# Lecture 15

# Single Molecule Localization Microscopy

18 March 2015

Rainer Kaufmann

rainer@strubi.ox.ac.uk

# localization microscopy – one technique, many acronyms



# single molecule localization microscopy

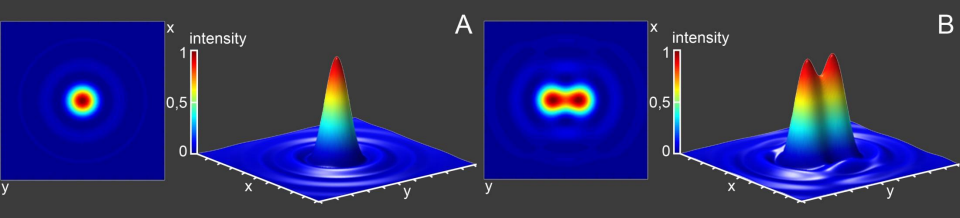
#### Outline:

- introduction and general idea of single molecule localization microscopy
- first approaches: "original" (F)PALM and STORM
- dSTORM, SPDM, GSDIM using standard fluorophores
- 3D
- live-cell (4D)
- quantitativ analysis using the additional single molecule information
- alternative approaches
- conclusion

# introduction to localization microscopy

problem in light microscopy: resolution limited by diffraction

$$\Rightarrow I_{im}(x_2, y_2) = PSF \otimes P_{fl}(x_1, y_1)$$



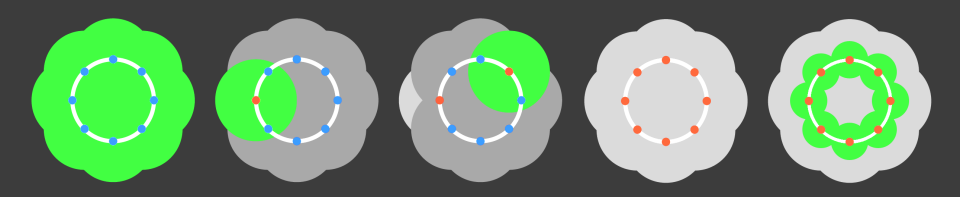
Rayleigh criterion: 
$$D = 0.61 \frac{\lambda}{NA}$$

# introduction to localization microscopy

### general idea:

look at signals of single molecules individually instead of all fluorophores at the same time

- → this allows a very precise determination of the molecule position
- → reconstruct super-resolution image from position data of the detected molecules



# introduction to localization microscopy

setup

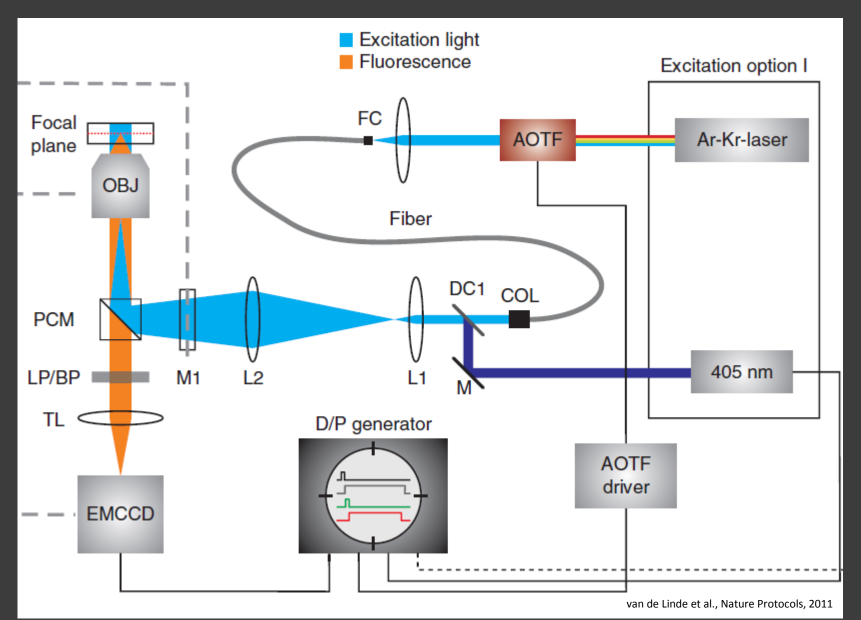
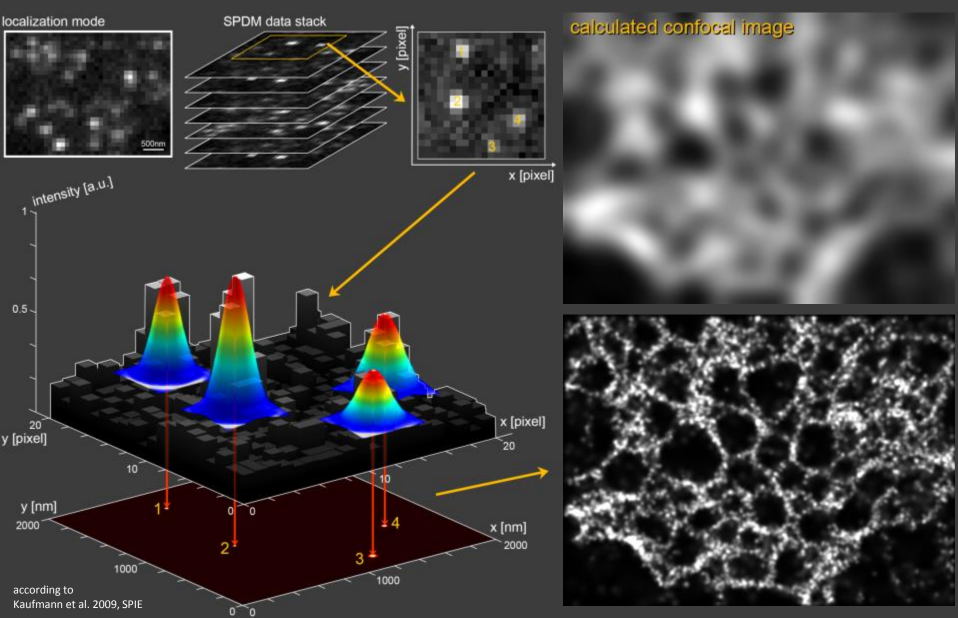


image reconstruction



position determination

#### localisation accuracy $\sigma$ of a single molecule is depended on

- width of the PSF s
- number of detected photons N
- background intensity b
- size of the pixels on the camera a

$$\sigma^2 = \frac{s^2 + a^2/12}{N} + \frac{8\pi s^4 b^2}{a^2 N^2}$$

typical model function: 2D Gaussian + linear background

$$I(x,y) = I_0 \exp\left[-\frac{(x-x_0)^2 + (y-y_0)^2}{2s^2}\right] + b$$

resolution

#### structural resolution in localization microscopy is dependent on:

- the localization accuracy of the individual molecules
- density of detected molecules (sampling theorem Nyquist resolution)

$$\Rightarrow structural\ resolution = \sqrt{(2.35\ \bar{\sigma}_{xy})^2 + (2\ \bar{d}_{NN})^2}$$
$$= \sqrt{(2.35\ \bar{\sigma}_{xy})^2 + 4/\rho}$$

 $\bar{\sigma}_{xy}$ : mean localization accuracy

 $ar{d}_{NN}$ : mean distance to next neighboring molecule(s)

 $\rho$ : local density of detected molecules

resolution

#### structural resolution in localization microscopy is dependent on:

- the localization accuracy of the individual molecules
- density of detected molecules (sampling theorem Nyquist resolution)



