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Multiplex-multiphoton microscopy and computational strategy for biomedical imaging

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Abstract

We demonstrate the benefit of a novel laser strategy in multiphoton microscopy (MPM). The cheap, simple and turn-key supercontinuum laser system (SCLS) with its spectral shaping module, constitutes an ideal approach for the one-shot microscopic imaging of many fluorophores without modification of the excitation parameters: central wavelength, spectral bandwidth and average power. The polyvalence of the resulting multiplex-multiphoton microscopy (M-MPM) device is illustrated by images of many biomedical models from several origins (biological, medical or vegetal), generated while keeping constant the spectral parameters of excitation. The resolution of the M-MPM device is **quantified** by a procedure of point-spread-function (PSF) assessment led by an original, robust and reliable computational approach FIGARO. **The estimated values for the PSF width for our M-MPM system are shown to be comparable to standard values found in optical microscopy.** The simplification of the excitation system constitutes a significant instrumental progress in biomedical MPM, paving the way to the imaging of many fluorophores with a single shot of excitation without any modification of the lighting device.

Research Highlights

A new solution of multiplex-multiphoton microscopy device is shown, resting on a supercontinuum laser. The one-shot excitation device has imaged biomedical & vegetal models. Our original computational strategy measures usual microscopy resolution.

I. Introduction

The assets of multiphoton microscopy (MPM) are now well-established in biomedical sciences, especially thanks to its near infrared range (NIR) of excitation, less dangerous and less absorbed than UV or visible excitation light (Diaspro, 2002; Mertz 2009). In routine use, standard commercial multiphoton systems include a tunable mode-locked titanium-doped sapphire laser excitation (Ti: Sa, 150 fs, 80 MHz, 700-1000 nm) delivering a 10 nm-spectral bandwidth at the full-width at half maximum (FWHM). This laser technology has proved its skills in MPM and has been overwhelmingly implemented on commercial MPM systems (Girkin et al., 2005, Lefort, 2018). In case of suited emission thresholds, a spectral detection is optimal for imaging many fluorophores with a single shot of excitation. Otherwise, photomultiplier tubes (PMT) are better adapted but the simultaneous detection is limited to four fluorophores at best, providing a multishot excitation strategy. Indeed, the remote multiphoton excitation spectra of standard fluorophores are spread on large bandwidths covering about 300 nm in the NIR, sometimes without any overlapping. This property limits the dynamic imaging of samples across different species which can only be alternatively imaged with a multishot excitation (Boppart et

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3 al., 2019). Then, a numerical merging of resulting images is mandatory for obtaining a multiplex image
4 and artefacts due to misalignment during the recording can appear. This sequential procedure
5 drastically increases imaging delay, a critical parameter in the field of dynamic multiplex imaging or for
6 detecting phenomena at video rates (Boppart et al., 2019; Kirkpatrick et al., 2012; Abdeladim et al.,
7 2019). The technological transition from standard multishot MPM toward a one-shot strategy of
8 excitation for multiplex imaging thus still requires the implantation of an alternative excitation solution
9 for MPM.
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12 More confidential excitation solutions based on a supercontinuum laser system (SCLS) have existed in
13 MPM for fifteen years (Palero et al., 2005, Tada et al., 2007, Li et al., 2009; Tao et al., 2011; Tu et al.,
14 2016; Cui et al., 2017; Magnol et al., 2018; Eibl et al., 2018; Poudel et al., 2019). Such solutions are
15 ideal for multiplexing the excitation beam thus resulting in a so called instrument of “multiplex-
16 multiphoton microscopy” (M-MPM) system. Exogenous fluorophores such as enhanced genetically
17 modified proteins (GFPs) have been imaged with a SCLS (Yokoyama et al. 2007; Yokoyama et al. 2008;
18 Hou et al. 2011). But in the aforementioned works, SCLS are mostly generated by the spectral
19 broadening of Ti: Sa pulses into a photonic crystal fiber (PCF), a strategy complex technically, expensive
20 and not compatible with a turn-key use, mandatory for routine practice of M-MPM. Nonetheless, a
21 large spectral bandwidth of excitation is closely associated to the question of chromatic effects. This
22 setting is at the origin of an increase of the noise and blur level in the images. This can be detrimental
23 to a precise quantification for the PSF instrument, though necessary for efficient restoration
24 procedure. PSF assessment is a procedure well-established in optical microscopy which consists in: (1)
25 imaging standardized microspheres smaller than the sought resolution, (2) numerically modeling the
26 resulting bead images by 3D-Gaussian shapes and (3) estimating the resulting dimensions using the
27 FWHM along the 3 dimensions. Nevertheless, available numerical solutions are often characterized by
28 high sensitivity to noise and model restrictions (e.g. assuming zero background, or limited to 1D / 2D
29 treatments of image projections), which may make them inefficient in the context of M-MPM PSF
30 characterization (Zhu et al., 2013; Guo, 2011). An alternative strategy named Fourier ring correlation
31 (FRC) provides an estimate of the FWHM of the PSF (Koho et al. 2019), resulting in a PSF model
32 consisting in an isotropic Gaussian. This is characterized by a single FWHM value considering the main
33 directions of the PSF aligned with the X, Y or Z axis, which is not always experimentally true.
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38 In this paper, we demonstrate the benefit of an original, cheap, simple and turn-key SCLS coupled with
39 a spectral shaping module, devoted to biomedical MPM. The resulting M-MPM system is first
40 dedicated to the characterization of multiphoton excitation spectra of about ten different biomedical
41 endogenous fluorophores. Then, several models of application from life sciences are imaged with this
42 new device and its one-shot excitation strategy. With this novel M-MPM system, no modification of
43 the lighting device is required, regardless of the sample origin: biological, medical, chemical or vegetal.
44 The diversity of application models imaged without technical modification of the lighting device
45 emphasizes a significant simplification of the current routine use of MPM. The resolution associated
46 to this new excitation strategy is quantified by PSF assessment. Our original computational approach
47 FIGARO has been especially elaborated for M-MPM devices, likely to be confronted with deleterious
48 chromatic effects on the effective performance of the instrument. This computational solution delivers
49 a complete and precise 3D PSF estimation, robust to noise and anisotropic 3D blur effects which are
50 not negligible with a wide spectral excitation (Lau et al., 2018; Chouzenoux et al., 2019). The estimated
51 values for the PSF width for our M-MPM system are shown to be comparable to standard values found
52 in optical microscopy. Thus, the one-shot excitation strategy with a unique spectral bandwidth
53 removes all the needs for each sample imaged to adapt the central wavelength, spectral bandwidth
54 and average power of laser beam. Thereby, our original M-MPM system combining a SCLS with a robust
55 computational strategy leads to a substantial instrumental simplification in modern biomedical
56 imaging.
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II. Multiplex-multiphoton microscopy (M-MPM) in biomedical imaging

