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ABSTRACT

Multiphoton microscopy (MPM) is an imaging method involving a near infrared range of excitation. However, generating 3D microscopic images in high depths with a satisfying quality is complex due to physical distortions of the beam (optical aberrations and diffraction limit) along the optical path. The ability of computational approaches to restore reliably deep 3D images relies on the accuracy of the mathematical estimation of optical distortions introduced by the instrument. Instrumental distortions of the beam are characterized by the point spread function (PSF) evolution, especially in the optical axis and whose model can be fitted by a Gaussian shape. In this publication, we present our approach which relies first on the design of an optical phantom constituted by standardized microspheres, immobilized into a 3D volume of 2 mm deep. Then, 3D-volumes of images containing a single object are selected and isolated all along the sample depth using automatic morphological tools. Finally, our computational 3D-Gaussian shape fitting algorithm named FIGARO is applied on each individual PSF. With this novel pipeline, FWHM evolution of MPM has been measured for the first time in the 3 dimensions along a 2 mm depth. We have highlighted a highly significant effect of spherical aberrations in our MPM system, showing an increase with a factor 4 of the axial PSF dimensions in the depth while preserving the lateral PSF dimensions.

Keywords: Multiphoton microscopy, deep 3D-imaging, PSF evolution

1. INTRODUCTION

Three-dimensional (3D) imaging has been developed since 30 years for biomedical applications [1]. Imaging the whole organ in its physiological environment is of first interest for delivering correct diagnostics without any resection protocol and with a painless procedure. Magnetic resonance imaging (MRI) is probably the most known 3D and painless medical imaging method highly employed in hospitals. But the resolution of MRI is limited to the millimeter range and some details at the scale of the cell or at sub-cellular scale can be missed and can distort the end diagnostic. 3D imaging, in vivo and in live, with a sub-micrometer resolution for biomedical applications is still a current necessity for biomedical researches and corresponds to a hot topic for biomedical imaging engineers [2, 3].

Optical microscopy is the de facto solution for 3D biomedical imaging potentially in vivo and associated to a micrometer resolution. Today, the gold standard in optical microscopy is confocal microscopy where the samples imaged have to be labeled with fluorophores for revealing their presence [4]. This method is invasive due to the required labeling process and may be not the best solution for 3D and in depth imaging due to the use of UV or visible ranges of excitation. Such excitation range can be absorbed or scattered by the constituents of the imaged sample, which reduces the ability of such excitation wavelength to generate a deep image of biological targets. Moreover, UV or visible ranges are highly energetic, and can finally damage the imaged sample.
Multiphoton microscopy (MPM) appeared in 1990 as an alternative to confocal microscopy. MPM is a microscopy method based on a process of fluorescence emission similar to confocal microscopy. The specificity of MPM relies on the excitation strategy, based on the simultaneous interaction of two photons at least on the substance imaged. The 2 photons involved cumulate their energy for generating similar fluorescence emission to those involved in confocal microscopy. The two-photon approach opens the access to near infrared range (NIR) of excitation. Wavelengths in the NIR are less energetic, less absorbed and less scattered than UV and visible wavelengths. By consequence, NIR has a deeper penetration depth than UV and visible ranges and deep 3D and in vivo microscopy of biomedical samples become accessible [5]. Moreover, the simultaneous presence of two (or more) photons of excitation with the fluorophores is a phenomenon of low probability. This characteristic results in an optical sectioning inherent to MPM [6].

MPM can reveal the structure of biomedical samples until several millimeters depth depending on the optical properties of the imaged sample, and especially its transparency to wavelengths in the NIR. However, optical microscopy and MPM are optical methods sensitive to physical phenomena distorting the excitation beam: diffraction limit and chromatic aberrations. These physical phenomena can be reduced thanks to computational strategies. Indeed, each microscope device has its own transfer function, also named Point Spread Function (PSF), whose consequences are harmful on the image quality [7]. Typically, optical distortions transform a Dirac signal into a Gaussian signal and the thinnest Gaussian shape is associated to the best resolution. The PSF effect, resulting in a blur in the acquired images, raises an instrumental and mathematical challenge.

