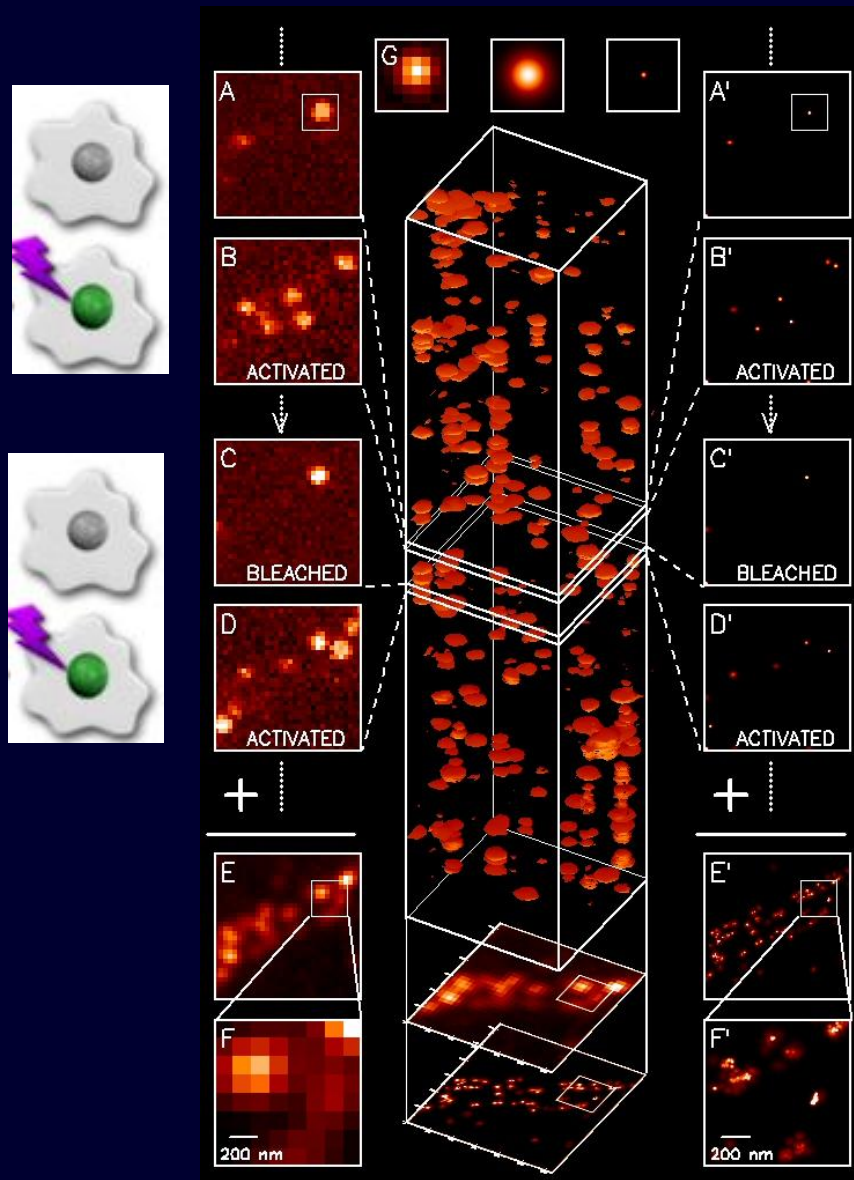
Zeiss



Olympus



Finding the Missing Link



E. Betzig, et al., *Science* **313**, 1642 (2006)

La Jolla Labs



Assembling the Rest of the Team

Jennifer
Lippincott-
Schwartz



George
Patterson



Rob Tycko, NIDDK

the microscope in the darkroom in Jennifer's lab



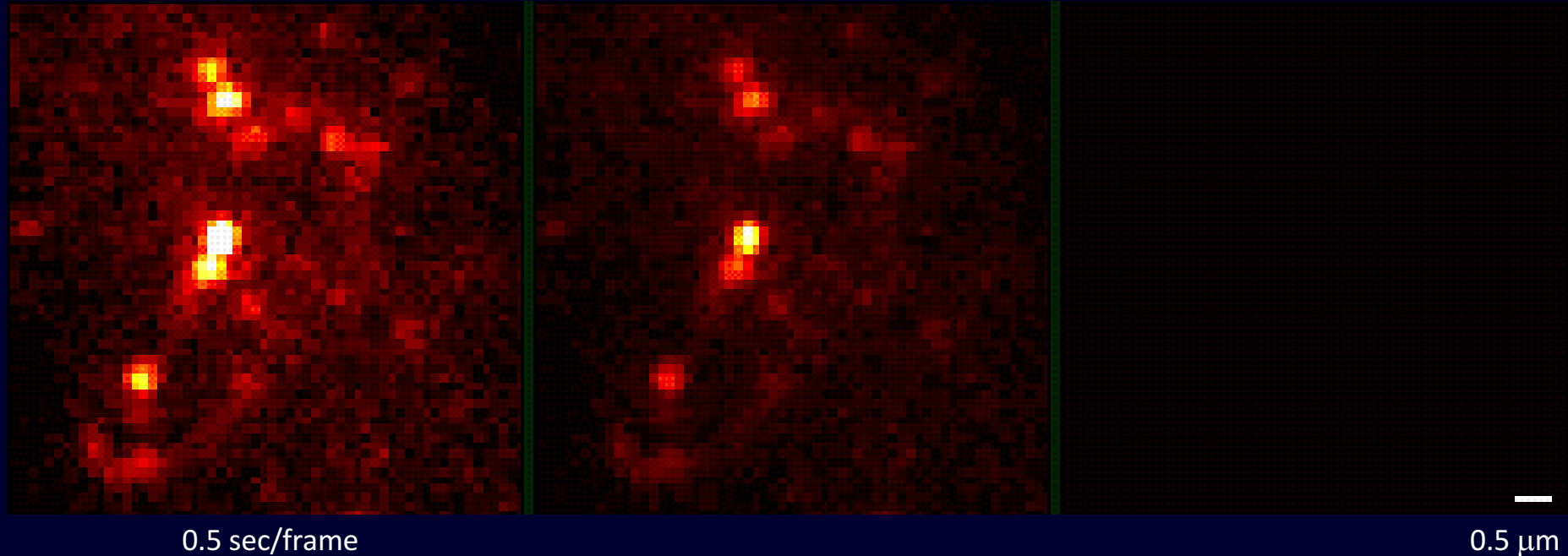
Photoactivated Localization Microscopy (PALM)