1. Experimental setup of M-MPM

Figure 1 summarizes the overall proposed M-MPM optical setup combining 4 technical modules numbered from A to D. The excitation device (Figure 1A) is composed by a SCLS which results from the spectral broadening of a monochromatic mode-locked seed laser centered at 1064 nm generating 5 ps pulse durations at 40 MHz, into a photonic crystal fiber (PCF) with a 6 μm core diameter. The seed laser peak power of 10 kW generates nonlinear processes into the PCF such as four-wave mixing, soliton effects and Raman scattering. This leads to the spectral broadening of the monochromatic spectrum from 410 nm up to 2.4 μm with a total average power of 2 W. At the fiber output, a parabolic mirror collimates the beam. A density filter manages the excitation average power before its injection into the spectral shaping module composed of a prism-line which plays the role of spectral selection. The resulting SCLS is spectrally and temporally characterized (Figure 1B) and then implemented on a standard configuration of upright multiphoton microscope stand (Figure 1C) from Thorlabs Inc. (Bergamo® II Series). Two PMTs with adjustable sensitivities, combined with spectral filters, detect the endogenous fluorescence emitted from the sample. The overall resolution estimation is performed by our original algorithm FIGARO for PSF assessment (Figure 1D).

The prism-line is composed of two prisms of SF57 flint dispersive glass from SCHOTT Advanced Optics, with an apex angle equal to 30°. An anti-reflection coating is centered at 900 nm on each effective surface. The first prism spatially disperses wavelengths; the second one collimates them. The distance d between prisms, here equal to 4.7 cm, and the incident angle ϑ of the beam on the first prism can be adjusted manually. This system governs the spatial spreading of wavelengths on the reflective element of the prism-line. In the current configuration proposed in Figure 1A, the back reflection mirror contains the entire spectrum spatially spread on 3.8 cm. It is preceded by two independent slits moveable with adjustable width that can be shifted all along the spectrum. With such method, a diffraction effect is visible on the spectrum when the slits are shifted. The resolution, measured with the knife-edge method (Siegman et al., 1991), equals 4 nm. **If the excitation window required is located further in the infrared range, the moveable slits can be translated correspondingly.**

2. Spectro-temporal characteristics of excitation pulses for MPM

The spectro-temporal characterization of excitation pulses was performed using a spectrometer (Ocean Optics, USB 2000+) and an autocorrelator (APE, Mini), positioned directly at the prism-line output (Figure 1B). Figure 2 presents the spectral and temporal characterizations of pulses measured at the prism-line output.

By increasing the distance between the two slits, central wavelength was tuned from 780 nm until 900 nm as shown in Figure 2A. For each spectral bandwidth broadened from 60 nm until 150 nm at FWHM, the corresponding autocorrelation trace was recorded and shown in Figure 2B. Figure 2C illustrates spectral bandwidth measurements at the FWHM and corresponding pulse duration as a function of the central wavelength highlighting femtosecond ranges. The NIR of excitation generated by our original excitation strategy combined with its range of pulse duration fits ideally with optimal spectro-temporal requirements of MPM. Therefore, our SCLS and its shaping module cumulate the best spectral properties for M-MPM of numerous fluorophores regardless of the biological origin of samples thanks to the cheapest, simplest and fastest managing protocol.

3. Endogenous multiplex multiphoton fluorescence excitation spectra of biological substances

Living samples contain some structuring biochemical substances which play the role of endogenous fluorophores. Their presence at the intracellular level and in the extracellular matrix is at the inception of the label-free imaging ability of MPM. As an example, Table 1 lists non-exhaustively a few of these fluorescent label-free substances and their origins. We can notice here that a unique fluorophore can be excited thanks to either a one-, two- or three-photon absorption process provided an adapted wavelength range of excitation.