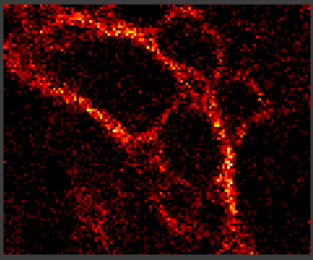
 $\widetilde{d}_{N4N} = 261 \text{ nm}$ 

500nm

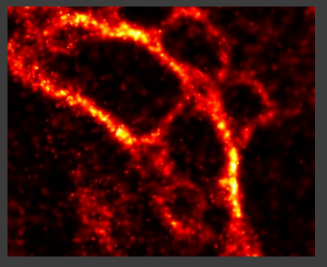
image reconstruction



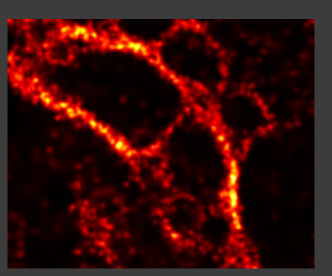
scatter plot



histogram with equal bins



visualisation of  $\sigma_{\chi \gamma}$ 



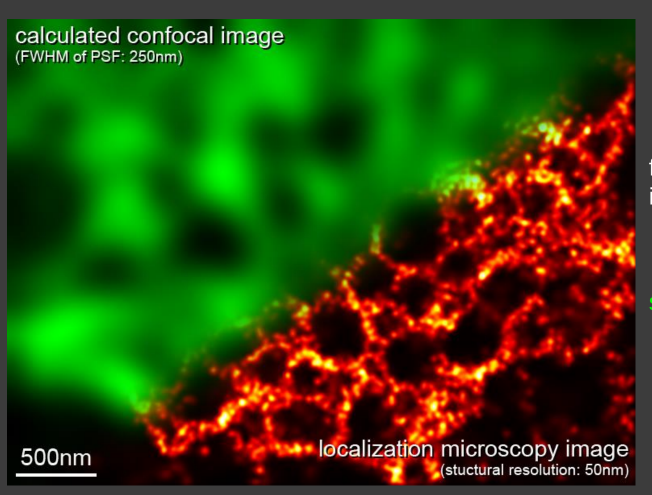
visualisation of structural resolution

more about visualisation of localization microscopy data:

Baddeley et al., Microscopy and Microanalysis, 2010

summary

enhanced structural resolution down the 20 nm range





fluorophores are detected individually



#### single molecule information

- positions
- number of det. photons
- ...

# (F)PALM and STORM

# (some) history of localization microscopy

#### localisation of single molecules / point-like objects

Burns et al., 1985 theoretical paper about super-resolution distance

measurements using spectral characteristics

Betzig, 1995 first measurements with SNOM under cryo conditions

Bornfleth et al., 1998 CLSM measurements of 3D distances < 60 nm using

fluorescent markers of different wavelengths (@ RT)

Heilemann et al., 2002 using single molecule live time instead of colours to measure

distances of 40 nm

#### localisation of many molecules to reconstruct structural information

2006: (PALM, FPALM, STORM) – photo-switchable / photo-activatable dyes

2008: (dSTORM, SPDM, GSDIM) – using standard fluorophores

# (F)PALM – (fluorescence) photo activated localization microscopy

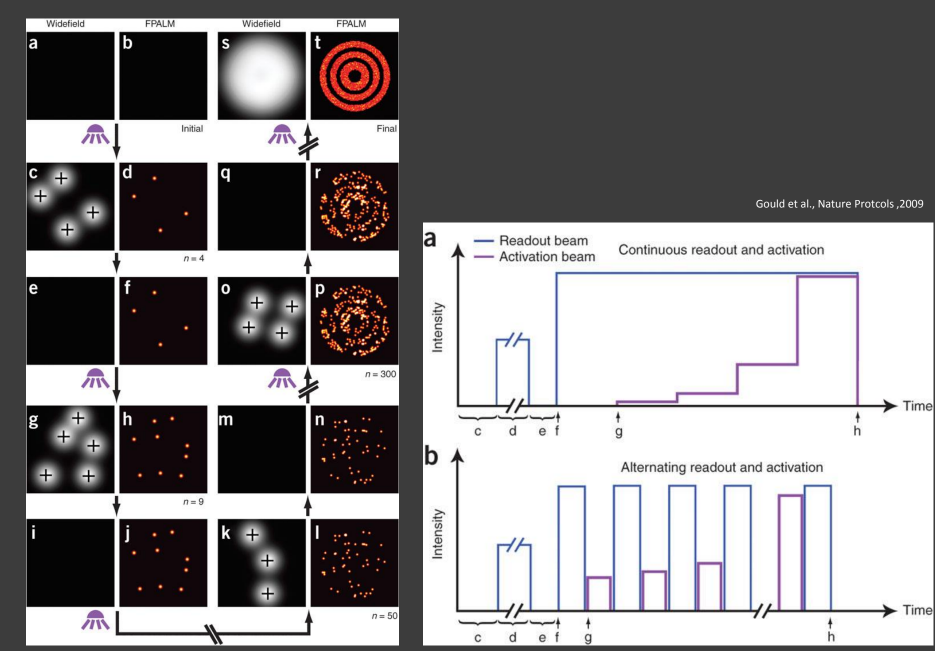
uses photo-activatable fluorophores (e.g. PA-GFP, caged Fluorescein, ...)

- at the beginning all fluorophores are "dark" (not fluorescent at their excitation wavelength)
- fluorophores can be "activated" to a "bright" state
- after bleaching the molecules they do not reappear
- → irreversible process

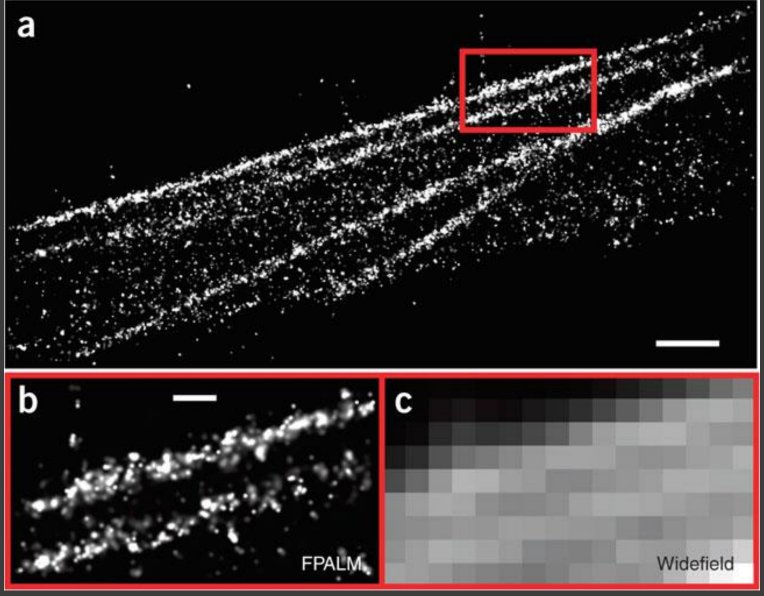
#### original publications:

- PALM: Betzig et al., Science, 2006
- FPALM: Hess et al., Biophysical Journal, 2006

# (F)PALM – (fluorescence) photo activated localization microscopy



# (F)PALM – (fluorescence) photo activated localization microscopy



Gould et al., Nature Protcols, 2009

### STORM – stochastic optical reconstruction microscopy

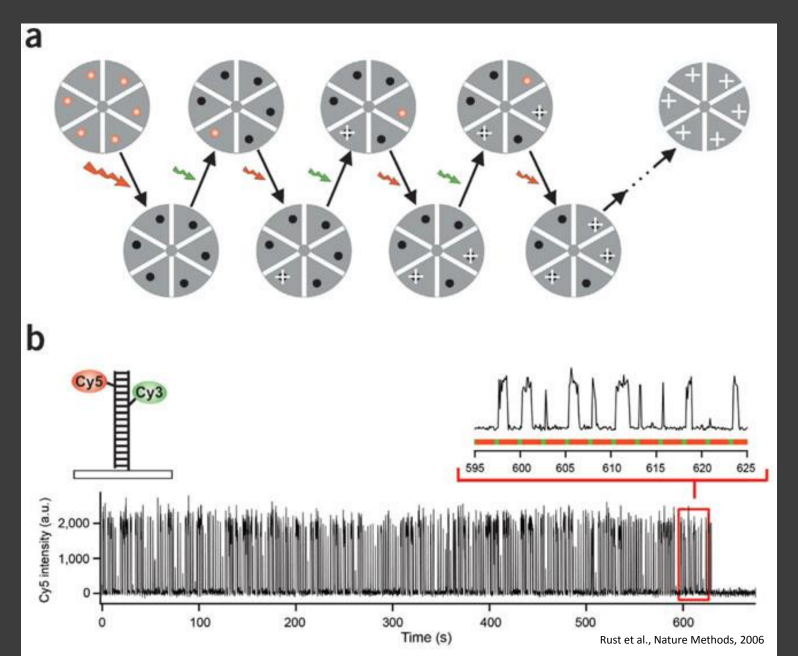
uses photo-switchable fluorophores (dye pairs (e.g. Cy3-Cy5) or proteins like Dronpa)

