MULTIPHOTON MICROSCOPY

Methods for Cell Analysis Course BioVis – Uppsala, 2014

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SciLifeLab BioVis platform

Information

Information given here about 2 Photon microscopy were mainly taken from these sources:

<u>Background information on 2-Photon microscopy:</u> http://micro.magnet.fsu.edu/primer/techniques/fluorescence/multiphoton/ multiphotonintro.html

<u>The microscopes:</u> Zeiss LSM 710 NLO; http://www.zeiss.com Olympus Fluoview 1000 MPE, http://www.olympusamerica.com

<u>Spectra-Physics Laser:</u> http://www.newport.com/store/selectcountry.aspx?newpurl=/ Lasers/361887/1033/catalog.aspx

Schematic drawing of LSM



Why use 2-Photon microscopy ?



Multiphoton

LSM/ widefield





The message to keep in mind

A multiphoton microscope gives you the opportunity to get images from deep (e..g. 500 nm) within (living) tissue, whilst photodamaging only the imaged volume.

A Multiphoton microscope is a point scanning system which excites fluorophores within the Focus volume only. Therefore you collect emission light from this volume only, enabling you to acquire optical slices, without the use of confocal pinholes.

Beside this, one is able to photomanipulate tissue/cells within a very small volume.

THE THEORY OF 2PM



Theory for 2PM : The 1Photon Excitation

Excited Singlet States



- Illuminate a fluorophore with appropriate λ of light
- 1 (excitation) photon absorbed gives 1 emission photon

Stokes shift

- BUT
- emission photon will have less energy i.e.
 longer λ than excitation photon
 AND



- it's λ and energy vary due to which S₀ level (0,1,2,3) the fluorophore relaxes
- \succ Fluorescence photons with different λ emission curve is bell shaped

Theory for 2PM : $\lambda \sim E$ - The Energy of a Photon

nm	Energy	$F = hc/\lambda$ $eV = 1.6*10^{-19}$ I	eV	nm
1800			12,39	100
1600 -	h:	Planck Constant: 6,626*10 ⁻³⁴ J*s	6,20	200
	c:	speed of light: 299792 458 m/s	4,13	300
1400 -	λ:	wavelength in nm	3,09	400
1200 -	eV:	electron Volt: 1,6*10 ⁻¹⁹ J, gain of energy	2,47	500
	+	when an unbound electron is accelerated	2,06	600
1000 -		by an electrostatic potential difference of 1V	1,77	700
800 -			1,54	800
600		1n 400nm = 2n 800nm = 2 nV	1,37	900
800 -		1p 400mm – 2p 800mm – 3 ev	1,23	1000
400 -			1,12	1100
200			1,03	1200
200	↓ ↓		0,95	1300
0 +			0,88	1400
0,0	0 1,0 2,0 3,0	4,0 5,0 0,0 7,0 8,0 9,0 10,0 11,0 12,0 13,0 14,0	0,82	1500
		ev	0.74	1600

Theory for 2PM : How to excite (Tryptophan)

Single-photon 1 photon, 280 nm 4.5 eV No laser for this...

A

В

 $8\pi h v^3$

 c^{3}

Two-photon 2 photon, 580 nm 2.13 eV x2 4.26 eV

Three-photon 3 photon, 840 nm 1.47 eV x3 4.41 eV



2-PM hypothesis introduced by Maria Göppert-Mayer, doctoral thesis 1931

Theory for 2PM : \lambda \sim E - The Energy of a Photon



Observe: range of overlap of potential Excitation 760nm : excite A488 & A633 *

for multicolor 2PM choose fluorophores so that they do overlap in excitation BUT NOT emission