In this context, several computational strategies have been developed especially for the numerical compensation of optical distortions in MPM. Most of them are based on the idea that, as soon as the PSF of the system is known, its contribution to the image distortion can be reduced thanks to mathematical inversion methods such as deconvolution. Many plugins (for example, in ImageJ software) are proposing tools for a basic restoration of images based on the deconvolution of the measured image by the experimental PSF. In this paper, we present an original pipeline with the aim to illustrate and quantify the non-stationarity of the PSF in MPM, and in particular the evolution of its width along the optical axis. The outline is as follows. First, the interest of computational strategies in MPM is reminded. Then, we present our main contribution, which is a novel computational strategy for the quantification of the axial PSF evolution in MPM. Finally, the illustration of the impact of axial PSF distortions is highlighted on images of muscle sample. We present an illustration of reconstruction of these images of muscle samples.

2. INTEREST OF COMPUTATIONAL STRATEGIES IN MPM

2.1 Evaluation of the resolution

MPM is an optical solution for the microscopic imaging of biological samples involving a minimally invasive procedure based on second and third order light-matter interactions requiring NIR of excitation. However, the use of wavelengths in the NIR is associated to deleterious physical effects of light diffraction and chromatic effects whose impact on the image quality concerns essentially the resolution. In optical microscopy, the resolution is an essential parameter whose quantification is fixed by standard protocols. Considering that a Dirac signal is transformed into a Gaussian shape due to the specific transfer function of the optical microscope, the resolution estimation amounts to measuring the width of this Gaussian signal. Experimentally, the Dirac signal is reproduced by fluorescent microspheres having a diameter smaller than the optical resolution. The spheres fluorescence goes through the optical microscope which applies its specific transfer function and acquisition parameters. By this procedure, the image of a microsphere becomes an image of the spread function; this transformation is thus named point spread function (PSF). Figure 1 illustrates the definition of the PSF and its impact on the images generated in optical microscopy.
The true signal of the object, named $\bar{x}$ on Figure 1, passes through the instrument which induces a convolution represented here by the linear operator $H$ associated with an acquisition noise $w$. This convolution results in the observed volume $y$. By consequence, the observed signal is the convolution between the specific transfer function of the instrument and the original volume $\bar{x}$. Specific computational strategies can be developed in order to remove or reduce the instrumental contribution to the image. The main strategy consists in computing a deconvolution of the image by the PSF measurement. As an analytical PSF model is usually not available or not representative of the real optical aberrations, a step of PSF fitting is often necessary. Once the PSF is available, the image restoration is performed so as to get the closest image to the true object [8, 9].

2.2 Quantification of resolution associated to a NIR of excitation in MPM

Experimentally the resolution is a parameter that can be measured thanks to microspheres, standardized object with a known shape. Here, we are illustrating the resolution damages induced by the diffraction limit thanks to two kinds of colored microspheres: green and orange microspheres having a diameter of 0.2 µm. Table 1 presents the measured resolution thanks to a standard commercial solution of PSF estimation and image restoration: the commercial software Huygens, SVI. The two microspheres are imaged with our multiphoton microscope in standard excitation condition. 10 microspheres have been imaged for each measurement.

| Microsphere | Raw PSF $PSF$ (µm) | Raw PSF $PSF_z$ (µm) | PSF after deconvolution $PSF_{xy}$ (µm) | PSF after deconvolution $PSF_z$ (µm) |
|-------------|------------------|------------------|----------------|
| Green       | 0.58             | 1.82             | 0.31           | 0.78             |
| Orange      | 0.61             | 2.13             | 0.34           | 1.35             |

The diffraction limit has a nonhomogeneous impact on the image resolution depending on the considered plan. In the optical plan (xy), the resolution is not significantly damaged by physical phenomena. Nevertheless, the resolution in the...
optical axis (z) is hugely dependent on the diffraction limit: 17% of increase for a raw PSF measurement between a green and an orange emission as shown in Table 1. By consequence, the NIR of excitation is necessarily associated to a damaged resolution compared, for example, to confocal or widefield fluorescence microscopy methods which involve a UV or visible excitation range.