lysosomes, COS-7 cell, Kaede-tagged CD63

single molecule frames

integrated image

PALM image

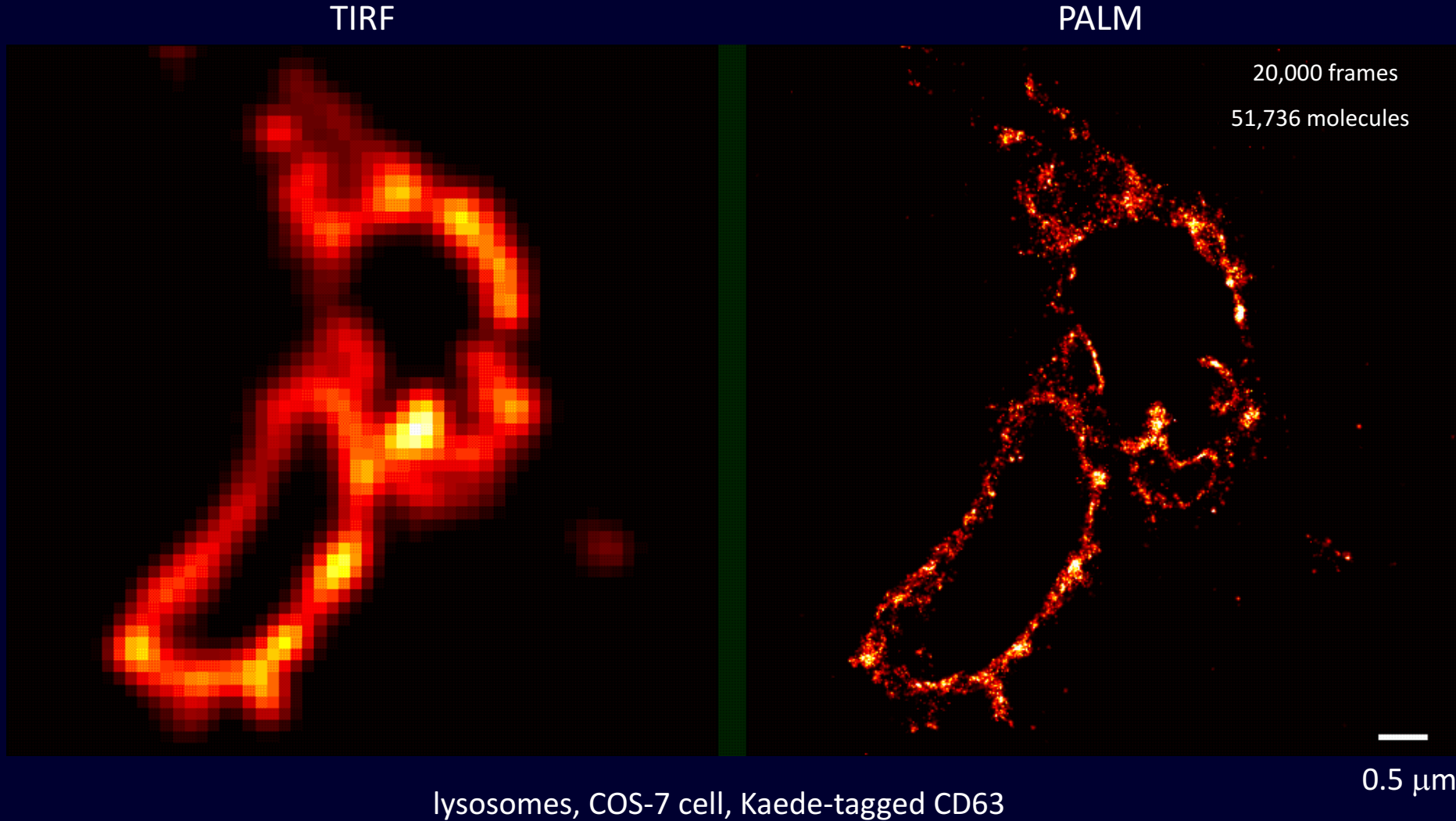


~80 nm cryosection:

- low autofluorescence
- immobile PA-FPs
- image internal organelles

E. Betzig, *et al.*, *Science* **313**, 1642 (2006)

Photoactivated Localization Microscopy (PALM)

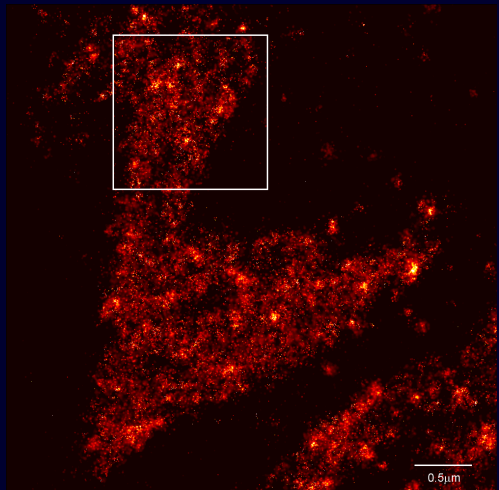


E. Betzig, *et al.*, *Science* 313, 1642 (2006)

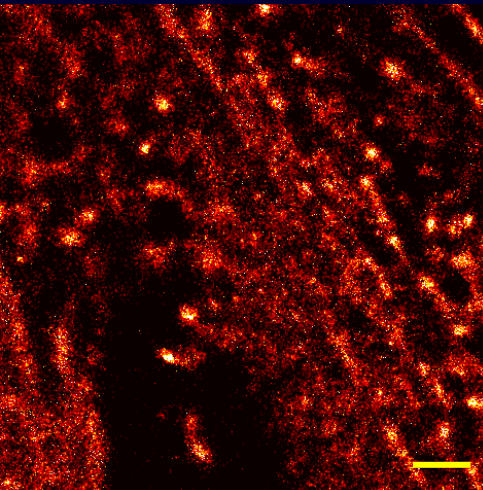
A High On/Off Contrast Ratio is Essential for High Resolution

paxillin, focal adhesions

EosFP > 2000:1

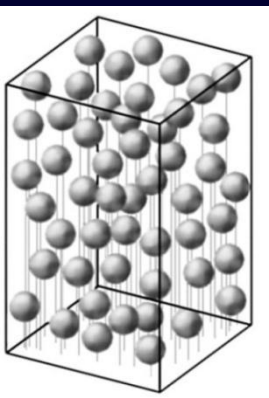
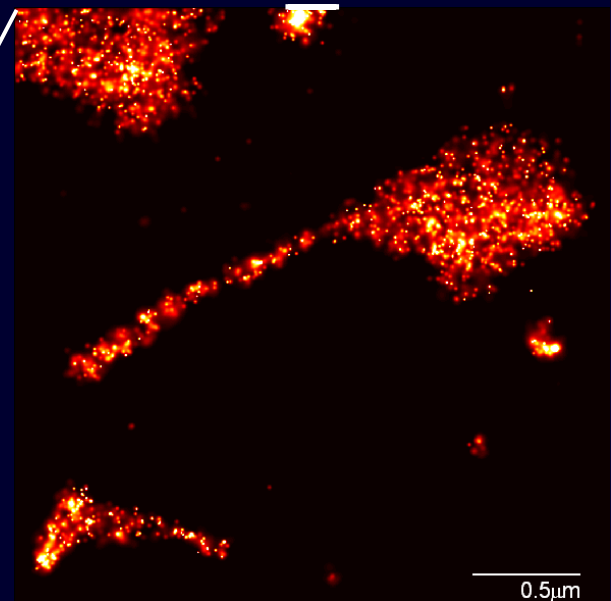


