Challenges in 3D Live Cell Imaging

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Abstract: A short overview on 3D live cell imaging is given. Relevant samples are described and various problems and challenges—including 3D imaging by optical sectioning, light scattering and phototoxicity—are addressed. Furthermore, enhanced methods of wide-field or laser scanning microscopy together with some relevant examples and applications are summarized. In the future one may profit from a continuous increase in microscopic resolution, but also from molecular sensing techniques in the nanometer range using e.g., non-radiative energy transfer (FRET).

Keywords: 3D cell cultures; light scattering; phototoxicity; wide-field microscopy; laser scanning microscopy; structured illumination; light sheet microscopy; FRET

1. Introduction

Experimental and pre-clinical life cell approaches traditionally use two-dimensional (2D) cell cultures, which are easy to establish, but frequently provide results of limited significance, since cells are lacking a physiological microenvironment. In contrast, three-dimensional (3D) cell cultures, e.g., multicellular tumor spheroids (MCTS), maintain tissue-like properties and therefore provide a more realistic background for experimental studies, e.g., screening of pharmaceutical agents [1,2]. However, imaging of 3-dimensional specimens is challenging, since the sample thickness commonly exceeds the depth of focus of a conventional detection system, and light scattering considerably impairs the image quality. Therefore, methods based on optical sectioning, e.g., confocal laser scanning microscopy (CLSM) [3,4], Optical Sectioning Structured Illumination Microscopy (OS-SIM) [5], or light sheet fluorescence microscopy (LSFM) [6,7] are applied preferentially. Here, images are recorded plane by plane, and resulting 3D plots are calculated offline. A problem for CLSM related methods as well as for OS-SIM is that for imaging each plane the whole sample has to be illuminated, so that upon recording of the whole specimen phototoxic damages are likely to occur [8]. Furthermore, photobleaching may increase in the course of an experiment and falsify the experimental results. Altogether, 3D imaging creates a large number of data (“big data”), which have to be handled appropriately.

2. 3D Samples

Two-dimensional cell cultures have a well-established protocol in biomedical research and provide a simple, fast, and cost-effective tool for e.g., drug discovery assays. However, mammalian cells commonly grow within a complex three-dimensional microenvironment with a different gene expression and protein synthesis pattern [9]. Therefore, various 3D models have been established to better mimic the natural cell environment.

Cells embedded in hydrogels, e.g., agarose, which are readily accessed by optical sectioning methods like CLSM or OS-SIM may provide more realistic studies of cell morphology, e.g., to examine the influence of mechanical signals on cell behavior [10]. Multicellular (tumor) spheroids have gained significance in preclinical studies, as they appear to be more appropriate for studies of cell physiology, cell metabolism, or tissue diagnostics. 3D cultivation techniques commonly prevent cell attachment to surfaces, using hanging drop
methods, liquid overlay methods or agitation-based approaches [11]. Cell cultivation in a solid matrix, e.g., agarose gel, may permit 3D cell growth under realistic conditions [12,13]. Cell spheroids are generally characterized by an external proliferating region and an internal quiescent zone (caused by the gradient of nutrient and oxygen diffusion), which may surround a necrotic core in larger spheroids [14].

Organoids are a type of 3D cell culture containing organ-specific cells that have been grown from a range of organs, including kidney, breast and liver (for a review see e.g., [15,16]). However, current 3D systems often lack a vasculature, which might support tissues with oxygen and nutrients, remove waste and build up an immune system. Nevertheless, organoid development is a rapidly growing field, and complex 3D systems including fully vascularized brain organoids [17] or organs-on-a-chip [18] have been reported in recent studies.

Traditionally, biopsies from a histological laboratory are routinely fixed and either embedded in paraffin, or frozen as thin sections, stained and mounted on glass slides. These procedures, however, introduce artifacts and severely limit the information, as only a small fraction of a specimen is used for microscopy. Novel approaches of nondestructive slide-free pathology are investigated, which allow deep volumetric microscopy of whole biopsy specimens [19]. Imaging of whole organisms is an important tool in developmental biology as well as drug screening [20]. Microscopy of small organisms requires either high transparency or the application of optical clearing techniques (see below), if viability is not a main criterion.

Measurements of 3D (cell) cultures often need specific sample holders, e.g., glass or plastic tubes or even micro-capillaries, which may be rotated for multi-view applications (see e.g., [21]).