2.3 Quantification of resolution damages associated to deep imaging in MPM

Even though MPM has a degraded resolution in the optical axis compared to standard confocal microscopy methods, the interest of MPM remains in its NIR of excitation. Indeed, biological samples contain constituents which are less absorbing and scattering NIR of excitation than UV or visible excitation. Thus, deeper 3D imaging windows are accessible thanks to MPM. In that situation, we have quantified the resolution deterioration into a sample of 2-mm-depth. We have measured the PSF evolution into a sample of gel containing fluorescent microspheres with a diameter of 0.2 µm. The microspheres are spread homogenously into the 2 mm depth of the sample. Then, we have imaged the samples and cropped each PSF individually. Then, we have measured the PSF at the full width at half maximum (FWHM). Figure 2 illustrates the evolution of the PSF dimensions in the optical plan (xy) and in the optical axis (z) in function with the depth of the PSF into the sample.

Figure 2. Measurement of the PSF FWHM in function with the depth of the PSF.

The evolution of the PSF dimensions in a 2-mm-deep sample is shown in Figure 2. On this graph, the PSF in the optical plan (xy) does not follow any significant evolution depending on their level of depth under the sample surface. However, the evolution of the PSF dimension, measured in the optical axis is hugely increased all along the 2-mm of depth. At the sample surface, the axial PSF is equal to 1.9 µm at the FWHM. This value is within the expected standards. At 0.5 mm under the sample surface, the axial PSF has increased with a factor 2. At 2 mm under the sample surface, the wideness of the axial PSF has increased with a factor 4. This result, combined with the lateral PSF estimation, illustrates the consequent impact of spherical aberrations occurring in MPM. Spherical aberrations represent thus the more harmful physical phenomenon for optical resolution in MPM.
2.4 Illustration of resolution damages associated to deep 3D imaging in MPM

For illustrating the damages of resolution associated to deep 3D imaging of biomedical samples in MPM, the whole muscle is imaged with a standard solution of MPM. 2D images are recorded at many levels of depth with a fixed step of 0.1 µm. The stacks are then superimposed to generate the 3D image. Skeletal muscle is a good candidate for illustrating depth resolution damages, thanks to its striated structure, constituted by bundles of fibers, themselves constituted by a periodical assembly of myosin and actin merged alternatively and periodically. Different levels of periodicity can thus be imaged. Moreover, myosin is a protein organized in a noncentrosymmetric structure which generates a signal of second harmonic. The second harmonic generation (SHG) is emitted label-free by the structure and then collected by the detectors. Figure 3 presents the 3D image of the whole muscle. The image is generated with a field of view (FOV) of 1.6×1.6×0.7 mm³. Figure 3 left shows the 3D image with the standard xy optical plan. Figure 3 right illustrates the 3D image in the xz plan representing the optical axis.

Figure 3. 3D images of a whole muscle. Label-free imaging of the myosin organization. Left image represents the front view of the image of the muscle sample, corresponding to the xy optical plane in the foreground. Right image represents the side view of the image of the muscle sample, corresponding to the xz plane illustrating the resolution deterioration in the optical axis.

Figure 3 right highlights several artefacts on the image. First, the 2D images which have been superimposed are revealed with vertical and bright lines. Moreover, the emitted SHG is less precisely defined on the image in the depth and appears with bright trails.

2.5 Conclusion about the need in computational strategy in 3D MPM

MPM is the exclusive solution offering 3D images of biological samples at millimetric depths and subcellular scale with a sub-micrometer resolution. But this optical method suffers from significant optical artefacts and especially spherical aberrations which damage the axial resolution of the 3D image, quantified by the PSF dimension in the optical axis (Figure 2). A realistic computational strategy for computational restoration of 3D MPM images generated along millimetric depths must consider the axial evolution of PSF. In this context, the quality of PSF estimation becomes the essential parameter for a successful image restoration in MPM. Noise and blur levels increasing with the depth are rendering this topic more complex when the PSF is located deeply under the sample surface. The depth variant property of the PSF imposes to resort to an adapted computational strategies considering this variation in order to remove the PSF contribution to the image, especially in the optical axis. The solution that we have developed is based on a PSF fitting model which considers the spatial variation of the axial PSF in depth.
3. EXPERIMENTAL AND COMPUTATIONAL STRATEGY FOR PSF FITTING IN MPM WITH A 3D MULTI-PARAMETRIC SOLUTION