Table 1. List of endogenous fluorescent substances from life sciences.

Kingdom	Samples	Molecules	Substances
Animals	Organs (lung, arteries, muscle...)	Proteins	Elastin, Collagen, Myosin
	Cells	Proteins (<i>enzymes</i>)	NADH, FAD
		Amino acids (<i>in proteins</i>)	Tryptophan, Phenylalanine, Glycine, Tyrosine...
Plants	Stem / leaves	Organic substances	Cellulose, Lignin, Chlorophyll
	Flowers / leaves	Proteins (<i>enzymes</i>)	NADH, FAD
	Cells	Amino acids (<i>in proteins</i>)	Tryptophan, Phenylalanine, Glycine, Tyrosine...

We now determine experimentally multiphoton excitation spectra of the most relevant autofluorescent substances in biomedical researches from small molecules such as amino acids to macromolecules such as proteins and organic substances. The experimental protocol proposed in (Kao et al., 2004) has been followed: each biological sample was composed of a unique extracted or synthesized solid state substance (micro-crystal or powder), placed on a microscope slide imaged with the setup presented in Figure 1. The excitation bandwidth delivered by our SCLS was fixed at 10 nm at FWHM and central excitation wavelength was tuned between 680 and 1000 nm with a step of 10 nm and a constant average power equal to 5 mW on the sample. Table 2 lists the substances tested and the spectral range of the emission filter associated. Each bandpass filter selected for this set of experiments has a spectral bandwidth of several tens of nanometers. This solution is adapted for the characterization of individual fluorophores. In a less favorable situation where several fluorophores are involved simultaneously with some overlapping emission regions, spectral detection could be favored, thus avoiding a risk of spectral crosstalk at the detection. This last solution presents however few limitations: the point-by-point spectral acquisition requires a longer acquisition duration and is less adapted in case of low signal to noise ratio emission. An intermediary solution consists in choosing emission filters detecting a peripheral part of the emission spectra, in ranges spectrally not overlapped, at the price of a weaker detected intensity and potentially degraded by noise. In the current situation, for each point of each spectrum, ten measurements were performed. The resulting excitation spectra were then normalized and averaged. Figure 3 shows the multiphoton fluorescence excitation spectra of these substances as a function of excitation wavelength.

Table 2. Multiphoton processes and detected spectral emission ranges of main endogenous fluorescent biological substances.

Substance	Multiphoton process	Detected range of fluorescence
Elastin, Flavin (FAD), NADH	Two-photon fluorescence	495-540 nm
Glycine, Tyrosine, Collagen	Two-photon fluorescence	420-500 nm
Tryptophan	Three-photon fluorescence	260-340 nm

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4 These seven biological substances highlight multiphoton fluorescence excitation spectra spread on 230
5 nm, between 690 and 920 nm. No spectral overlapping of multiphoton fluorescence excitation exists
6 between several of these substances such as amino acids and metabolic indicators. The one-shot
7 imaging of fluorophores presenting such a spectral gap is not possible technically with a standard Ti:
8 Sa system having a 10-nm spectral bandwidth of excitation at the FWHM. For optimal excitation
9 conditions, the one-shot imaging of these seven substances would require an excitation spectrum
10 continuous on the range between 680 and 920 nm.
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16 4. Square and cubic dependences of multiphoton fluorescence

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18 The law dependence of fluorescence intensity emitted in function with excitation average power
19 deposited on the sample was measured for each fluorescent biological substances (Table 2). The average
20 power was mastered with an optical density. The spectral properties remained constant during the
21 whole experimental session: a central wavelength fixed at 900 nm and a spectral width of 150 nm at
22 the FWHM; the excitation spectrum and the pulse duration were checked for each average power and
23 remained not significantly modified. Figure 4 shows the logarithmic plots of the normalized
24 fluorescence intensities measured for elastin and tryptophan in function with the logarithmic plot of
25 the average power. The quadratic behavior is observed in Figure 4 for elastin and demonstrates the
26 two-photon process involved; a similar quadratic evolution, not shown in Figure 4, was observed for
27 FAD, NADH, glycine, tyrosine, and collagen. The cubic evolution of fluorescence intensity depending
28 on average power for tryptophan demonstrates the three-photon process involved.
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32 5. Multiplex-multiphoton images of samples from life sciences