PA-GFP < 75:1

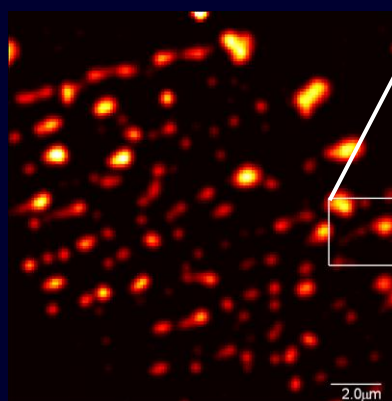


Eos FP and caged Q-rhodamine support Nyquist-defined sub-20 nm resolution

caged Q-rhodamine, > 1000:1



diffraction limited TIRF



E. Betzig, et al., *Science*
313, 1642 (2006)

From Rags to Riches, Thanks to HHMI

Janelia Research Campus



The Boss: Gerry Rubin

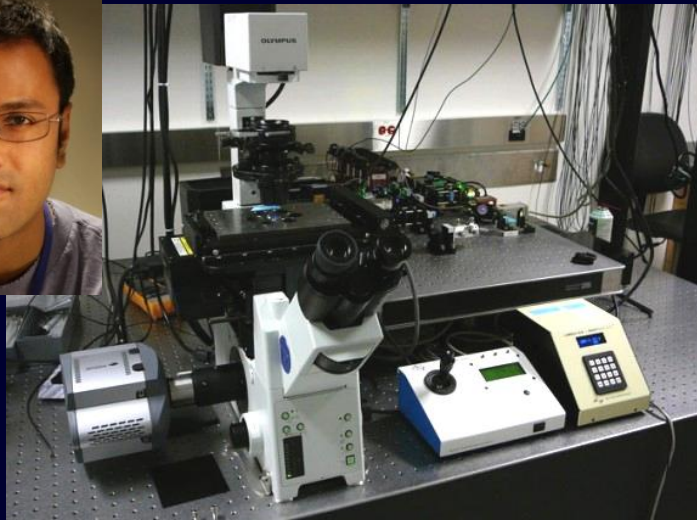


Endless Coffee



Hari Shroff

my PALM



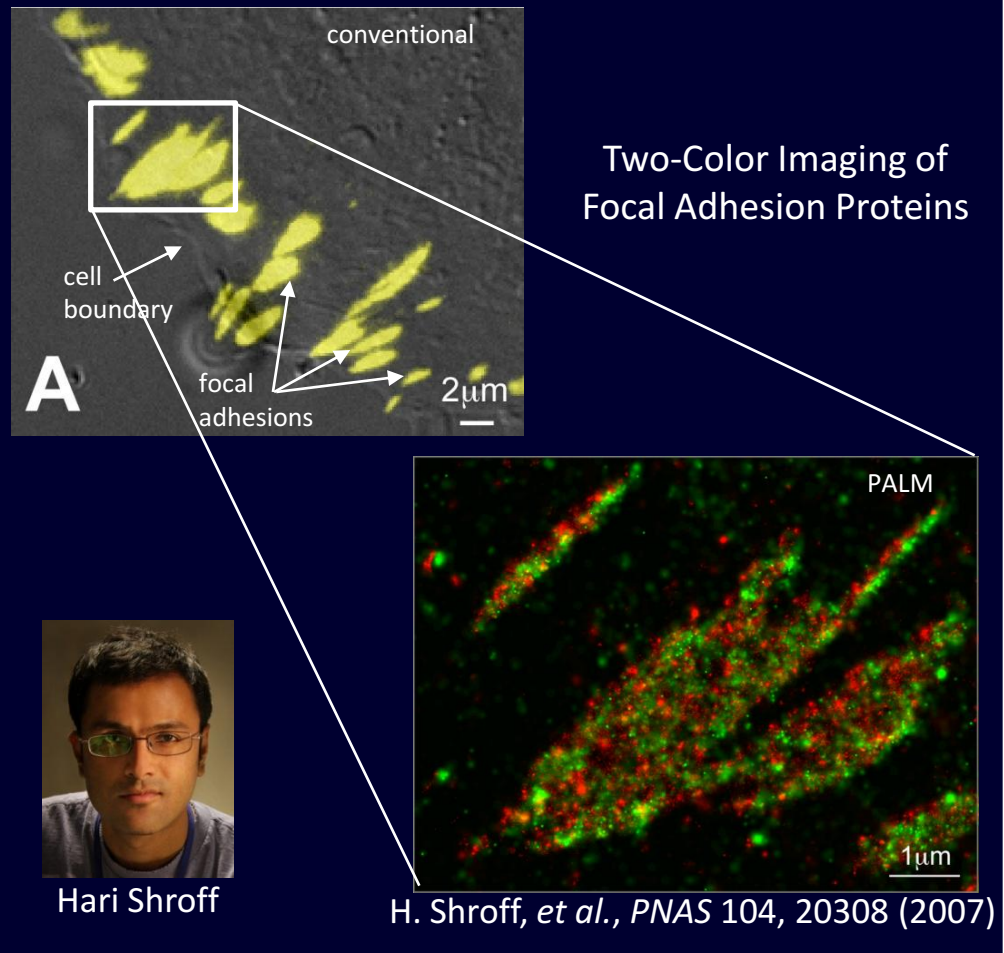
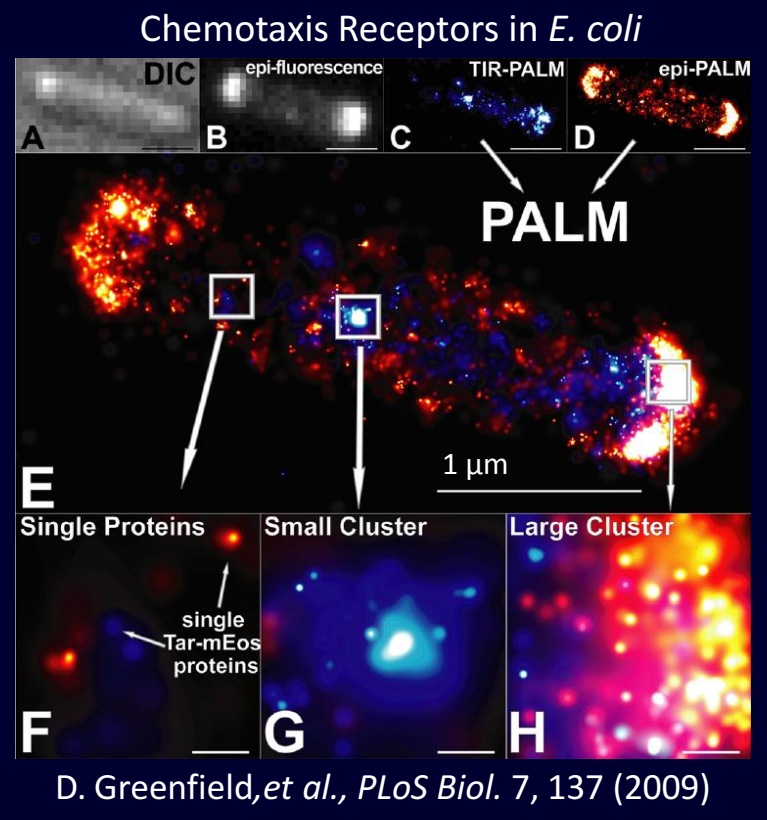
Gleb Shtengel



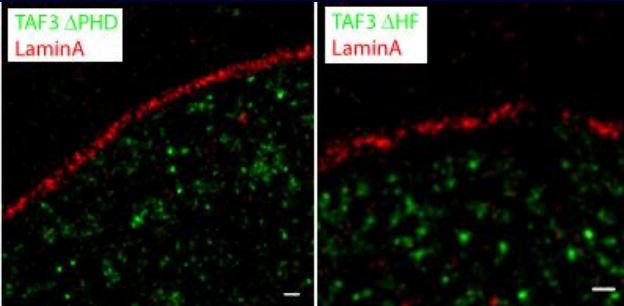
Harald's iPALM



PALM Application Examples

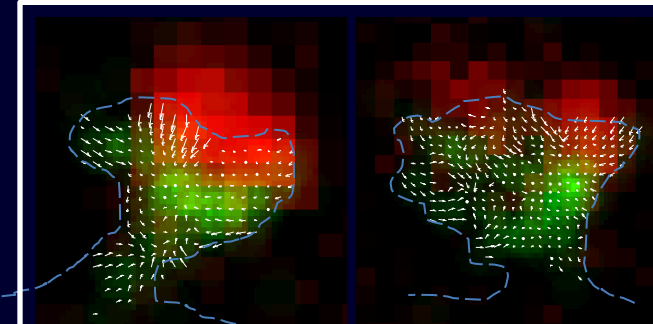


Regulation of Gene Expression During Myogenesis



J. Yao, et al., Genes Dev. 25, 569 (2011)

Actin Polymerization in Dendritic Spines



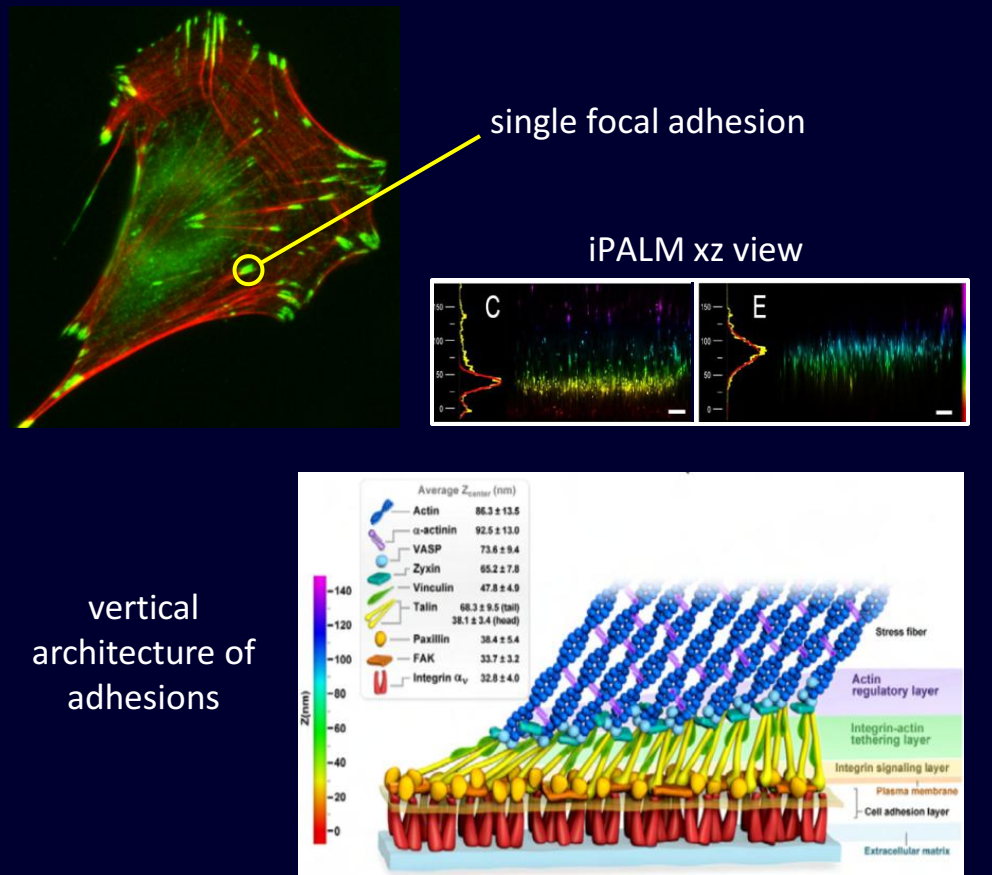
N. Frost, et al., Neuron 67, 86 (2010)