- fluorophores can be switched many times between a "bright" and a "dark" state
- → reversible process

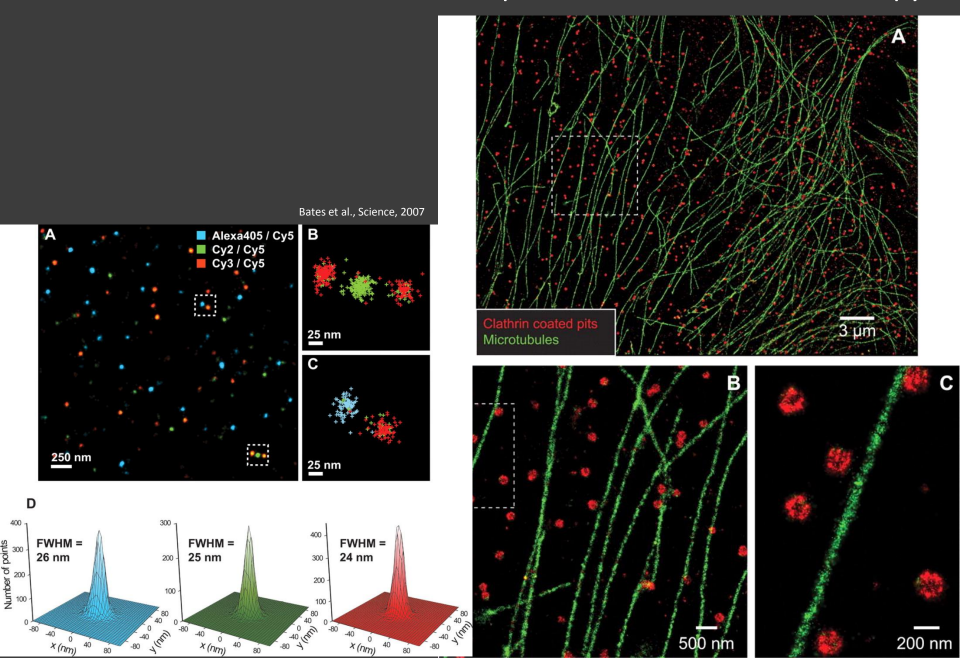
#### original publication:

Rust et al., Nature Methods, 2006

# STORM – stochastic optical reconstruction microscopy



# STORM – stochastic optical reconstruction microscopy



direct STROM

spectral position determination microscopy

ground state depletion microscopy followed by individual molecule return

uses standard fluorophores (e.g. Alexa and Atto dyes, GFP, YFP, RFP, ...)

- switching mechanism based on a light induced long-lived "dark" state
- stochastic recovery to "bright" (fluorescent) state is used for optical isolation of the single molecule signals

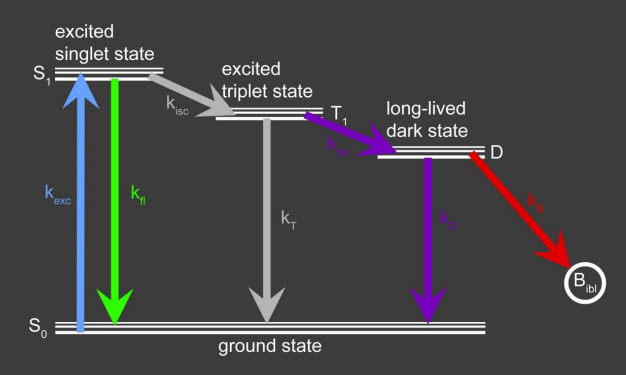
#### original publication:

- dSTORM: Heilemann et. al., Angewandte Chemie International Edition, 2008
- SPDM: Lemmer et al., Applied Physics B, 2008
- GSDIM: Fölling et al., Nature Methods, 2008

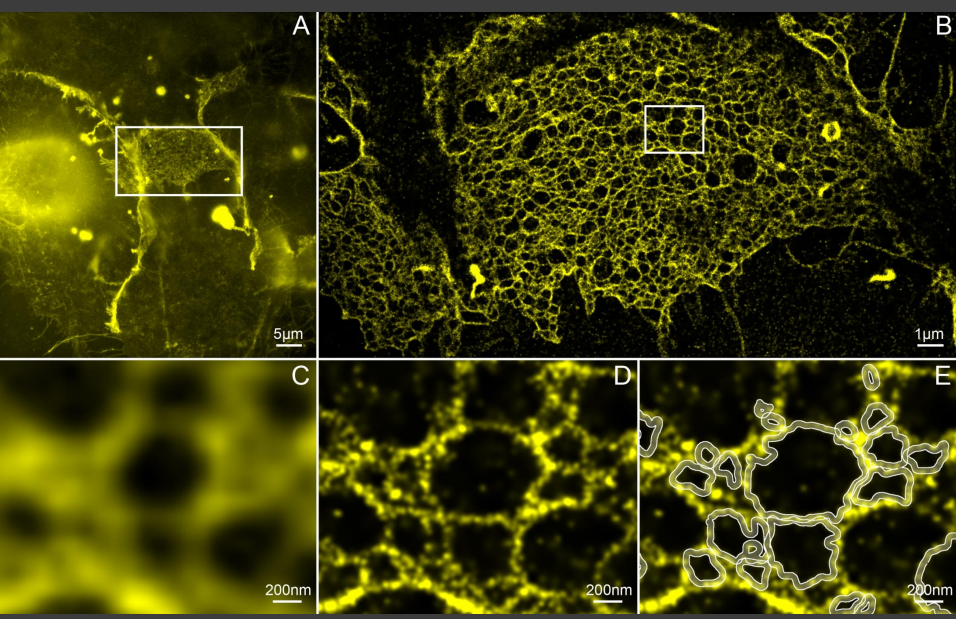
light induced long-lived (ms – 100 s) dark state

critical parameters for driving fluorophores into the long-lived dark state:

- illumination intensity
- wavelength
- embedding medium



statistical recovery of fluorophores from the light induced long-lived dark state can be used for optical isolation of single molecules