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34 In this section, we demonstrate the adaptation of our original excitation strategy to generate
35 multiphoton images of several models of application from different fields of life sciences. The spectral
36 properties delivered by our system remained constant during the whole experimental session and
37 regardless of the sample imaged. Its central wavelength was fixed at 900 nm with a spectral width of
38 150 nm at the FWHM. The resulting excitation beam was thus implemented into our microscope stand
39 for generating multiphoton images of different models from life sciences. Figure 5 presents the
40 multiphoton images generated with this excitation beam. Two kinds of samples were imaged, one from
41 the plant kingdom (douglas wood) and one from the animal kingdom (mouse kidney). The origin of the
42 fluorescence signals was determined thanks to two information: the spectral range of emission and
43 the spatial localization of the emitted signal on the image. For douglas wood, the endogenous
44 fluorescence detected between 420 and 500 nm was located on the image from the inner part to the
45 peripheral of the wood cell wall; the fluorescence detected between 604 and 678 nm was located in
46 the peripheral region of the wood cell wall. The yellow signal represents the overlapping regions of the
47 wood cell wall. The spatial organization presented in Figure 5A-C respectively corresponds to the lignin
48 and cellulose localizations (Thomas et al. 2013, Donaldson et al. 2018). For the kidney sample,
49 extracellular matrix components, flavins and lipofuscin (Zhang et al., 2018) were imaged (Figure 5D-F).
50 Their two-photon fluorescence signals were detected respectively between 604 and 678 nm and
51 between 500 and 550 nm. The TPA spectra of these four different fluorophores ranged between 690
52 and 950 nm. We notice here that no photodamage characterized by a reduction in fluorescence levels
53 was identified all along the period of image recording. For both application model, the selection of the
54 excitation wavelengths with the SCLS from 750 nm to 850 nm has switched off the signal emitted from
55 the red channels (between 604 and 678 nm). Reversely, the selection of the excitation wavelengths
56 from 850 nm to 1000 nm has switched off the signal emitted from the blue channels (between 420 and
57 550 nm). The simultaneous generation of the red and blue fluorescent signals for each application
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3 model was only possible thanks to a broad excitation spectrum covering a spectral range from 750 to
4 1000 nm.
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7 8 6. Discussion about M-MPM in life sciences 9

10 Figure 5 highlights the production of multiphoton fluorescent images of application models from plants
11 and animals kingdom with a unique and unchanged excitation spectrum. Central wavelength, spectral
12 bandwidth and average power were kept constant all along the experimental period. For the first time
13 and regardless of the samples origin, the imaging of many fluorescent substances was achieved
14 without any modifications on physical lighting parameters, thanks to the original and easy handled
15 one-shot excitation strategy (Figure 1).
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18 The multiplex and one-shot excitation strategy offered by the original M-MPM system overcomes
19 several technical constraints associated to the standard excitation strategy of MPM. First, M-MPM is
20 associated with the smallest time delay required to generate multiphoton signals. As described in
21 (Diaspro et al. 2002), according to a basic principle of optics, the focal position is wavelength
22 dependent. This phenomenon has a straightforward consequence resulting in an axial offset between
23 focal plans recorded with excitation wavelengths spectrally pulled apart. With a standard multishot
24 excitation, this trouble often occurs and is associated to a variable image blur between focal plans. The
25 correction of this drawback requires a preliminary procedure of quantification which is not standard
26 and depends on the microscope objective and the sample medium. The image blur, not always easy to
27 detect or to correct during the recording or numerically after the image merging, results in a mismatch
28 between positions of the same object. The SCLS, generating images all along the focal spot, allows the
29 adjustment of focal position in live for each spectral window detected. The axial correction of the focal
30 position become realizable before recording the image, without modification of the excitation
31 parameters. Second, regardless of the sample origin, the one-shot excitation approach removes the
32 constraints about central wavelength selection and wavelength dependence of average power. With
33 a Ti: Sa laser, the wavelength adjustment is a time consuming procedure, associated to a wavelength
34 dependent average power. The disparity in average power of a Ti: Sa laser can reach a factor 20
35 between its maximum at 800 nm and its minimum at 1000 nm. Such variations can flawed quantitative
36 measurements and can be harmful for sample integrity if no careful attention is paid on the average
37 power mastering. Finally, thanks to the one-shot excitation strategy, users do not need any more to
38 adjust the lighting device for each experiment.
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44 The use of a single and fixed large band excitation spectrum, regardless of the biomedical model
45 imaged from life sciences, represents a significant headway in the simplicity of use, reduced cost and
46 dynamic performance offered by multiphoton microscopes. This instrumental simplification drives
47 MPM towards dynamic M-MPM. This new device is associated to a new progress in biomedical imaging
48 and in a new routine practice of microscopic characterization of biomedical models.
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53 III. Computational strategy for resolution assessment 54

55 A large spectral bandwidth of excitation might be associated with chromatic effects that could be
56 detrimental for image resolution. We quantify here the resolution performance thanks to a 3D PSF
57 assessment procedure. Several numerical solutions of PSF estimation exist today in optical microscopy,
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the most popular being the commercial software “Huygens”¹ by “Scientific Volume Imaging” having a multiphoton module, and the free plugin of Image J “MetroloJ”². We have recently developed a novel computational strategy named FIGARO, implemented as an ImageJ plugin³, especially dedicated to the robust 3D resolution assessment of M-MPM.