iPALM: Ultrasensitive PALM in 3D

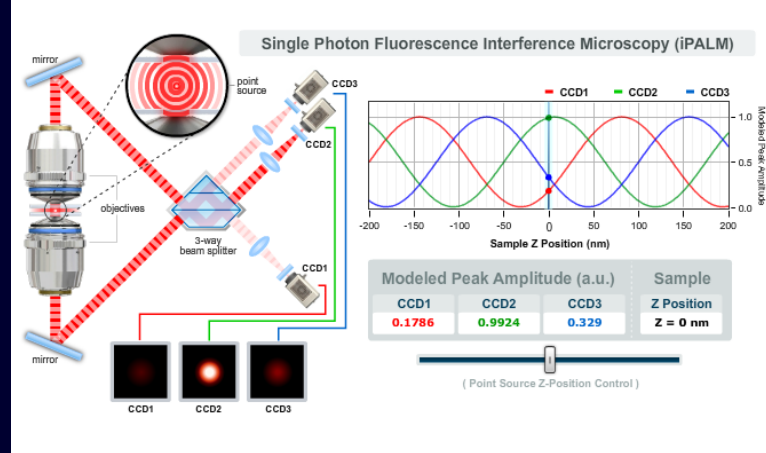
Harald Hess



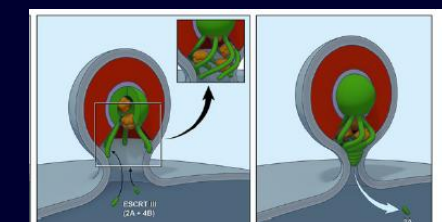
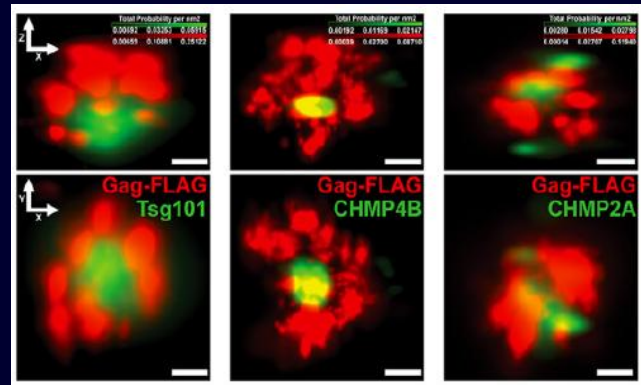
iPALM schematic



three phase single molecule interferometry



ESCRT machinery at HIV budding sites

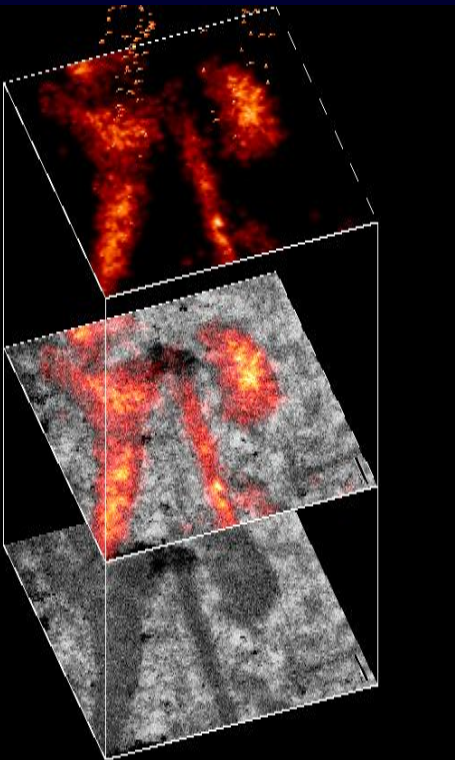


P. Kanchanawong, et al., *Nature* **468**, 580 (2010)

S.B. Van Engelenburg, et al., *Science* **343**, 653 (2014)

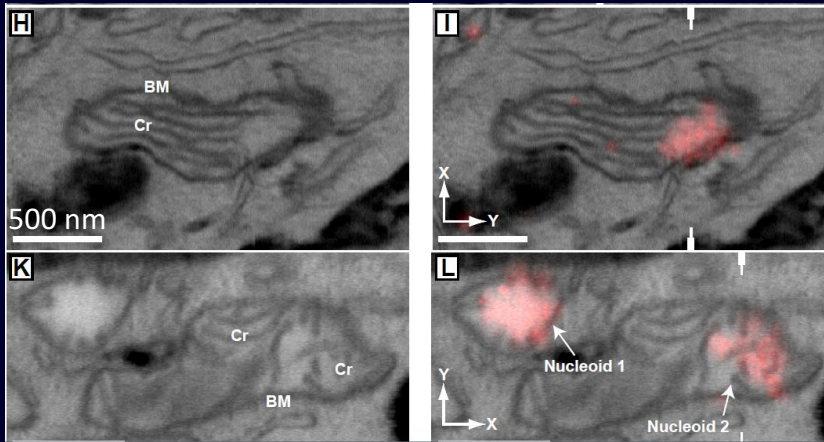
Correlative Electron Microscopy and PALM

first correlative EM
with super-resolution:
mitochondria

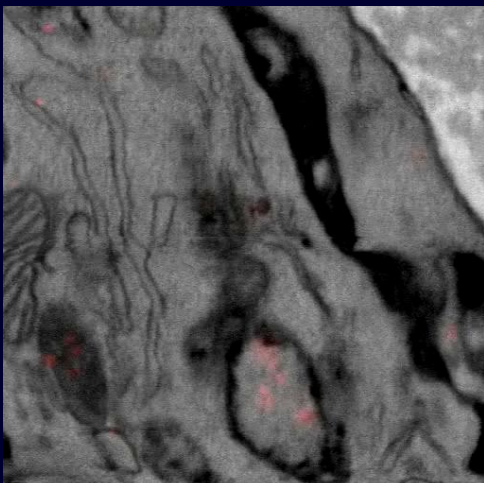


E. Betzig, et al., *Science*
313, 1642 (2006)

3D correlative EM/PALM
mitochondria (B&W – FIB SEM)
mitochondrial DNA (red - iPALM)

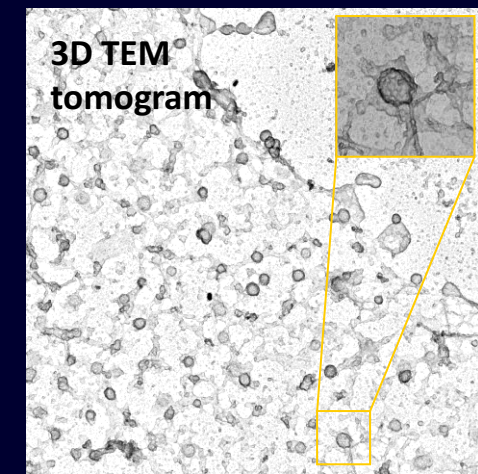


scrolling plane-by-plane thru 3D

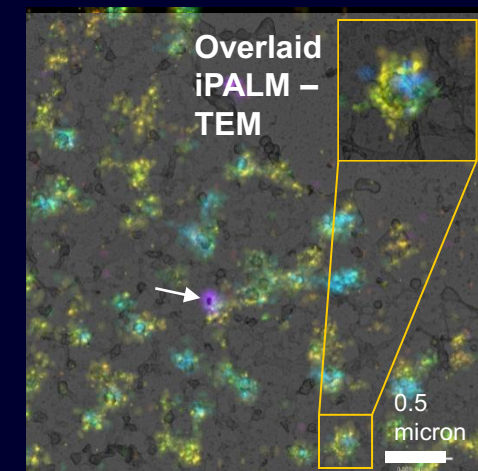


B.G. Kopek, et al., *PNAS*, 109, 6136 (2012)

cell membrane (B&W - TEM)
& clathrin (color - iPALM)