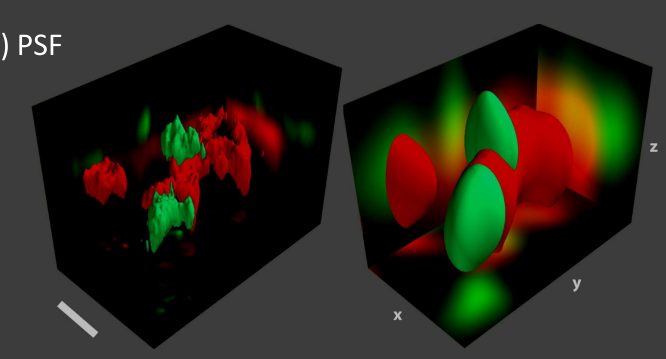


EYFP-Cld3

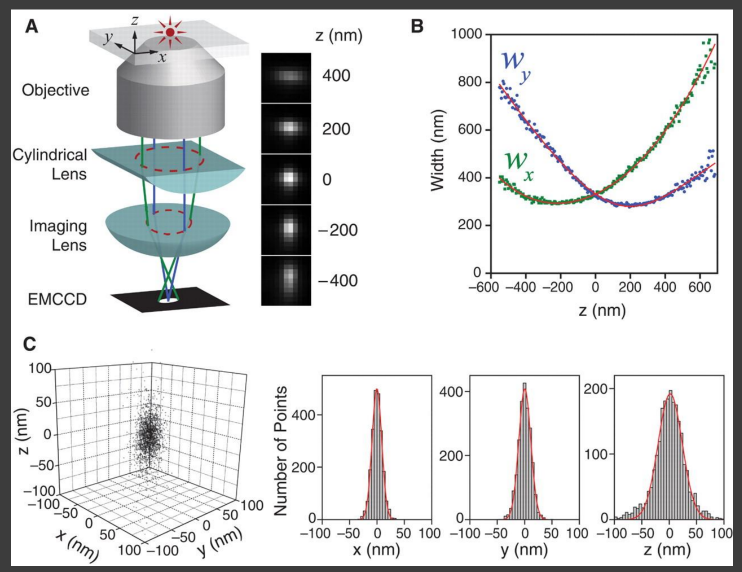
Kaufmann et al., PLoS ONE, 2012

# 3D

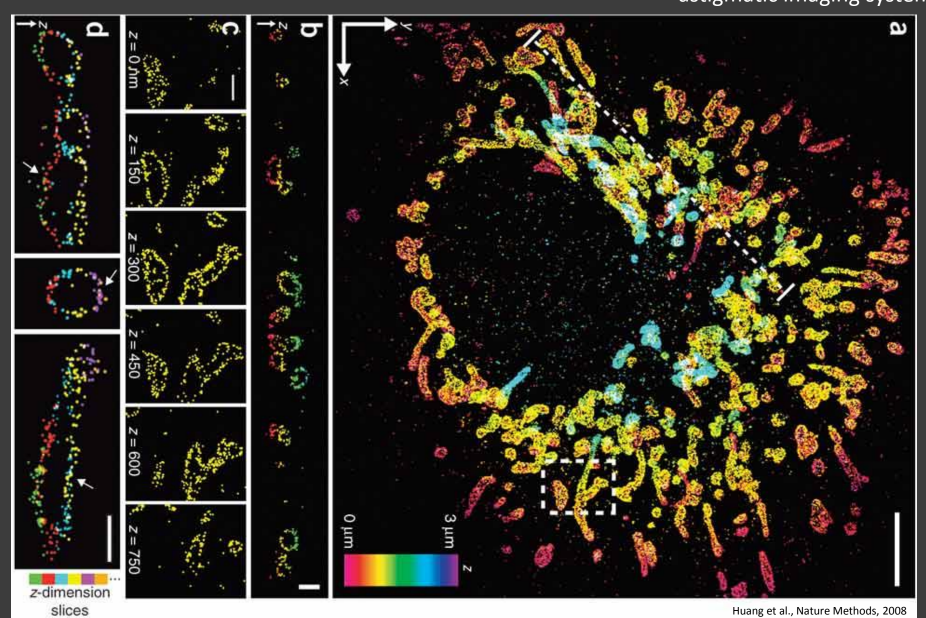
astigmatic (elliptical) PSF
biplane
double helical PSF
iPALM