1. Principle and interest of an original computational strategy: FIGARO algorithm

We have used our original computational strategy FIGARO, delivering a PSF fitting directly applicable to the 3D-images of normalized and calibrated fluorescent microspheres. This strategy achieves a trade-off between accuracy and simplicity governing the success of PSF estimation. Mathematically, an observed raw data crop, containing a single microsphere, is stacked into a vector y . The fitting process is then based on a computational estimation of a noiseless, scaled and centered version p of y with a multivariate Gaussian shape, and scaling/shift/center parameters a , b , and μ respectively, at each voxel position $x \in \mathbb{R}^3$. More precisely, the corresponding entry p is assumed to be close to $a + b g$, where g is the multivariate Gaussian probability density function defined by Equation 1.

$$\forall x \in \mathbb{R}^3, g(x, \mu, C) = \sqrt{\frac{|C|}{(2\pi)^3}} e^{\left(-\frac{1}{2}(x-\mu)^T C (x-\mu)\right)} \quad (1)$$

Here, μ denotes the mean of the Gaussian distribution, i.e. the center of the PSF in our case, while C is a 3x3 symmetric positive definite precision matrix which characterizes its width and orientation, $|C|$ being its determinant. For an optimal description of the raw observed data y , the key point of our numerical strategy relies on a fitting of parameters (a, b, p, μ, C) . This fitting task is performed using FIGARO algorithm. It is worth noting that our PSF fitting procedure does not require any pre-processing of the dataset, as it performs the denoising and fitting task jointly, assuming a Gaussian additive noise model, whose variance level is estimated automatically. The mathematical description, and numerical validation of FIGARO is detailed in our preceding publications (Lau et al., 2018; Chouzenoux et al., 2019).

Figure 6 displays the FIGARO results, starting from the image of a single microsphere cropped from native image volume. Figures 6A and 6B represent 2D projections of the acquired bead profile marginalized in XZ and XY plans, respectively. The FWHM (bleu ellipses) and center (red dots) of the PSF are computed from the μ and C values estimated with FIGARO algorithm. Figures 6C to 6F present the FIGARO intensity profiles allowing to quantify the instrument PSF.

2. Experimental PSF estimation

The experimental resolution estimation of our M-MPM device is now evaluated, using our computational strategy FIGARO. We also display comparisons with the two standard PSF estimation solutions: MetroloJ and Huygens. In that purpose, yellow-green fluorescent microspheres with a 0.2 μm diameter of Invitrogen™ (reference F8811) were mounted between a microscope slide and a coverslip. Such objects serve usually as a reference standard for the calibration of the green channel of laser scanning microscopes and emit a fluorescence spectrum between 500 and 545 nm at the FWHM with a maximum peak at 515 nm. **With the same protocol as involved in section II.2, we have**

¹ <https://svi.nl/HuygensSoftware>

² <https://imagejdocu.tudor.lu/plugin/analysis/metroloj/start>

³ https://imagejdocu.tudor.lu/plugin/analysis/figaro_psf_3d_optical_microscopy/start

determined the multiphoton excitation spectrum of these microspheres. At the FWHM, the related excitation spectrum covers the range of wavelengths between 870 and 960 nm. The excitation beam from our SCLS were centered at 900 nm with a spectral bandwidth of 150 nm at FWHM. The 3D images were generated thanks to the XY scanning device coupled with the z-axis scan (Figure 1). The resulting 3D native image was composed of a stack of 2D squared images of 4096×4096 pixels, for a total field of view equal to $163.84 \mu\text{m} \times 163.84 \mu\text{m}$. The pixel size on the image equal to $0.04 \mu\text{m} \times 0.04 \mu\text{m}$, cumulated with a z-stack having a $0.1 \mu\text{m}$ step-size on a total depth of $10 \mu\text{m}$. These settings were adjusted in order to combine an accurate acquisition of PSF image, being constituted by about ten points along each of the 3 dimensions, associated with an acceptable fitting duration of few minutes. All images were loaded in tiff format with ImageJ and 6 individual microspheres exploitable for PSF measurements were manually cropped from the native image. They were then analyzed for PSF estimation with three different computational strategies: MetroloJ, Huygens and FIGARO. Table 3 gathers the resulting PSF parameters values in the three directions of interest X, Y and Z.

Table 3. PSF measured from 6 microspheres in the three dimensions X, Y and Z. Three strategies compared: MetroloJ, Huygens and FIGARO.

Bead number	MetroloJ			Huygens			FIGARO		
	PSF X (μm)	PSF Y (μm)	PSF Z (μm)	PSF X (μm)	PSF Y (μm)	PSF Z (μm)	PSF X (μm)	PSF Y (μm)	PSF Z (μm)
1	0.004	0.038	0.006	0.304	0.310	7.82	0.300	0.323	2.275
2	0.109	0.038	1.38	0.510	1.078	1.969	0.364	0.674	2.572
3	0.129	0.136	1.688	0.328	0.313	12.564	0.205	0.249	2.85
4	0.04	0.038	0.009	0.356	0.343	4.063	0.599	0.458	2.627
5	0.007	0.038	0.007	0.339	0.340	6.736	0.459	0.275	0.969
6	0.106	0.022	0.051	0.306	0.320	3.529	0.334	0.681	2.572
Average Value	0.07	0.05	0.5	0.4	0.4	6.1	0.4	0.4	2.3
Standard deviation	0.05	0.04	0.7	0.08	0.29	3.5	0.13	0.18	0.6

3. Discussion about PSF estimation and related computational strategies

The resolution performance is an essential parameter in microscopy, quantified by a standardized practice of PSF estimation. The values, resulting to a protocol of PSF estimation, seem to highly depend on the mathematical procedure being used. Here, three different PSF estimation procedures have been compared for the evaluation of the resolution performance of our M-MPM. For each PSF estimation, the value of standard deviation has been calculated.