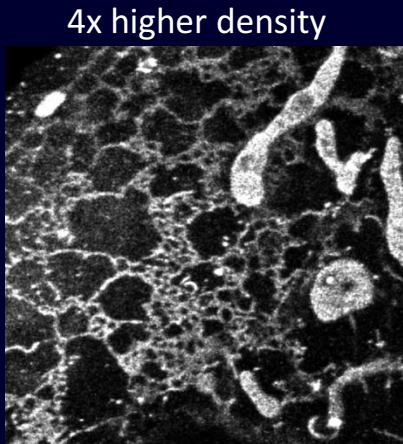
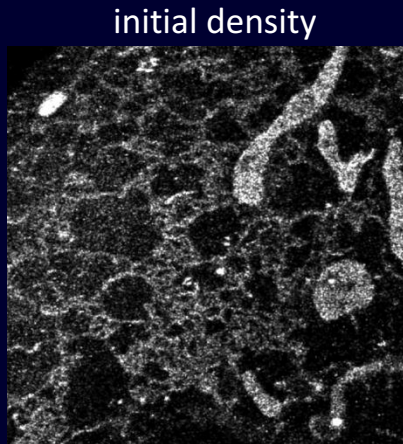


K. Sochaki, et al., *Nat. Methods*, 11, 305 (2014)

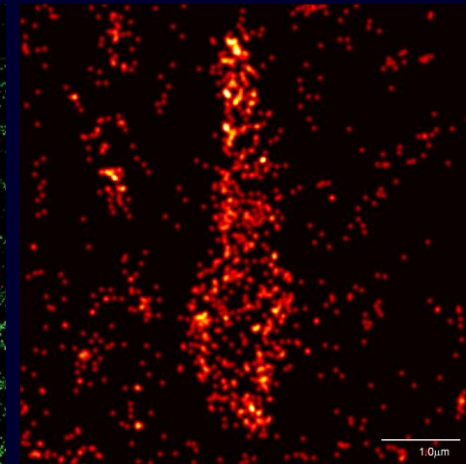
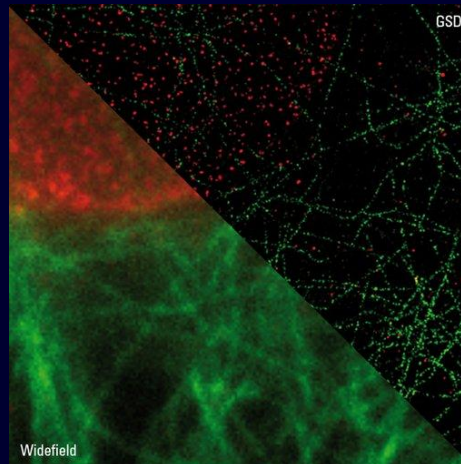


Caveats with Super-Resolution Microscopy: Fixed Cells

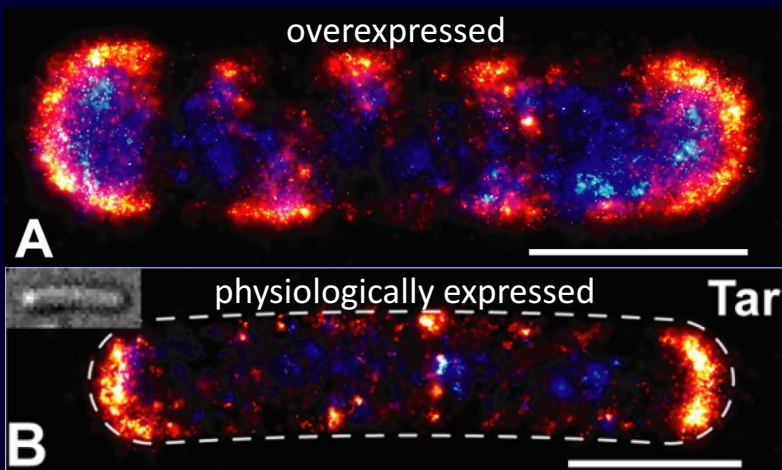
extremely high labeling densities required



exogenous dyes: limited affinity & high background



overexpression of protein



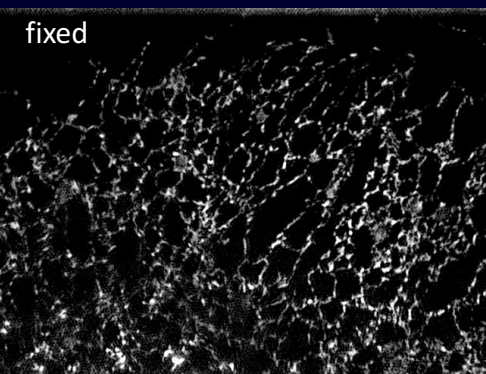
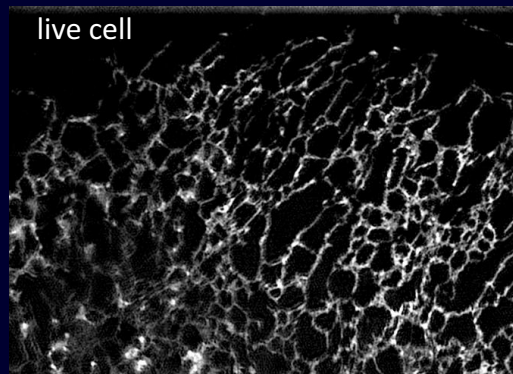
B

physiologically expressed

Two fluorescence microscopy images of fixed cells showing physiologically expressed protein. The signal is more evenly distributed and localized to specific cellular structures. A scale bar is present at the bottom right.

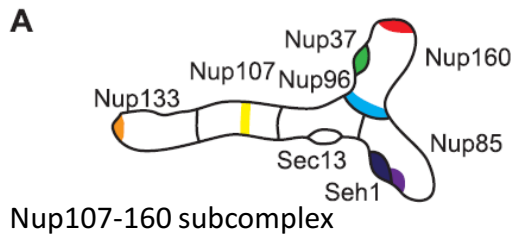
Tar

fixation artifacts, endoplasmic reticulum

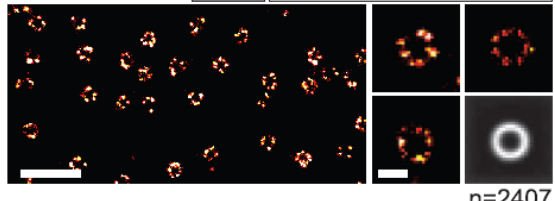


Particle Averaging Improves Resolution of Stereotypic Structures

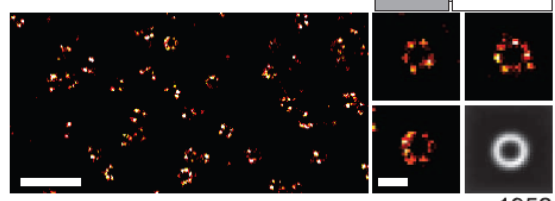
nuclear pore complex proteins



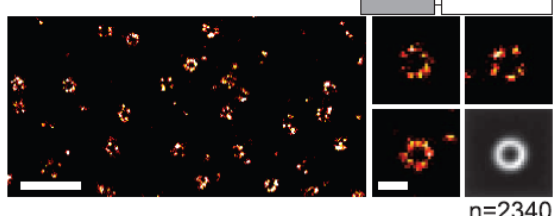
C EGFP-Nup107 (107-N)



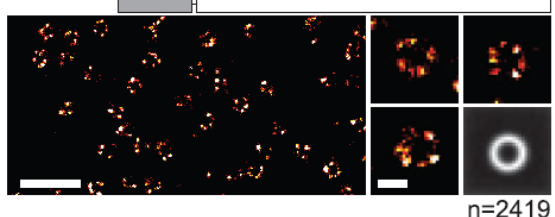
E EGFP-Nup37 (37-N)