#### astigmatic imaging system

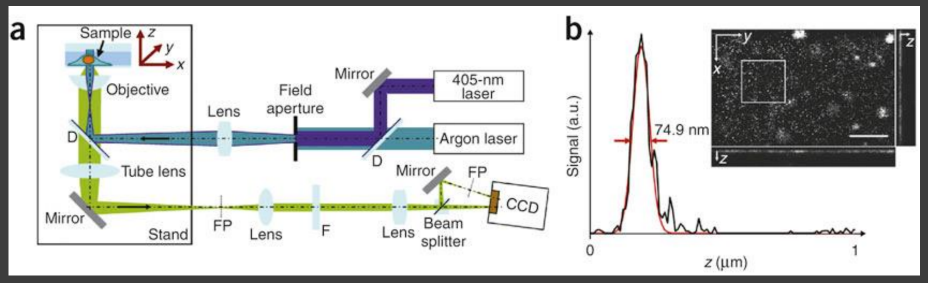


resolution lateral: 30 nm axial: 50 nm



imaging of two different axial plane simultaneously

→ fitting of 3D-PSF yields 3D position of the fluorophore



Juette et al., Nature Methods, 2008

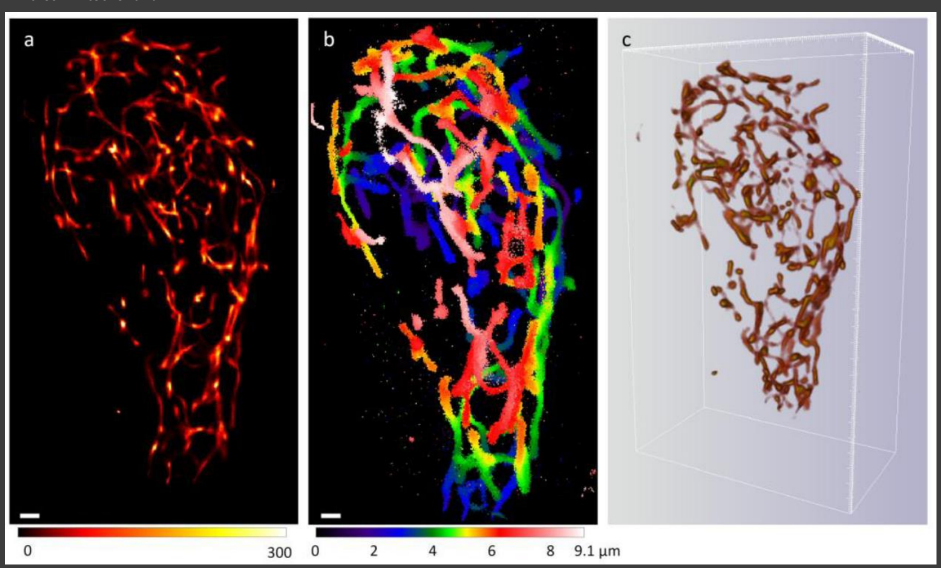
resolution

lateral: 30 nm

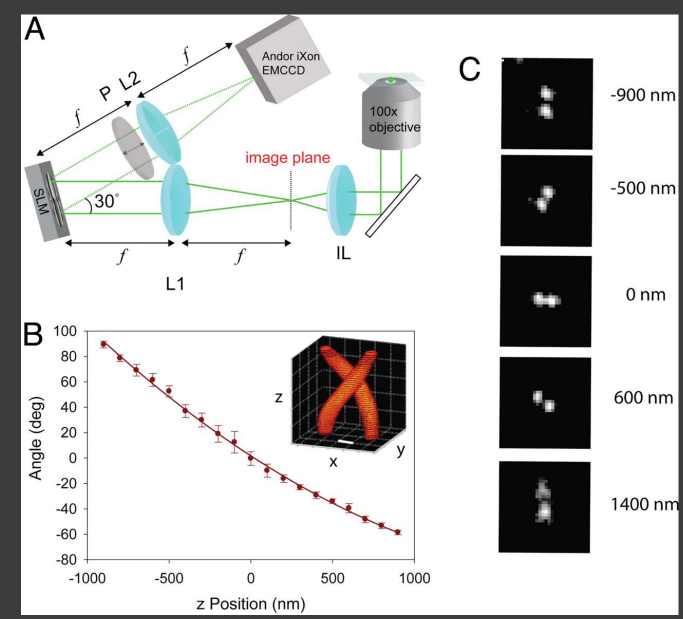
axial: 60 nm

#### biplane imaging

mtEos2-mitochondria

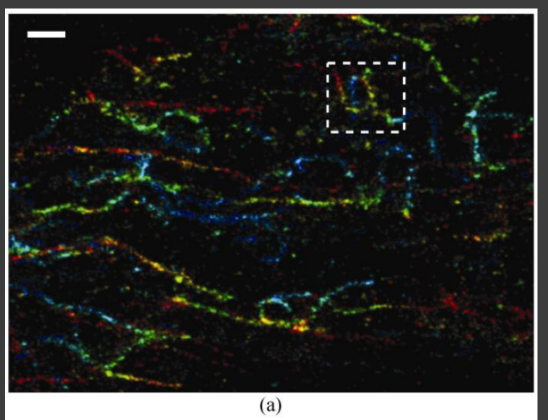


#### double helical PSF



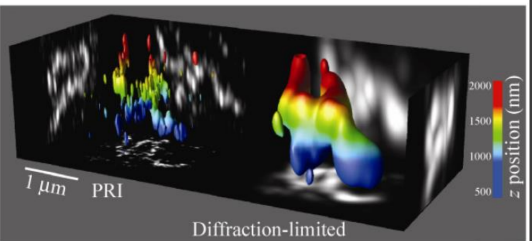
fitting of two 2D Gaussians

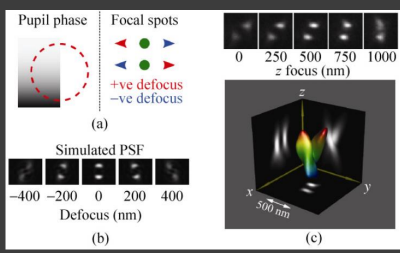
→ 3D position of the molecule

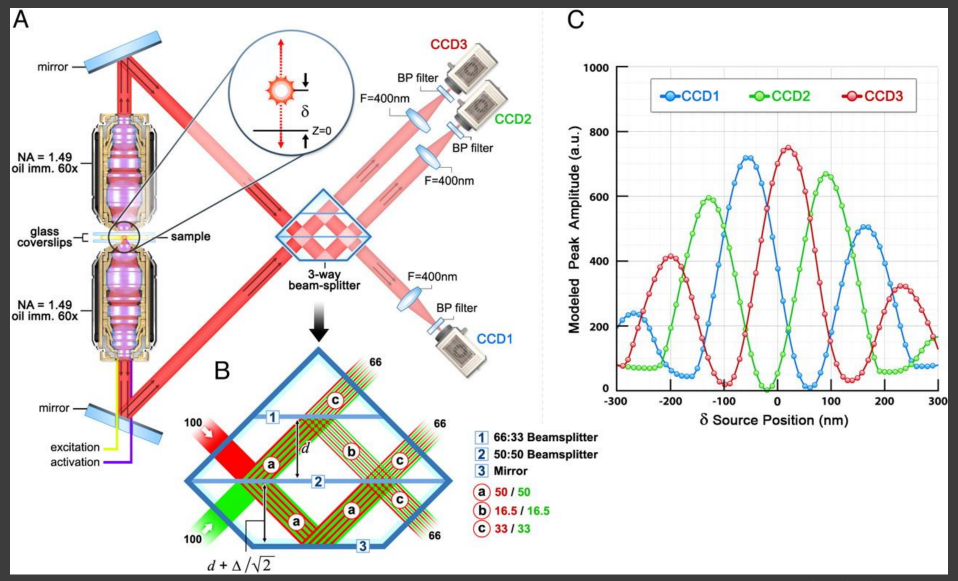


3D double helical PSF

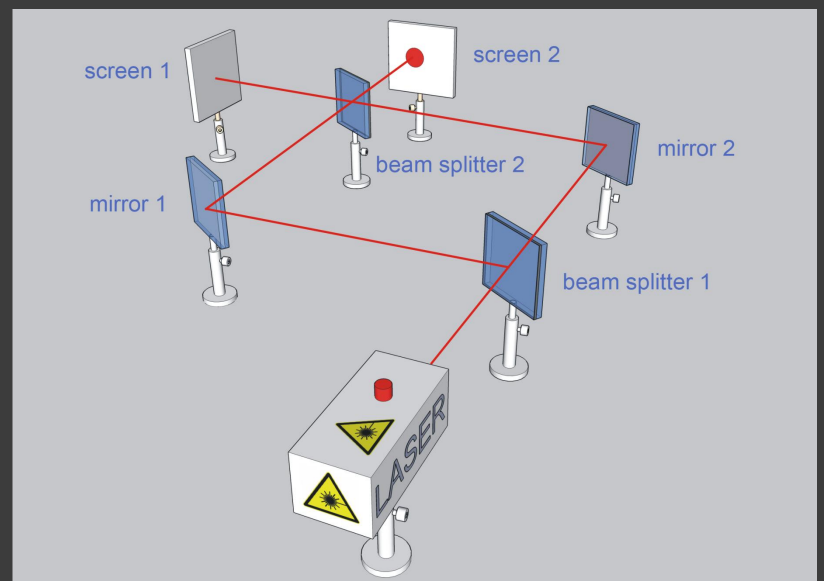
resolution xy: 30 nm z: < 100 nm

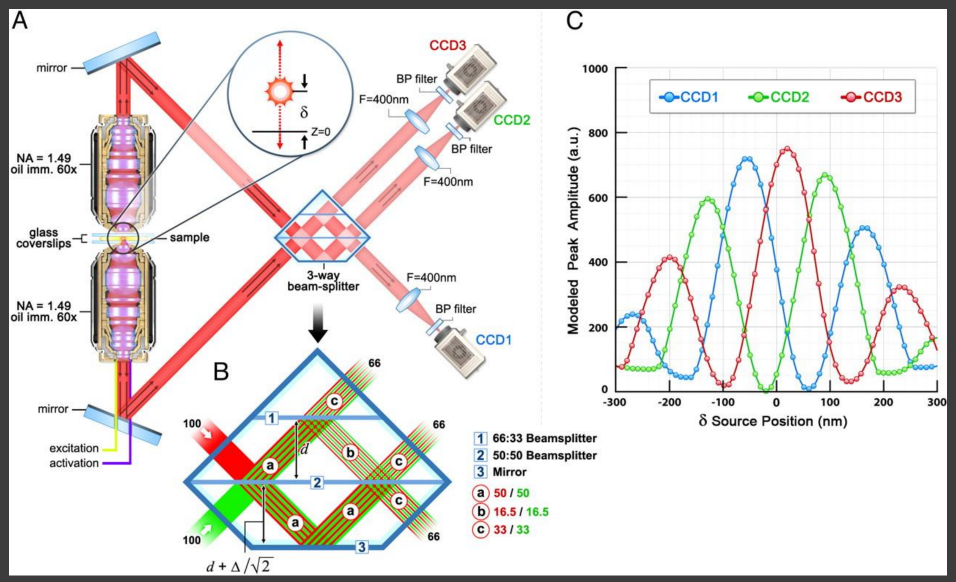


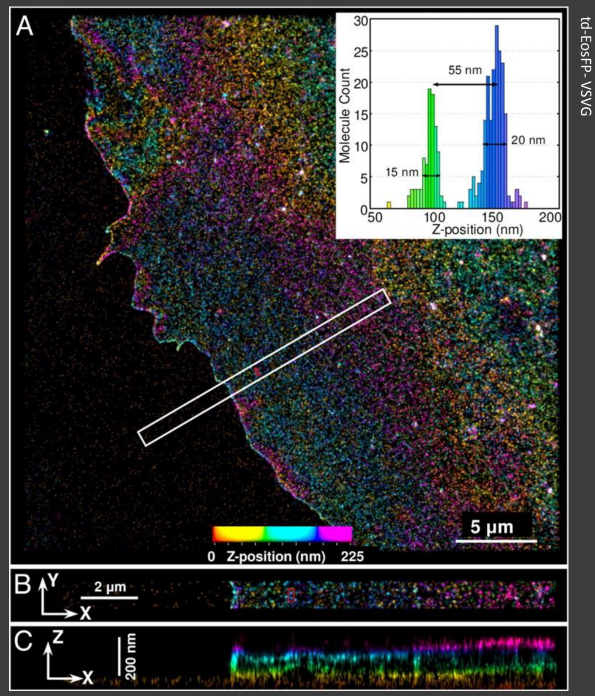




#### Mach-Zehnder-Interferometer





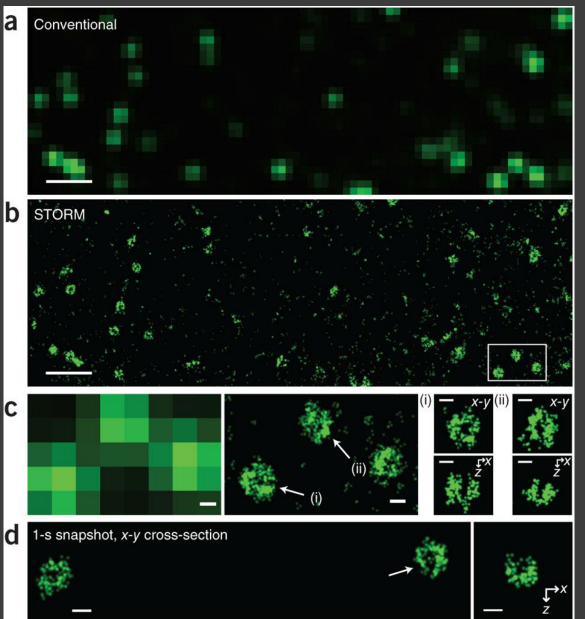