* has to be checked on microscope

Fluorochrome	Absorption		Emission	
Alexa Fluor 350	720-800		440	
Alexa Fluor 488	720-800		515	
Alexa Fluor 546	720-840		569	
Alexa Fluor 568	720-840		596	
Alexa Fluor 594	720-850		610	
Alexa Fluor 633	720-900		647	
AMCA	780-800		444	
bis-MSB	680-750		420	
Bodipy ear	900-950		512	
Calcium Crimson	etra	N	615	
Calcium green	780-850	red	531	
Cascade Blue	750-800		420	
Coumarin 307	780-800		530	
CY2	780-800		506	
CY3	780		565, 615	
CY5	780-820		670	
Dansyl Hydrazine	700-750		440	

Dealing with fluorescence in 2P



The 780nm NIR Laser might/will excite all three fluorophores, the Instrument has to unmix the mixture of Blue/Green/Red, or we have to use better fluorophore combination

Reminder – simultaneous vs sequential scanning





Simultaneous Excitation Resulted in artifact Due to bleeding through on "green" image,where the "blue" appears and on the red image where the "green appears"

Sequential scanning Does not show such Artifacts, therefore in THIS sample the Excitation are far apart.

Multicolor imaging in 2P



Simultaneous scan excites several fluorophore at once, emission is guided by filter and beamsplitter to PMTs. If FL-green bleed over into PMT of FL-red it will be seen here (in red). Sequential scan excites and collects one fluorophore at a time. ! Be sure that 488 does result in emission of FL red in the "green range"... Test that...



Lambda Scan with LSM – linear unmixing



Emission wavelength

430

537 547



Linear Unmixing determines the relative contribution from each fluorophore for every pixel of the image. And recalculates an image for Fluorophores used

02	
02	
0.1	
0.1	
0.1	
0.2	
17	
15	
50.5	
163.9	
255.0	
243.7	
10.6	
25.0	
109.3	
92.4	
77.0	
026	
635	
104.4	

WHY USE 2P? - to see deeper



Nikon instruments

See deeper – scattering problem

NIR light : 700-1100nm travelling through Specimen to focal plane will not scatter and disperse* as much as light of shorter λ (350-633 nm for FL microscopy)

excitation of fluorophores in greater depth

Problem: different fluorophores need its own NIR Laser?
Solution: Laser can be tuned from e.g. 690 to1040 nm, fluorophores have wide excitation range in 2PM

> *(due to different refractive indices of the various components in specimen) See also : Optical Clearing



specimen

Blue light gets easily scattered by particles. Otherwise Sinatra c/would nt sing " Blue skies, smilin' at me, nothin' but blue skies do I see"

See deeper – absorption problem



Tissue optical window: 700nm-900nm (absorption of hemoglobin/tissue component and water)

See deeper



XYZ images of mouse brain sections expressing GFP, comparing single-photon 488 nm excitation and two-photon 910 nm excitation.

With single photon excitation, tissue can be observed only to a depth of about 90 µm, but with two photons, observation to a depth of about 320 µm is possible.

Items displayed in color are vertical cross sections of 3-dimensionally constructed images.

Specimens provided by:

Kimihiko Kameyama, Tomoyo Ochiishi, Kazuyuki Kiyosue, Tatsuhiko Ebihara Molecular Neurobiology Group, Neuroscience Research Institute, National Institute of Advanced Industrial Science and Technology, Japan

Brochure, OLYMPUS, FV1000MPE

WHY USE 2P? - small excitation volume, no pinhole



Matyas Molnar

Small focus spot

Multiphoton LSM objective specimen excitation and emission focus events 2-photon classical microscopic system Ex~P_{avg} $Ex^{(P_{avg}/A)^2=I^2}$ That's why Multiphoton is also named Nonlinear. Chance for 2PM event drops drastically with distance to focus

Two-Photon event occurs only in focus volume

> All emission light is directly from focus

Resolution is similar (or worse) to LSM▶ 0.3x1µm ellipsoid (high NA objective)

Penetration depth depending on specimen and optical parameter but might be up to nearly 1mm

These features will be important for various live cell imaging techniques, like bleaching, photodamaging, uncaging ...

Small focus spot

Laser of LSM scans through specimen

Laser of 2PM scans through specimen



excitation/emission and photodamage/heat

occurs within specimen also outside the focal plane occurs within specimen only in the focal plane

Small focus spot

What is the chance that 2 photons hit the same fluorophore at almost the same time?