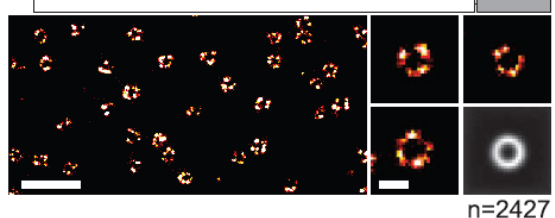
G mEGFP-Seh1 (Seh1-N)



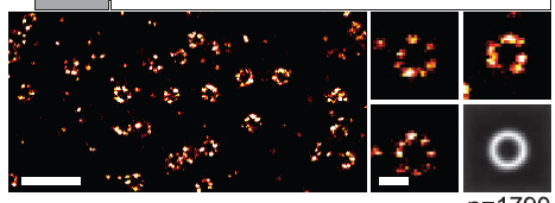
B mEGFP-Nup133 (133-N)



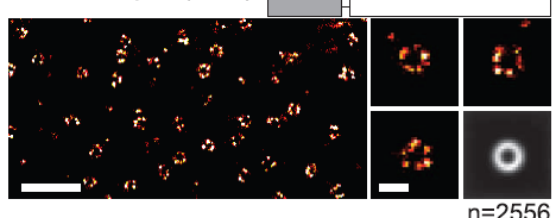
D Nup160-mEGFP (160-C)



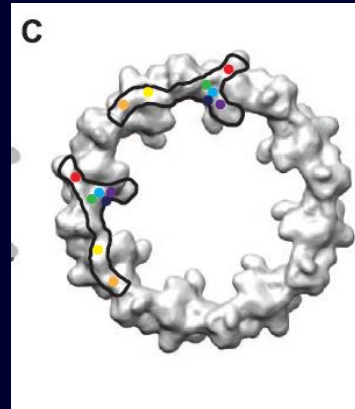
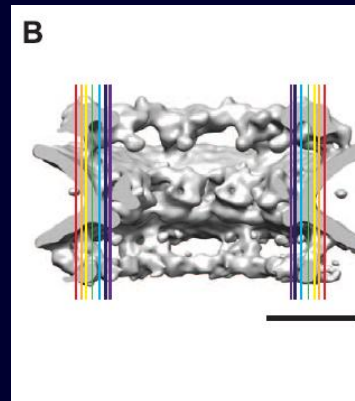
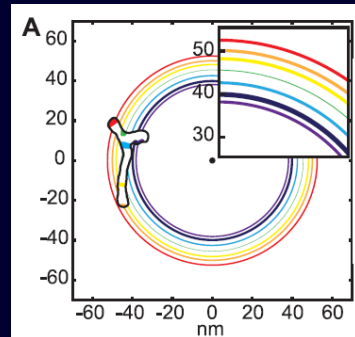
F mEGFP-Nup160 (160-N)



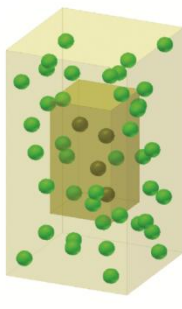
H mEGFP-Nup85 (85-N)



positions
determined
to < 1 nm



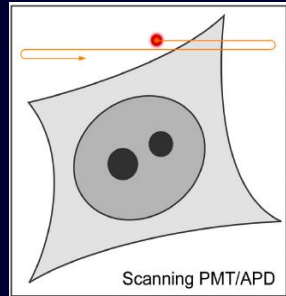
Caveats with Super-Resolution Microscopy: Live Cells



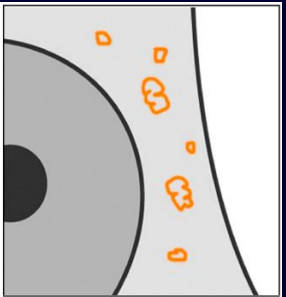
Nyquist criterion:

N -fold resolution increase in D dimensions $\rightarrow N^D$ -fold more photons collected

STED / RESOLFT



Excitation PSF
+
STED pulse PSF
=
Effective PSF
(PSF shaping)



reported resolution (nm)

xy: 20 nm

xyz: 30 nm

photon increase required

100

intensity (W/cm²)

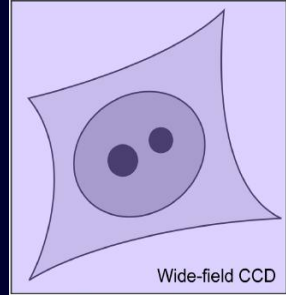
$10^4 - 10^9$

acquisition time (sec)

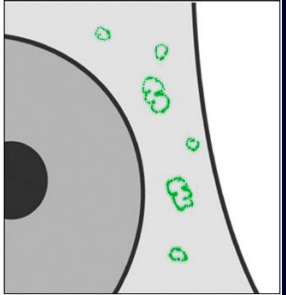
> 60

~1,000

Localization



Time series (few 1,000 exposures)
Wide-field CCD



xy: 20 nm

xy: 10 nm,
z: 20 nm

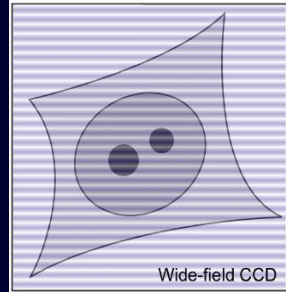
100

$10^3 - 10^4$

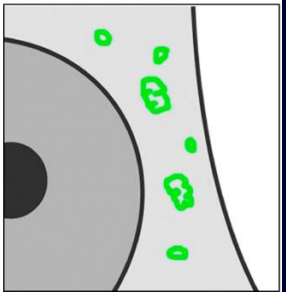
>20

1,500

SIM



5 phase shifts
5 phase shifts
5 phase shifts
Interference of exciting light with sample structure (Moiré effect)
Mathematic reconstruction



xy: 100 nm

xy: 100 nm,
z: 370 nm

4

$10 - 10^2$

0.1 - 1

~10

Live Cell Structured Microscopy

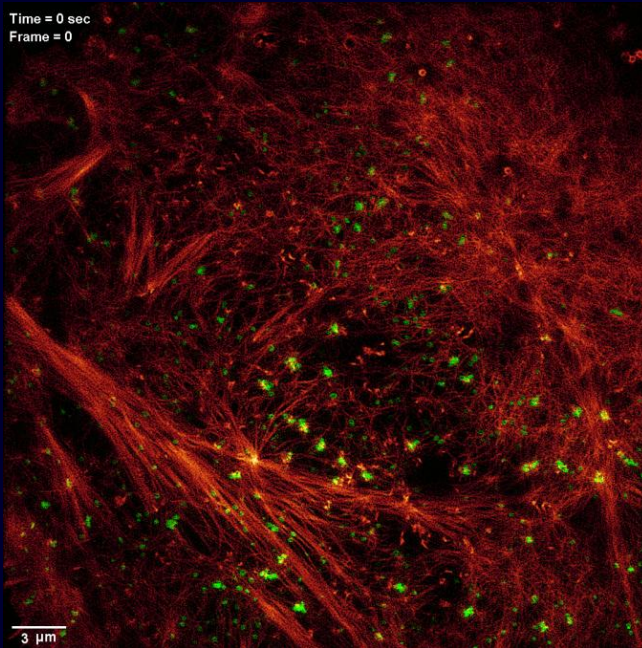
endoplasmic reticulum

2D SIM, 98 nm resolution
0.1 sec acquisition, **1800 frames**



clathrin coated pits and cortical actin

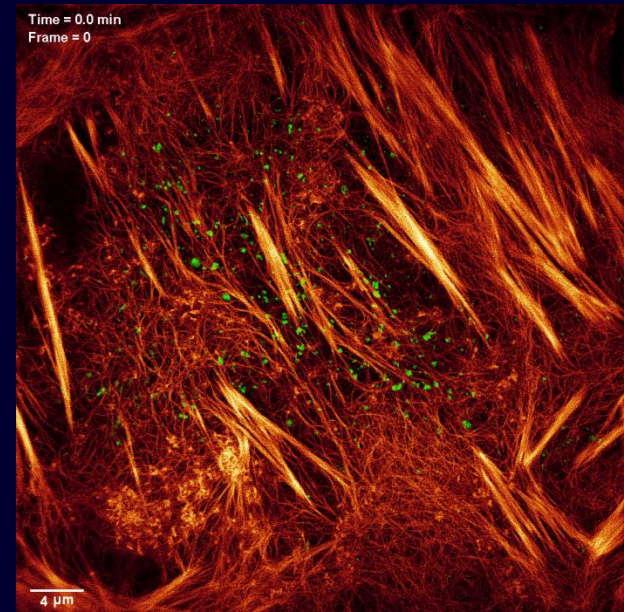
TIRF-SIM, 82 nm resolution
0.5 sec acquisition, 90 frames