3D iPALM

resolution: 50 nm in all 3 directions

two examples for "live-cell" applications

## live-cell STORM (dSTORM)



### resolution

2D

spatial: 25 nm

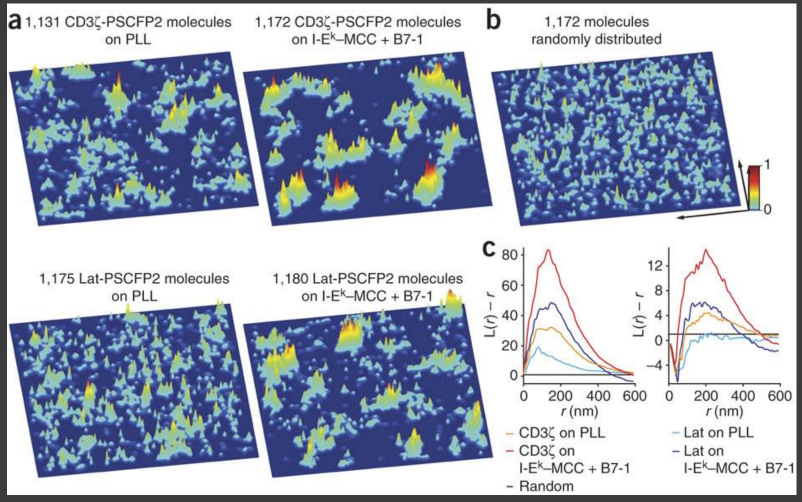
temporal: 500 ms

3D

spatial: xy: 30 nm, z: 50 nm

temporal: 1-2 s

### hsPALM



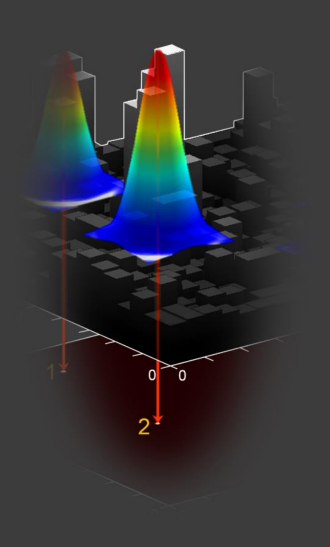
Lillemeier et al., Nature Immunology, 2009

2D, spatial resolution: 60 nm, temporal resolution: 4-10 s

how to get a lot more information from the data

## the additional single molecule information

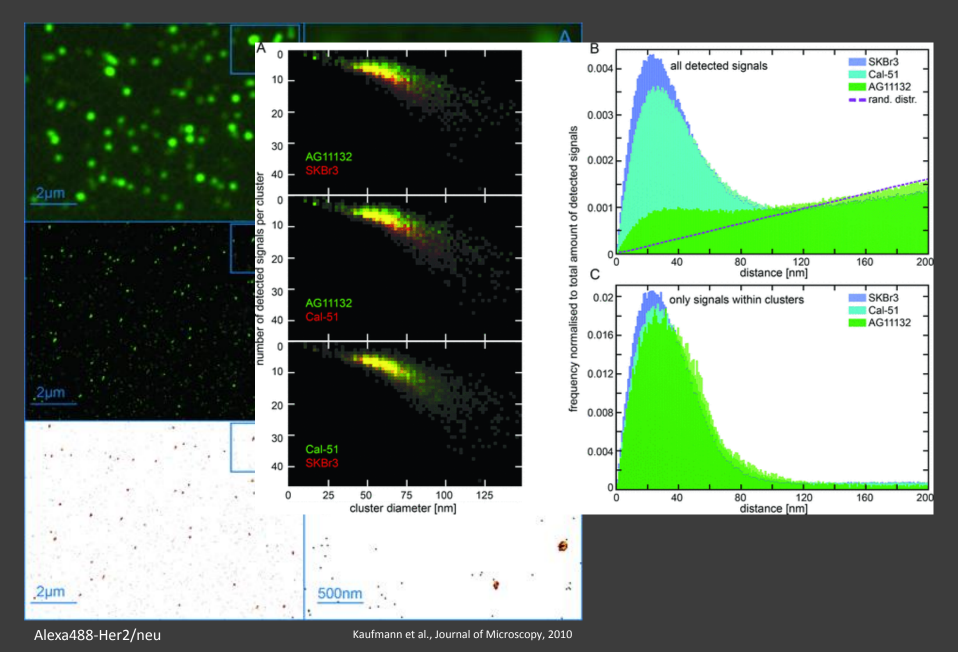
### remember?



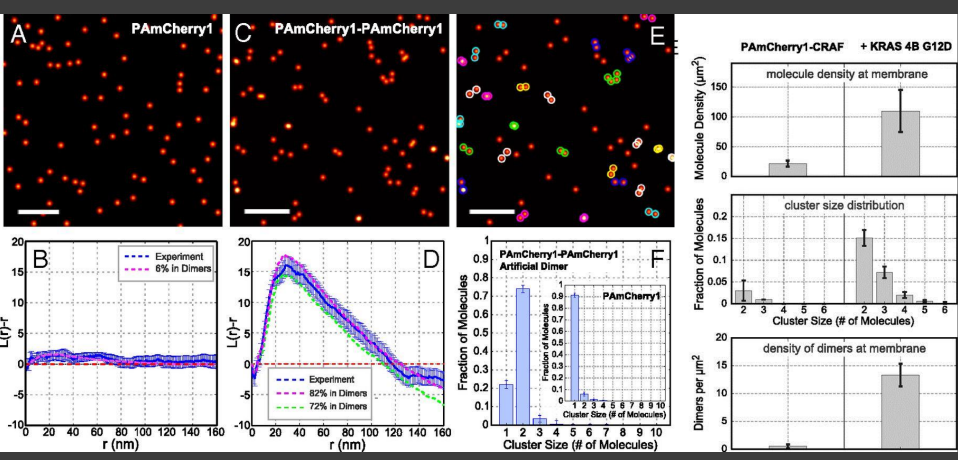
all the molecules in the image have been detected one by one

- → position of each molecule
- → number of detected photons
- → shape of the PSF
- → polarisation
- → wavelength
- → dynamics (in living cells)
- **→** ...

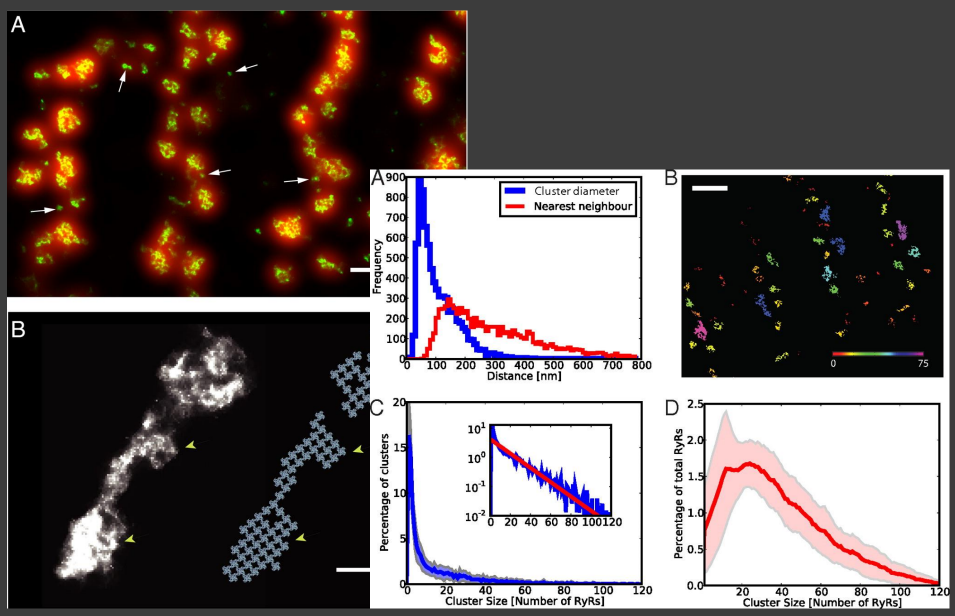
## statistical analysis of small protein clusters