- a matter of time and area
- The probability of observing a two-photon absorption event on a bright sunny day is 1 per 10,000,000 years, whereas the one-photon absorption takes place every second
- <u>Time \rightarrow the virtual state</u>
- $\rightarrow \Delta t$ of intermediate virtual state = 10 attosec (10⁻¹⁷ s)
- ightarrow1 attosecond (10 $^{-18}$ s) is the time window
- \rightarrow light travels 3 hydrogenatoms within 1 attosec
- Area \rightarrow the fluorophore

 \rightarrow quite small target

Problem: Light can not travel faster than speed of light

Solution: More photons are needed (high density of

photons)

We need a million times more photons than in single photon fluorescence and "good" objectives



More photons please

Problem:

1 million times more photons? Very strong laser There is no continuous wave laser to achieve this.

Solution: A moderate Laser with high photon intensity pulses

- Iow average power (0.3 2.5 W)
- high peak power (30-300 kW) pulses 50-100 fs wide
- pulse frequency 80 Mhz (1pulse/ 12,5ns)

This laser is dangerous when used (Class 4)!

Problem: Many fluorophores but one Laser
Solution: To excite a wide range of fluorophores the laser is tuneable for e.g. 700-1040 nm

Pulsed NIR Laser is tuneable for excitation wavelength twice the 1Photon-excitation wavelength





Laser pulse is far from focal plane, photon density is low, no chance for two photons to hit a fluorophore in one time

Laser pulse is closer to focal plane, photon density is more concentrated but still low, no chance for two photons to hit a fluorophore in one time



Laser pulse reached the focal plane, photon density is high, high probability for 2 photons to hit one fluorophore within 10 attosec



The lucky ones emit fluorescence like they were hit by 1 high energy photon instead of 2 low energy photons

Excitation / emission occurs only in Focal plane /spot





Laser pulse leaves focal plane, NO incident of two photons hitting one fluorophore



Laser pulse disperses in tissue, NO incident of two photons hitting one fluorophore



REMEMBER

Excitation / emission occurs only in Focal plane /spot <u>confocal image</u> <u>without</u> <u>a pinhole</u>

Repeat again

Recapitulate:

- NIR Laser to reach deep

- Excitation of "normal" fluorophores via 2P effect

NIR is tuneable over range e.g. 690 nm – 1040 nm
 2P is only happening in focal volume
 -Ex/Em/photodamage only at focal volume

Applications:

Living animals

Manipulation of "precise" small volumes Non-linear effects

Relax...

Light travels 300.000.000 m / s
→ 7.4 x around earth in one s (40.075 km circumference)



circumference of human hair 53-565 µm

The Femtosecond Laser:

Pulses of 100fs

 $1 \text{ fs} = 1 \times 10^{-15} \text{ sec}^{-15}$

http://www.nsf.gov/od/lpa/news/03/images/wire_on_hair_big.jpg

http://antwrp.gsfc.nasa.gov/apod/ap070302.html

Multiphoton microscopy Objectives and Detectors

Light must come in to depth Light must get collected from the depth

Bring back home the photons

Laser \rightarrow <u>Objective</u> \rightarrow Excitation Emission \rightarrow <u>Objective</u> \rightarrow <u>Detector</u>



Multiphoton objectives



Long Working distance (2mm) including (!) High Numerical Aperture

(good resolution/focus,narrow depth of focus)



all photons to the focus for high chance of 2P-Ex

The Olympus XLPlan N 25x, NA 1.05

- High transmittance and correction for broad range of e.g. 400 nm to 1000 nm
- Water dipping (remember in vivo imaging) / cover slip
- Correction collar (!) to compensate for different refractive indices (water 1.3, specimen 1.34-1.4)
- 34 degree angle at lense top for better accessibility to specimen for manipulation



High NA + Long WD = expensive objective

Multiphoton detectors - NDD



Using the "long way" gives more flexibility, the confocal filterfree scanhead can be finetuned what range of light shall be collected, but the way is long (equals 32 cm glas!) and hence light is lost...