Mats
Gustafsson,
1960-2011

early endosomes and cortical actin

Nonlinear SIM, **62 nm resolution**
1.5 sec acquisition, 34 frames



Dong
Li



Lin
Shao

The Challenges and Importance of Studying Live Cell Dynamics

tradeoffs, tradeoffs, tradeoffs

spatial resolution

photo-toxicity

imaging depth

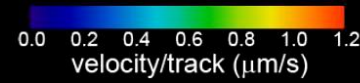
temporal resolution

Life is Animate

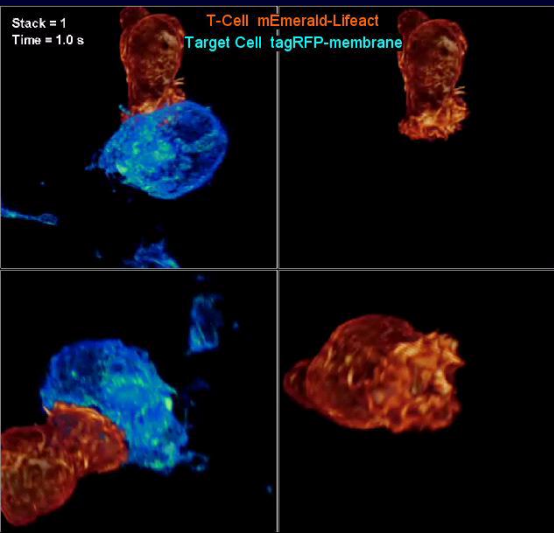
dividing HeLa cell

prometaphase

00:00:00

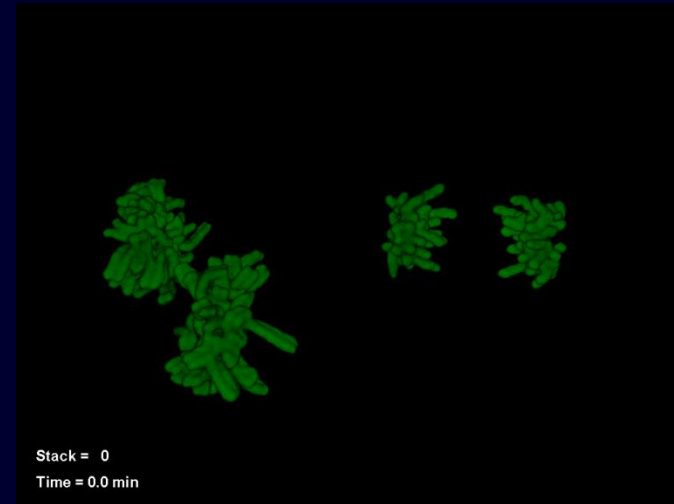
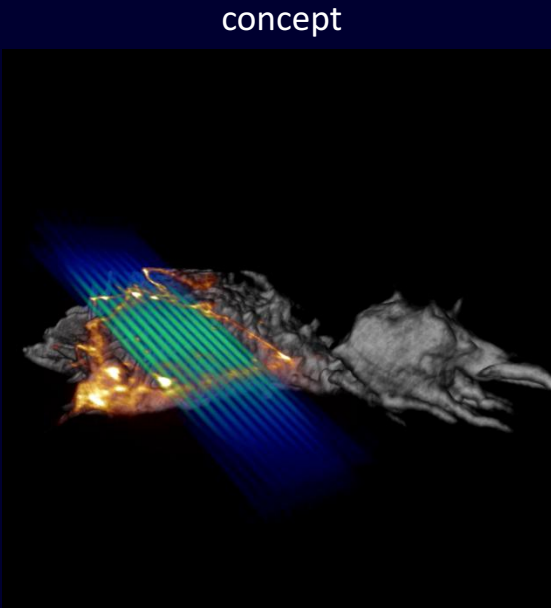


Lattice Light Sheet Microscopy: Non-Invasive 4D Live Cell Imaging

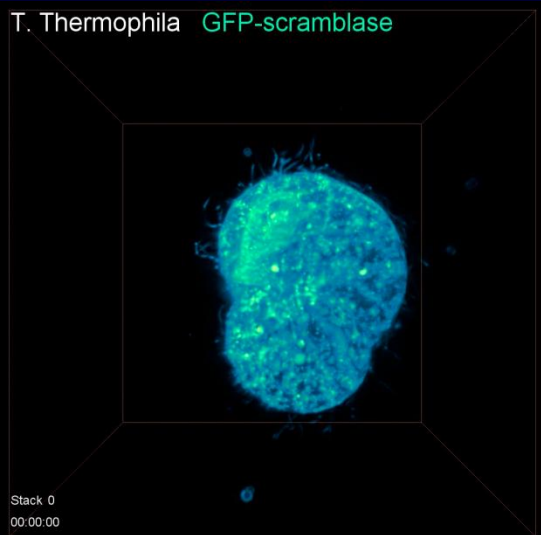


T cell and its target cell

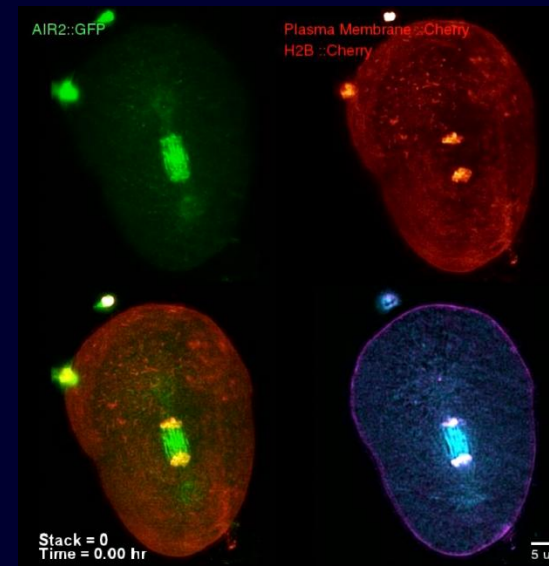
chromosomes, mitos, and ER during mitosis



Tetrahymena thermophila



C. elegans early embryo



Bi-Chang Chen



Wes Legant



Kai Wang

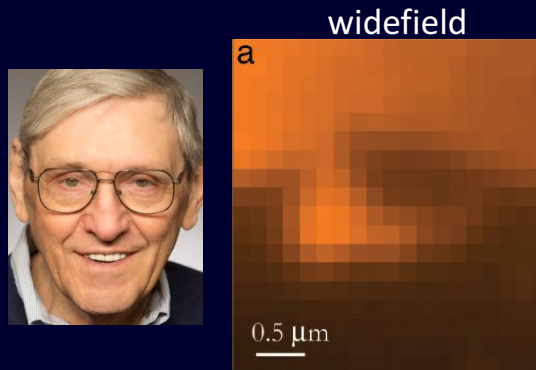
B-C Chen, et al., *Science* **346**, 1257998 (2014)