The standard methods of PSF assessment with “MetroloJ” and “Huygens”, highlight guarded and highly dispersed results of resolution. Those values illustrate the limits of the underlying mathematical strategies in those two tools, which are facing their limits in the situation of M-MPM. Mathematically, both Huygens and MetroloJ process the 3 directions X, Y and Z separately, using standard curve-fitting strategy, and thus obtain only 1D characterizations of the PSF along those 3 directions. This procedure is not precise enough, and unsuitable for noise levels and highly anisotropic blur associated to M-MMP.

A relevant PSF estimation strategy must consider 3D data and deal with essential mathematical properties such as guaranties of convergence and background not assumed to be equal to zero. Our computational strategy FIGARO, especially devoted to M-MPM resolution estimation, has specifically integrated these properties into its mathematical protocol. With this strategy, the resolution estimation equals $0.4 \mu\text{m}$ in the X and Y directions and $2.3 \mu\text{m}$ in the Z direction. Our computational strategy has highlighted its strong relevance for the analysis of PSF data coming from M-MPM, with a

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3 low dispersion, and has demonstrated a better resistance to noise and blur than standard strategies.
4 Our FIGARO strategy allows to distinguish the signal coming specifically from microspheres to the noise
5 generated by the large spectral bandwidth of excitation. The estimated resolution values were in
6 adequacy with standard resolutions of optical microscopes (Diaspro et al. 2002): a planar sub-
7 micrometer resolution (optimal value theoretically estimated at 361 nm) and an axial micrometer
8 resolution (optimal value theoretically estimated at 1.58 μm). Our M-MPM device thus does not suffer
9 from chromatic aberrations more than standard optical microscopy systems, emphasizing lateral and
10 axial resolution performance correctly preserved compared to standards usually encountered with a
11 monochromatic excitation.
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14 V. Conclusion

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16 Multiphoton microscopy (MPM) is a technical solution well-adapted for dynamic imaging or for
17 revealing complex biomedical structures. But the quasi-monochromatic excitation systems usually
18 implemented in commercial multiphoton microscopes limits at four the quantity of fluorophores that
19 can be imaged simultaneously at best or imposes a multishot imaging approach of each individual
20 fluorophore of interest. Such a sequential procedure is time-consuming and associated with several
21 engineering locks, likely corrupting image information. In that context, a large spectral bandwidth of
22 excitation has gained interest. Implemented on a multiphoton microscope, this new excitation strategy
23 results into a so called multiplex-MPM (M-MPM) system. In this paper, we have highlighted an original
24 engineering solution of M-MPM based on a simple, cheap and reliable supercontinuum laser source
25 coupled with a passive spectral shaping module. This one-shot excitation strategy turns out to provide
26 a new solution with temporal performance favorable to generate multiphoton processes. The
27 polyvalence of the resulting M-MPM system was illustrated by images of biomedical models from
28 several origins (biological, medical or vegetal) while keeping constant the spectral parameters of
29 excitation. To characterize the resolution performance of this new device, we have developed an
30 original and novel computational strategy. This approach, especially devoted to M-MPM devices, rests
31 on a rigorous mathematical approach based on the more recent and sophisticated methods of
32 numerical estimation. A preserved optical resolution was shown compared to standard optical
33 microscopy systems. Henceforth, many new perspectives are now opened in dynamic biomedical
34 microscopy thanks to M-MPM associated with a robust computational strategy.
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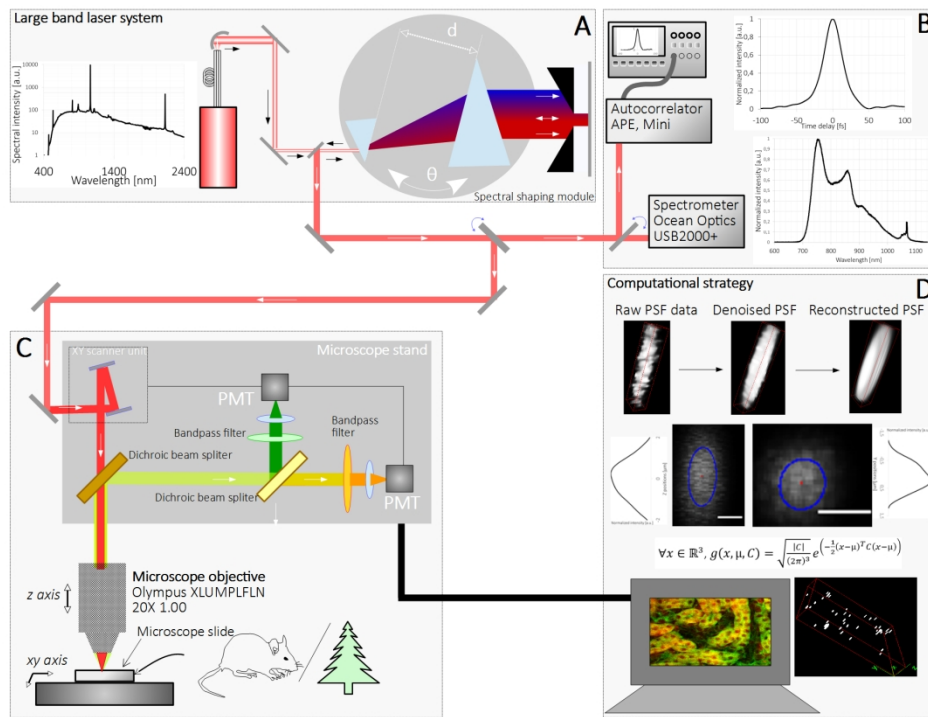
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Experimental setup of the M-MPM device with a SCLS and its spectral shaping module (A, B) for multiphoton imaging of samples from life sciences (C) combined with our original computational strategy for PSF assessment (D).