Using the NDDs as "short cuts" avoids loss of light. NDDs filter light via "old days" filtercubes and therefore lack in flexibility.

Multiphoton detectors - NDD

Loss of emission light: NDD vs LSMD I



Alexa 488, MaiTai 780nm, 5% (quite high), spectral range emission 500-550nm, no/open pinhole, digital gain etc for NDD (no over/under exposure)

Multiphoton detectors - GaAsP



With the very sensitive GaAsP detector right behind the objective we are able to collect more light from weakly fluorescent specimen (higher signal to noise ratio)
➢ one detector with no filter
➢ no distinction between different fluorophores...
➢ Efficience 40 % for 400 -700 nm

GaASP NDD

NDD

Loss of emission light: NDD vs GaAsP

Bring back home the photons - summary

FL emission is shorter in λ and get more scattered and dispersed than NIR Ex light
➢ Loss of emission light i.e. signal light

light gets lost via the optical pathways

Loss of emission light i.e. signal light

To compensate this loss Detectors should have

- better sensitivity
- proximity to specimen
- more





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Comparison of CLSM and 2P

	LSM	Multiphoton	
light source	laser UV to VIS	tuneable 50-100fs pulsed IR laser	
depth of visualization	up to 100 μm depending on tissue	up to 1000 μm depending on tissue	
XYZ resolution	via focal plane of objective, pinhole and wavelength	Similar (or worse) as LSM, No pinhole needed	
volume of exitation	throughout the Illuminated tissue	only the focal plane	
sensibility	Loss of signals via optics > Descanned detectors	Enhance signal by use of > Non-descanned detectors	

Go deeeeeeper....

Zeiss LSM 710 NLO



A method called Optical Clearing is available Making visualization depth of e.g. 1 mm possible Using light of 300 – 633 nm for excitation



Dendrites of cortical projection neurons of a transgenic mouse expressing YFP via the thy1 promotor. This high-resolution image of the dendritic processes to a depth of of 430 µm was made using multiphoton excitation of 920 nm in the living animal. Specimen provided by Stephen Turney, MCB, Harvard University, USA



Υ

layer 1

layer II

layer III

layer IV

3-dimensionally constructed images of neurons expressing EYFP in the cerebral neocortex of a mouse under anesthesia.

Cross-sectional images down to 0.7 mm from the surface can be observed after attachment of a special adapter to the specimen. Objective: LUMPlanFL 60xW/IR

Hiroaki Waki, Tomomi Nemoto, and Junichi Nabekura National Institute for Physiological Sciences, National Institutes of Natural Sciences, Japan



Optical clearing



Removing the optical barriers (different RIs) makes the object invisible – transparent

- Left: water (n=1.3) and glass rod (n=1.5)
- Right: oil (n=1.5) and glass rod (n=1.5)

Optical clearing





Problem:

Biological tissue : poor light transmission due to interface lipid:water (PM :in/ex-cellular fluids) Solution:

Replace aqueous fluids with solvents which matches Refractive Index (RI) of lipids.

- Penetration of light into the tissue increases,
- Scattering of light decreases.

Optical Clearing Agents (OCAs) : aromatic hydrocarbons

- water insoluble but soluble in EtOH or MetOH.
- each clearing is preceded by dehydration (Et/MetOH)
- benzyl-alcohol-benzoate (BABB) (excellent)
- Methyl salycylate (wintergreen oil) (very good)
- Thiodiethanol (TDE) (good)
- Glycerin (poor clearing)
- OCAs have usually a refractive index of around 1.5, hence matching RI of glass, and immersion oil.

Optical clearing – go deeper

PBS 60x wi PBS 40x oil Glycerol BABB TDE 50µm **Depth** into tissue 100µm 150µm 180µm

Appleton et al, Journal of Microscopy, Vol. 234, Pt 2 2009, pp. 196-204

THANKS FOR YOUR ATTENTION!