3.1 Experimental pipeline

Our experimental strategy follows a protocol divided in five steps described thereafter. First, the whole muscle is resected from a wild mouse from tendon to tendon. Then, the muscle is fixed into a gel containing microspheres for PSF estimation. Individual images of microspheres are recorded and cropped. Simultaneously, the SHG signal from myosin of muscle is recorded. Each image of microsphere are then modeled with the FIGARO strategy (FItting GAussians with pRoximal Optimization). Finally, muscle images are deconvolved with the BP3MG solution (Block Parallel Majorize-Minimize Memory Gradient) considering the 3D evolution of the PSF depending on the depth of SHG recording [9].

3.2 Computational strategy

The computational estimation of the PSF Gaussian shape considers that each voxel \( n \in \{1, ..., N\} \) contained into the volume of interest must follow the shape \( y_n \):

\[
y_n = \bar{a} + \bar{b} \bar{p}_n + w_n
\]

\( w \) being the noise. Our strategy relies on the PSF fitting with a Gaussian shape whose expression is given by:

\[
\bar{p}_n \approx \frac{|\bar{C}|^1}{(2\pi)^{\frac{3}{2}}} \exp \left( -\frac{1}{2} (x_n - \bar{\mu})^T \bar{C} (x_n - \bar{\mu}) \right)
\]

Our computational approach proposes a new cost function for estimating \( (\bar{a}, \bar{b}, \bar{p}, \bar{\mu}, \bar{C}) \) from data, based on a new alternating proximal algorithm with convergence guarantees. A penalized least-squares criterion is considered. The minimization procedure follows an alternating minimization scheme with sounded convergence guarantees. Automatic procedure is also included for tuning the regularization parameter, in accordance with the noise statistics.

The mathematical details can be found in our previous publication [10]. Moreover, we have developed a version of this algorithm presented as a free plugin for Image J. The plugin is available by following:

[www.github.com/AlexisLauret/FIGARO](https://www.github.com/AlexisLauret/FIGARO).

3.3 Reconstructed image of a muscle structure considering a space variant PSF

The image size of the whole organ formed with the standard MPM protocol generates about 30 Go of raw data for a single image. No computational system can fairly well process simultaneously the whole image. By consequence, the illustration of our strategy is reduced at a box with dimensions of 22x10x10 \( \mu \text{m}^3 \). Figure 4 represents the resulting image which has been processed by our pipeline considering the axial variations of the PSF.
Figure 4. Muscle image of 22×10×10 µm³ which has been processed by our experimental pipeline considering the axial variations of the PSF. Left image represents the xy optical plan in the foreground (front view). Right image represents the yz plan illustrating the interest of the image restoration in the optical axis (side view).

Figure 4 illustrates the interest of our computational strategy for image restoration especially in the optical axis, by considering the axial space variations of the PSF. Right image represents the side view of the muscle, contained in the yz plan. This viewpoint illustrates the interest of the image restoration in the optical axis. Our computational strategy reveals the myosin organization all along the optical axis. Such a viewpoint for muscle structure has never been shown in any experimental studies of muscles.

4. CONCLUSION

Multiphoton microscopy is an optical imaging method especially adapted to biomedical imaging. Its ability to produce 3D images is unequaled by the other imaging methods. Today, the main strategy chosen for generating 3D images of biomedical samples consists in resecting the sample of interest and associated this step to a labelling process with labeled antibodies. Such a time consuming, and costly solution, associated to image slicing and confocal image recording, generates only a partial description of the 3D structure of the muscle, missing out on details concerning the 3 dimensional organization. This property is especially true in muscle imaging where the 3D image reconstruction is often associated to a physical sample slicing. Here, we present our solution combining two technological advances: an instrumental one with a computational one. Instrumentally, we benefit from the non-centrosymmetric organization of myosin, a structural protein of muscle. This organization generates a second harmonic signal; myosin can thus be image label free in the whole depth of the muscle. Our computational strategy proposes a robust solution for point spread function fitting, which is usually deteriorated by noise, especially in the case of multiphoton excitation. Associated to an adapted deconvolution strategy, this robust and reliable computational strategy shows a 3D organization of muscle which has never been demonstrated until now.
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