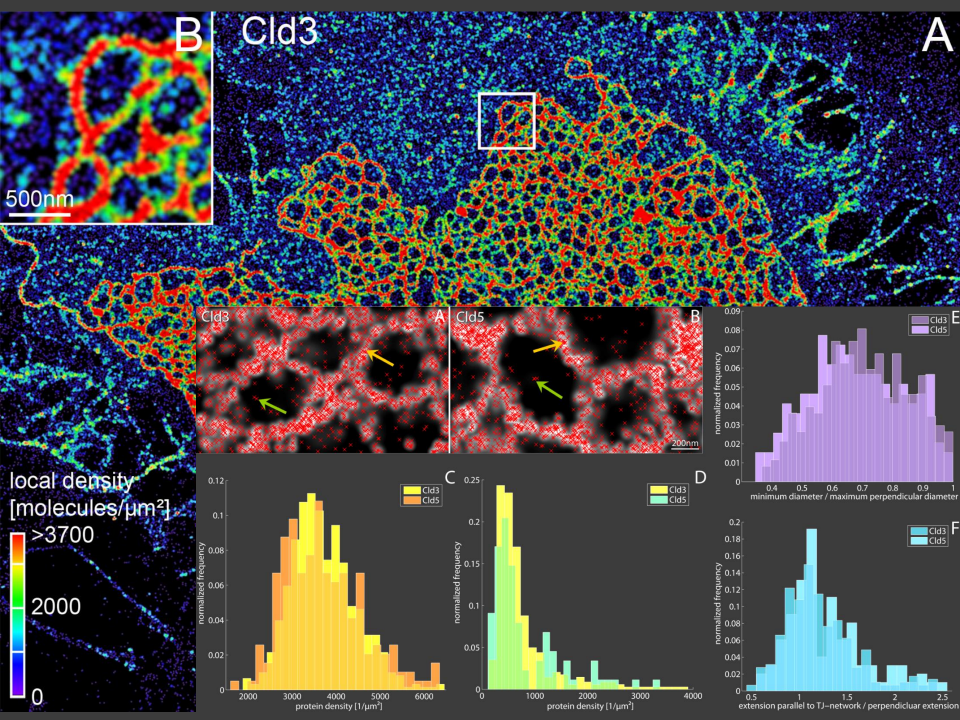
## analysis of protein clusters and molecule counting



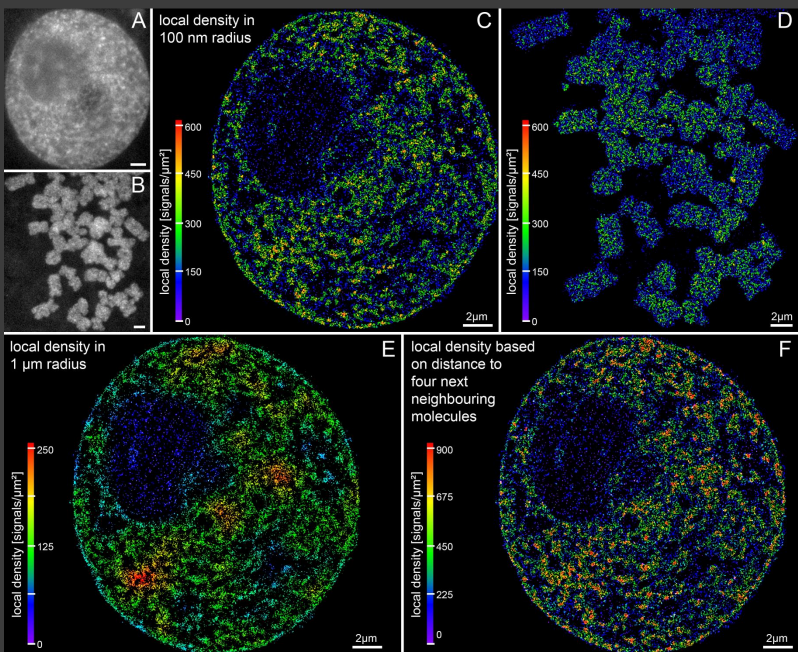
## statistical analysis of large protein clusters



Alexa647-RyR Baddeley et al., PNAS, 2009

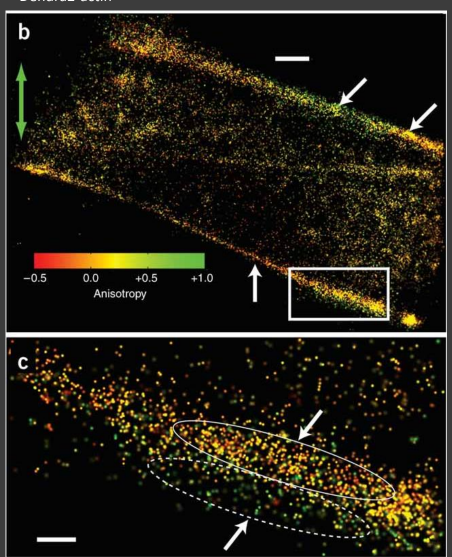


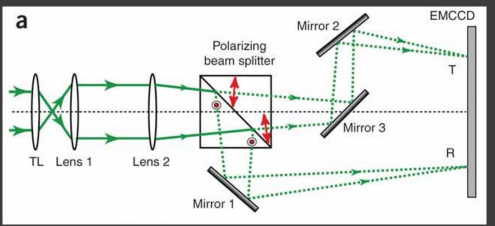
## visualisation of protein densities



## polarisation of the detected fluorophores

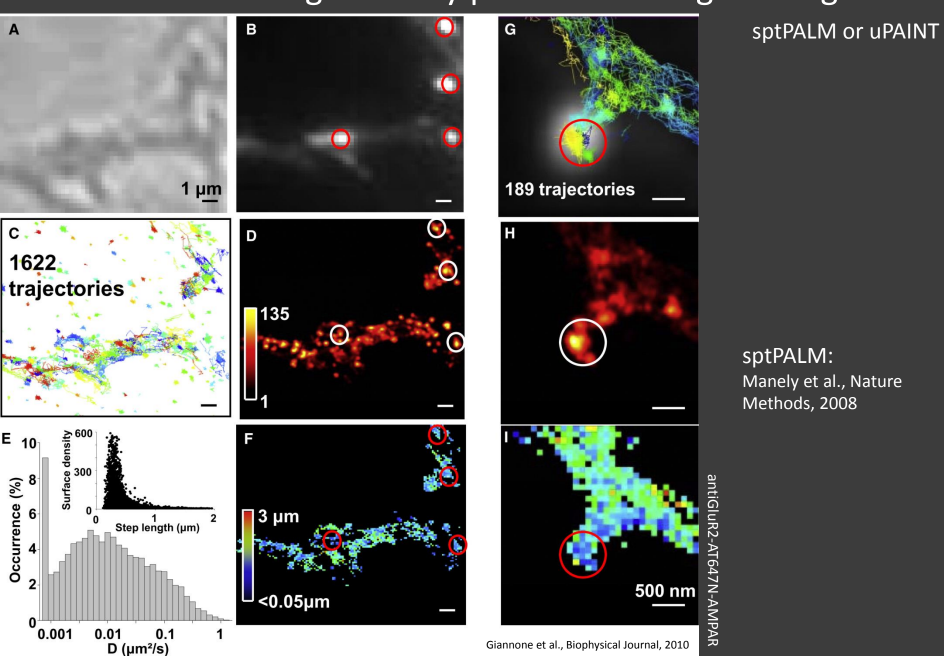
#### Dendra2-actin





Gould et al., Nature Methods, 2008

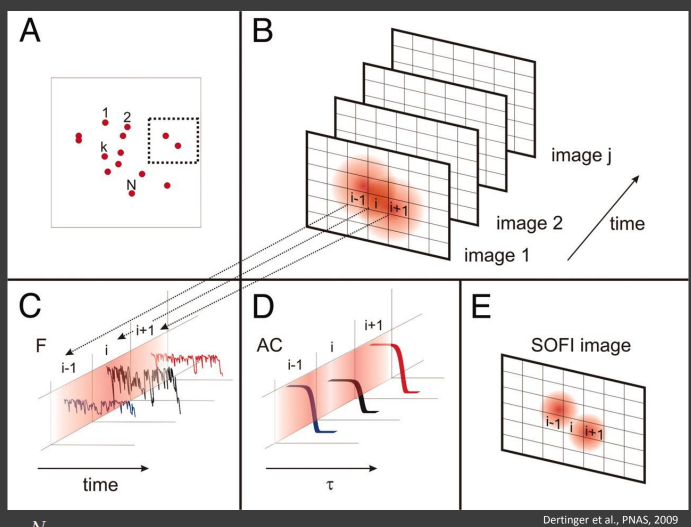
## high density particle tracking in living cells