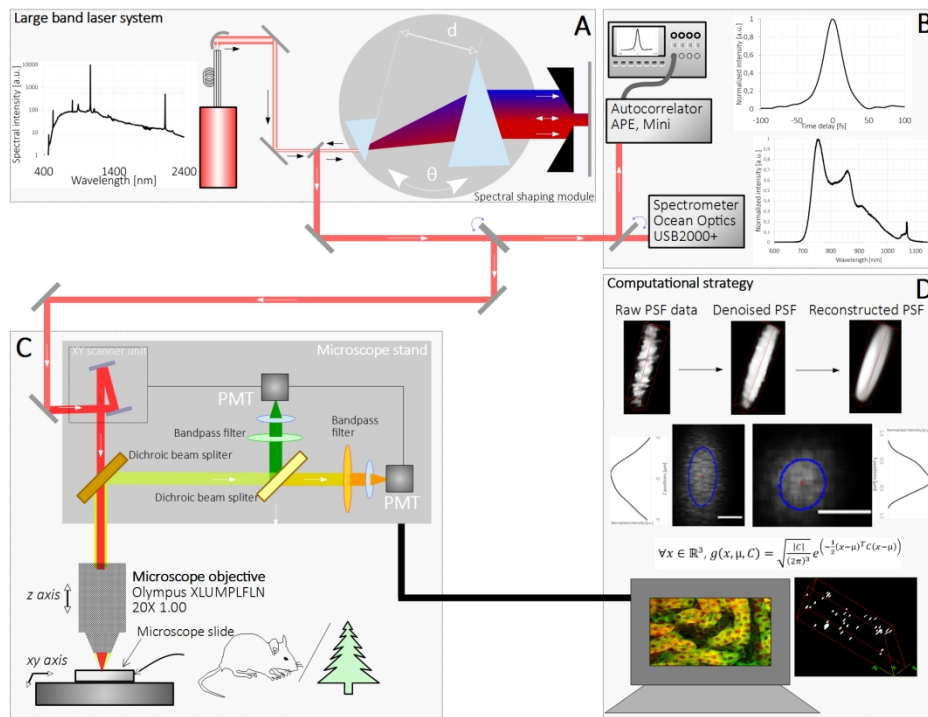


Figure 1. Experimental setup of the M-MPM device with a SCLS and its spectral shaping module (A, B) for multiphoton imaging of samples from life sciences (C) combined with our original computational strategy for PSF assessment (D).

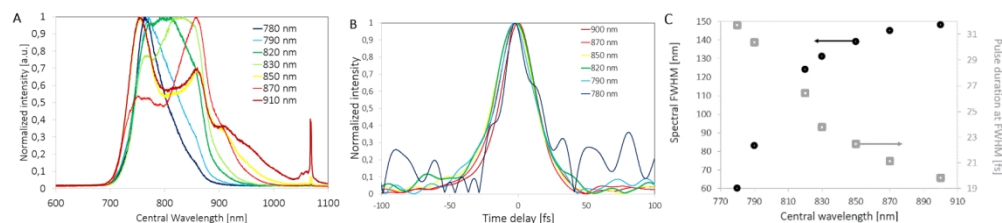


Figure 2. A. Spectral shape and B. Temporal shape of pulses central wavelengths between 780 and 900 nm and spectral bandwidths at FWHM between 60 and 150 nm. C. Spectral bandwidth at the FWHM (nm) and related experimental pulse duration (fs) as a function of central wavelength (nm). For each plot, an arrow points to the corresponding ordinate scale.

638x154mm (72 x 72 DPI)

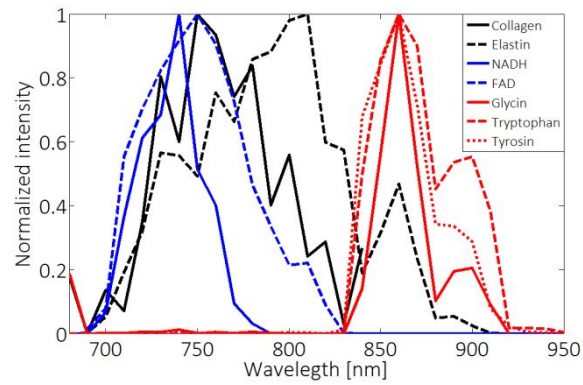


Figure 3. Experimental measurements of multiphoton fluorescence excitation spectra of endogenous fluorescent biological substances of interest in biomedical researches as a function of excitation wavelength: proteins (collagen, elastin), enzymes (NADH, FAD) and amino-acids (glycine, tyrosine and tryptophan).