Ultra-High Density 3D Localization Microscopy

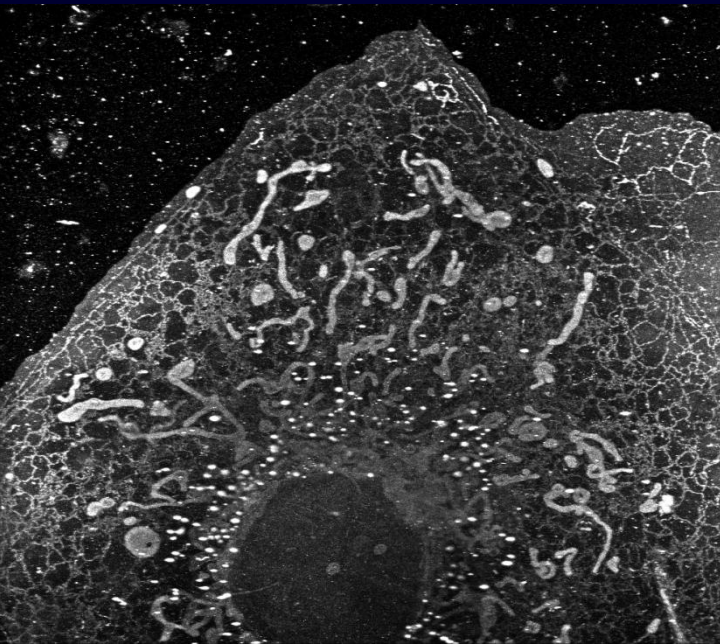


Wesley Legant

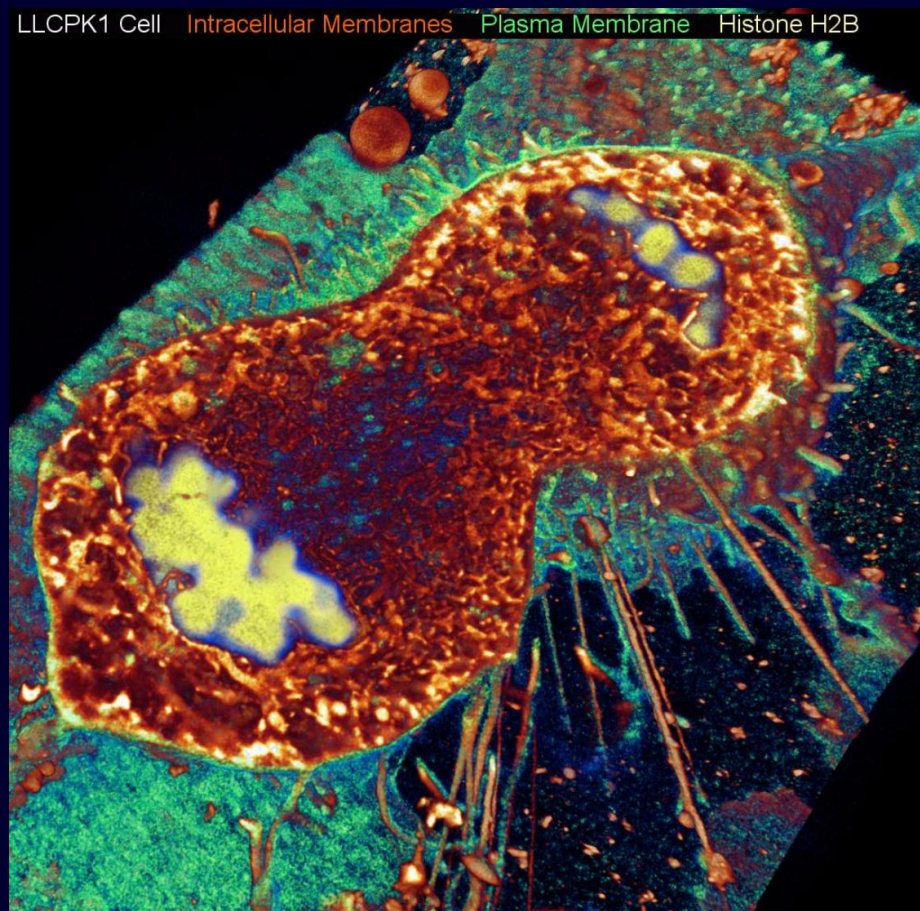
Points Accumulation for Imaging in Nanoscale Topography (PAINT)



A. Sharonov, R.M. Hochstrasser, *PNAS* 103, 18911 (2006)



3D PAINT with lattice: dividing cell



over 300 million localized molecules

Adaptive Optics (AO): Moving Cell Biology Away from the Cover Slip

non-scattering media: zebrafish embryonic brain



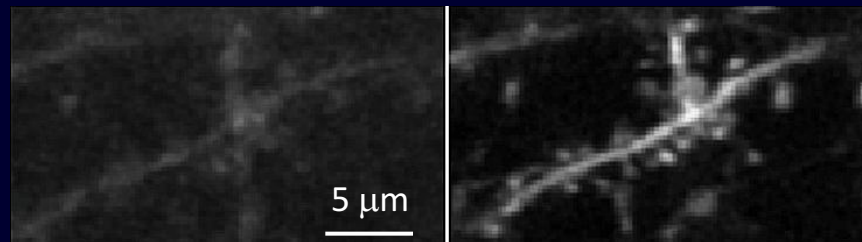
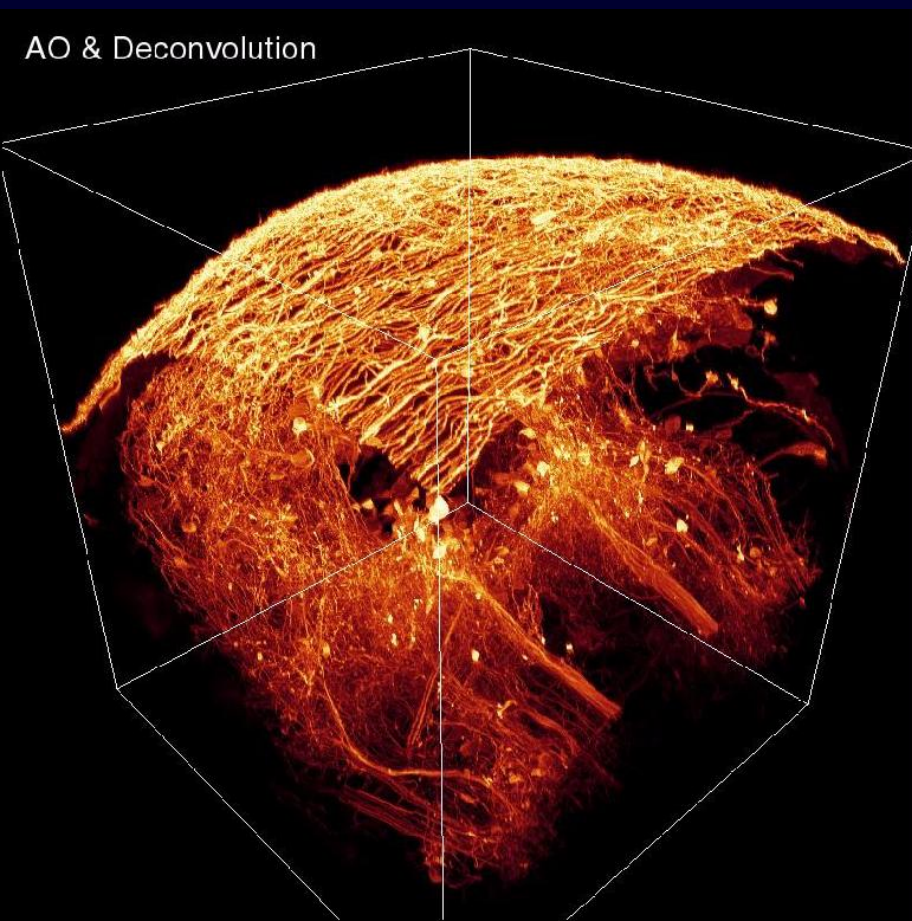
Kai Wang

scattering media:
mouse visual
cortex

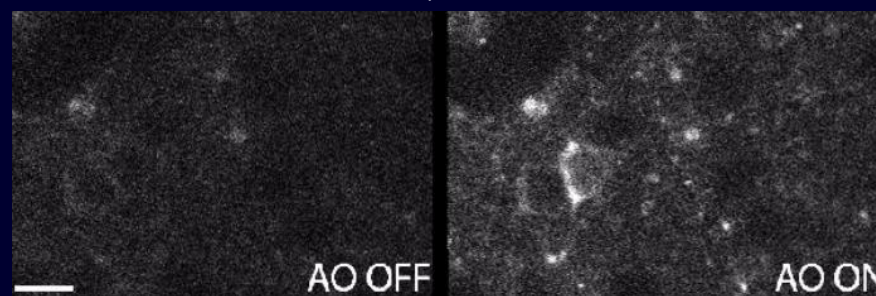


Na Ji

AO & Deconvolution



functional imaging of neural activity, 400 μm deep



The Beauty and Complexity of Living Systems