# alternative approaches

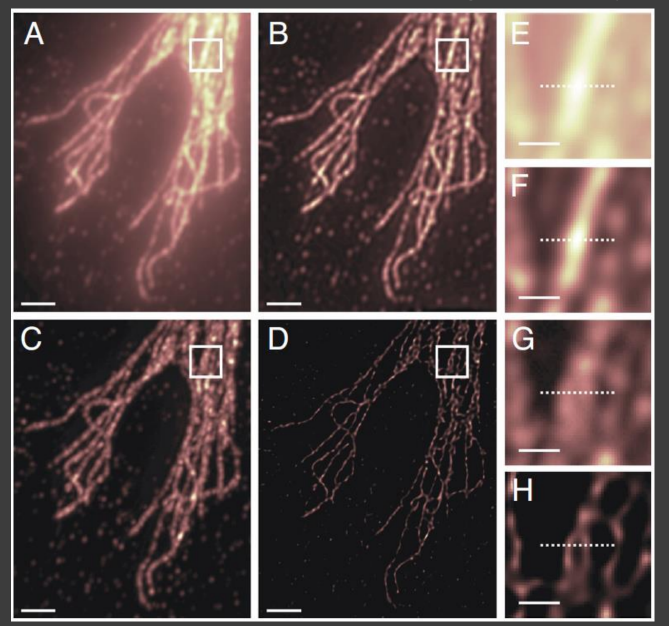
## SOFI - making the setup even more simpler

localization microscopy using a lamp!



$$\sum_{k=1}^{N} U(\mathbf{r} - \mathbf{r}_k) \cdot \varepsilon_k \cdot s_k(t) \qquad G_2(\mathbf{r}, \ \tau) = \sum_k U^2(\mathbf{r} - \mathbf{r}_k) \cdot \varepsilon_k^2 \cdot \langle \delta s_k(t + \tau) s_k(t) \rangle$$

# SOFI - making the setup even more simpler



lateral resolution: 70-100 nm

BUT!

no single molecule information

only resolution enhancement

Dertinger et al., PNAS, 2009

## 3B analysis localization microscopy

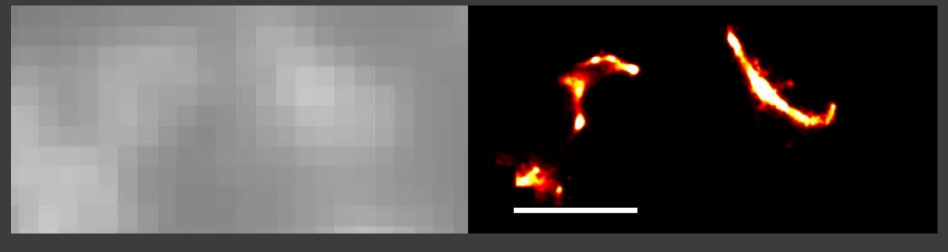
### similar approach as SOFI but some differences:

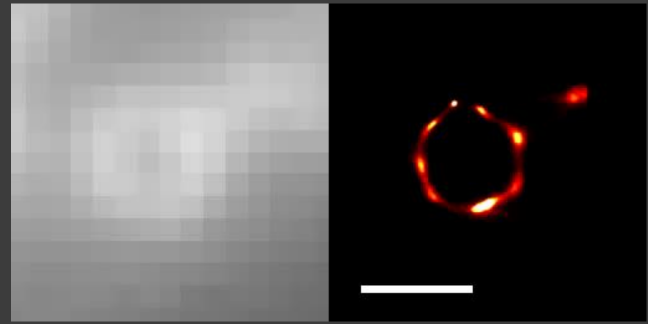
- + also based on very high molecule densities florescent in one frame
  - → very fast: only several hundred frames needed for reconstruction of an image with a resolution of 50 nm → time resolution: 4 s
- + single molecule information is still accessible
- extremely extensive computation effort
  - → regions larger than 2 x 2 μm would need to be processed for days on a conventional (core i7) CPU

## 3B analysis localization microscopy

wide-field

reconstruction (resolution: 50 nm)





Alexa488-podosomes

PALM: irreversible photo-activation

- quantitative analyses, particle tracking, counting
- → needs (in most cases) TIRF!

STORM, dSTORM, GSDIM, SPDM: reversible photo-switching

- → resolution, fast
- → also works without TIRF → imaging deeper inside cells

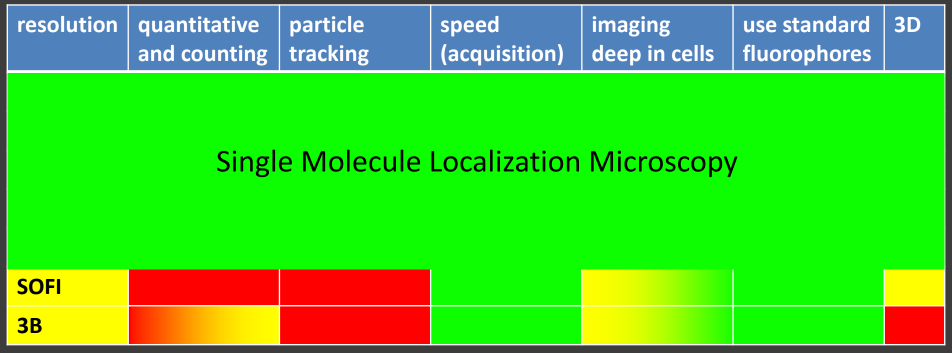
SPDM and GSDIM with FPs: (ir)reversible photo-switching

- → quantitative analyses using conventional FPs
- → also works without TIRF → imaging deeper inside cells

resolution	quantitative and counting	particle tracking	speed (acquisition)	imaging deep in cells	use standard fluorophores	3D
(F)PALM						
STORM						
dSTORM						
SPDM						
GSDIM						
SOFI						
3B						

referring to the original ideas of the methods

If you have a wide-field microscope with a laser for excitation of the fluorophores and one for switching/activating you can do ALL of these methods!



referring to the original ideas of the methods

If you have a wide-field microscope with a laser for excitation of the fluorophores and one for switching/activating you can do ALL of these methods!

### links

### original (F)PALM and STORM:

http://www.sciencemag.org/content/313/5793/1642.short http://www.nature.com/nmeth/journal/v3/n10/full/nmeth929.html http://www.sciencedirect.com/science/article/pii/S0006349506721403

### dSTORM, SPDM and GSDIM (with standard fluorophores):

http://onlinelibrary.wiley.com/doi/10.1002/anie.200802376/full http://www.springerlink.com/content/vx05p35kr3424228/ http://www.nature.com/nmeth/journal/v5/n11/full/nmeth.1257.html

#### 3D:

http://apl.aip.org/resource/1/applab/v97/i16/p161103\_s1?view=fulltext http://www.pnas.org/content/106/9/3125.short

### live-cell applications:

http://www.nature.com/nmeth/journal/v8/n6/abs/nmeth.1605.html http://www.nature.com/ni/journal/v11/n1/full/ni.1832.html

### statistical data analysis:

http://www.pnas.org/content/106/52/22275.short
http://onlinelibrary.wiley.com/doi/10.1111/j.1365-2818.2010.03436.x/full
http://www.plosone.org/article/info%3Adoi%2F10.1371%2Fjournal.pone.0031128

### links

### high density particle tracking:

http://www.nature.com/nmeth/journal/v5/n2/full/nmeth.1176.html http://www.sciencedirect.com/science/article/pii/S0006349510007137

#### Nat. Protoc.:

http://www.nature.com/nprot/journal/v4/n3/abs/nprot.2008.246.html http://www.nature.com/nprot/journal/v6/n7/abs/nprot.2011.336.html

### commercial systems:

http://zeiss-campus.magnet.fsu.edu/articles/superresolution/palm/introduction.html http://www.nikoninstruments.com/en\_GB/Products/Microscope-Systems/Inverted-Microscopes/Biological/N-STORM-Super-Resolution http://www.leica-microsystems.com/products/light-microscopes/life-scienceresearch/fluorescence-microscopes/details/product/leica-sr-gsd/

### algorithms:

http://www.super-resolution.biozentrum.uni-wuerzburg.de/home/rapidstorm/
http://code.google.com/p/quickpalm/

### summary and links:

http://www2.bioch.ox.ac.uk/microngroup/research/localization-microscopy.shtml