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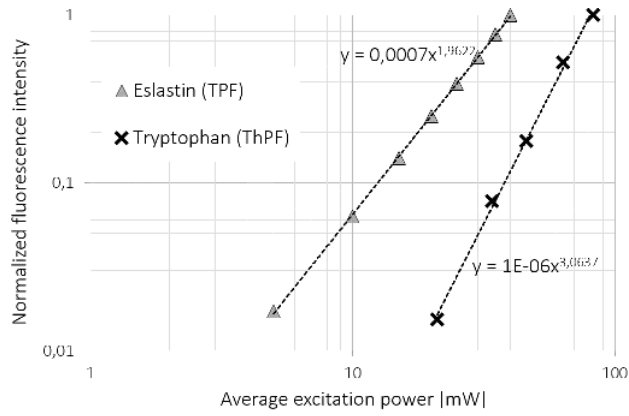


Figure 4. Normalized fluorescence intensities in function with average excitation power for elastin and tryptophan. Square and cubic dependences of multiphoton fluorescence are highlighted.

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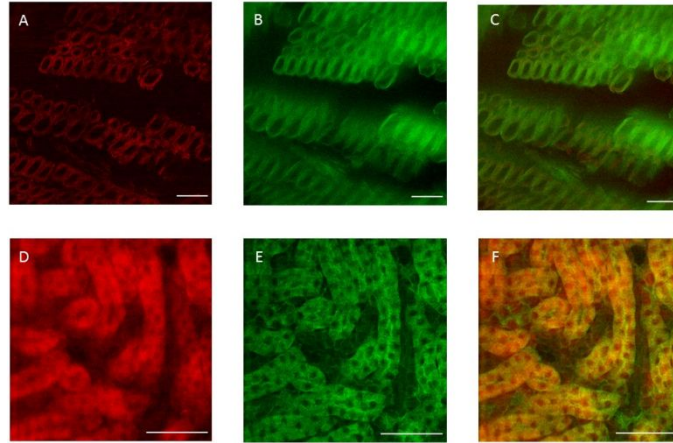


Figure 5. Illustration of multiphoton images of several biological samples obtained with a unique large band multiphoton excitation centered at 900 nm with a spectral bandwidth of 150 nm at FWHM. Images with 2048×2048 pixels were acquired with a pixel dwell-time of $3.6 \mu\text{s}/\text{pixel}$. A to C: Endogenous fluorescent Douglas wood detected between 604 and 678 nm (A), between 420 nm and 500 nm (B), and merged (C). D to F: Kidney autofluorescence detected between 604 and 678 nm (D), and between 500 and 550 nm (E), and merged (F). Scale bar: $50 \mu\text{m}$.

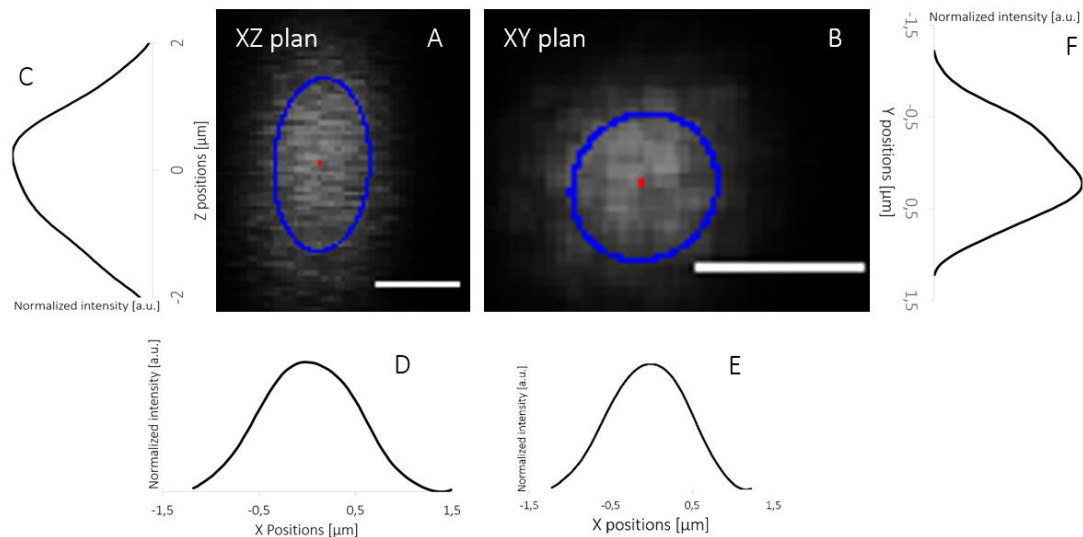


Figure 6. PSF characterization of our M-MPM device. Images of yellow-green fluorescent microspheres (diameter: 0.2 μm, detection range 500-550 nm); excitation centered at 900 nm, 150 nm at FWHM. A and B. XZ and XY 2D image of a single microsphere after estimating PSF with FIGARO plugin. Red dots and blue ellipses localize respectively the bead center and the computed FWHM of intensity with an analytical Gaussian model. C and D. PSF intensity profile in XZ plan. E and F. PSF intensity profile in XY plan. Scale bars: 1 μm.