3. Phenomena and Challenges

3.1. Light Scattering

Interaction of light with any kind of samples is described in terms of absorption and scattering. In particular, light scattering experiments with angular or spectral resolution have been used for more than 30 years for characterization of various types of cells [22,23] or for measurement of morphological changes in cells undergoing necrosis or apoptosis [24,25]. However, scattering reduces the quality of images due to light attenuation, blurring and a loss of contrast. Obviously, these problems are more severe for three-dimensional than for two-dimensional samples, since scattering does not only occur in a certain plane of detection, but also creates background signals from the whole illuminated volume. This is well documented by Figure 1 showing 3D spheroids of Chinese hamster ovary (CHO) cells of about 250 µm diameter expressing a membrane-associated Green Fluorescent Protein (GFP). Conventional fluorescence microscopy (Figure 1a) shows a completely blurred image, since information from the focal plane is superposed by out-of-focus images, and since pronounced scattering further reduces the image quality. The impact of scattering appears lower, if individual planes of the sample are selected either by confocal laser scanning microscopy (CLSM) or by light sheet fluorescence microscopy (LSFM). Nevertheless, a loss of fluorescence intensity occurs in the central parts of the CLSM image (Figure 1b) and along the direction of light propagation in the LSFM image (Figure 1c). Obviously, light attenuation is less pronounced in the LSFM image, where due to the anisotropy of Mie scattering [26] light is scattered preferentially into forward direction. However, some stripes in the direction of light incidence are often unavoidable. Scattering is becoming lower at higher wavelengths, which are used preferentially in multiphoton microscopy [27,28].
For reduction of light scattering optical clearing techniques matching the refractive indices of sample and surrounding medium have gained considerable importance. Therefore, these techniques are used increasingly for deep view imaging of skin, brain and other organs [29–31]. Currently available optical clearing techniques are not compatible with live cell imaging. However, efforts are being made to find biocompatible solutions, especially for ex-vivo applications [32]. While penetration depths are limited to 100–200 µm in non-cleared samples, they can be even larger than 0.5 mm in cleared samples, permitting e.g., to image entire neuronal networks in mouse brains.

3.2. Phototoxicity, Photobleaching

As reported above, 3D images are often based on optical sectioning, and information is summed up from z-stacks of individual exposures. Only for LSFM, each sectional image results from one illuminated plane, whereas for other wide-field and laser scanning techniques, the whole specimen has to be illuminated for each image section. This implies that light exposure for obtaining a 3D image is summing up and often exceeds the limit of non-phototoxic light doses. Tolerable light doses were determined in a previous manuscript [8] and ranged between 25 J/cm² (375 nm) and 200 J/cm² (633 nm) for cultures of native cells, thus increasing with illumination wavelength and corresponding to 4 min. up to about 30 min. of solar irradiance (around 100 mW/cm²). If cells were stained with a fluorescent dye or transfected with a fluorescent protein, typical non-phototoxic light doses were only around 10 J/cm², corresponding to 100 s of solar irradiance. In Figure 2, a maximum number of images is indicated for the case that cells are illuminated with 100 mW/cm² (corresponding to 1 nW/µm²) for 1 s (wide-field images) or 5 s (laser scanning image). While only about 20 layers of a 3D cell spheroid can thus be irradiated once by CLSM, each layer can be illuminated about 100 times by LSFM. This favors light sheet microscopy for long-term experiments in cell or developmental biology.

In previous studies [33] we found that non-phototoxic light doses did not depend on whether the light was applied continuously or in short pulses. This implies that for multiphoton imaging (see Section 4.1) the integral light dose and the wavelength of illumination appear to be the main limiting factors. Therefore, due to the longer wavelengths phototoxicity is generally lower for multiphoton than for single photon imaging.

Increasing sensitivities of novel detection systems (e.g., ultra-sensitive cameras) will increase the number of images measured at non-phototoxic light doses and will permit recording of fast dynamic processes, e.g., rapid cell migration, membrane or microtubule dynamics, mitochondrial motion as well as endo- or exocytosis.
4.1. Confocal Laser Scanning Microscopy

In Confocal Laser Scanning Microscopy (CLSM) samples are scanned point by point, resulting in a two-dimensional image within typically a few seconds. A spatial pinhole in a conjugate image plane is used to block out-of-focus light thus acquiring optical sections with high contrast, as depicted in Figure 3. Information (e.g., fluorescence) from individual

<table>
<thead>
<tr>
<th>Method</th>
<th>Experiment</th>
<th>Max. light dose [J/cm²]</th>
<th>Irradiance [mW/cm²]</th>
<th>Max. no. of images</th>
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<tbody>
<tr>
<td>Wide-field microscopy</td>
<td>Autofluorescence</td>
<td>25</td>
<td>100</td>
<td>250</td>
</tr>
<tr>
<td>Wide-field microscopy</td>
<td>Fluorescence marker</td>
<td>10</td>
<td>100</td>
<td>100</td>
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<tr>
<td>CLSM</td>
<td>Fluorescence marker</td>
<td>10</td>
<td>100</td>
<td>20</td>
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<tr>
<td>LSFM (N layers)</td>
<td>Fluorescence marker</td>
<td>10</td>
<td>100</td>
<td>N × 100</td>
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Figure 2. Maximum non-phototoxic light doses and maximum number of images for various methods of 3D live cell imaging. For autofluorescence experiments an excitation wavelength of 375 nm is assumed. An exposure time of 5 s is assumed for LSFM, and a time of 1 s for all other (wide-field) techniques (data partly reproduced from [8]).

A further phenomenon upon pronounced light exposure is photobleaching or fluorescence bleaching. This effect may be concomitant with modification or destruction of a specific fluorophore [34] and makes quantitative evaluation of fluorescence signals difficult. In some cases, intersystem crossing to a (non-fluorescent) excited triplet state occurs, and after deactivation of this state the corresponding molecules may fluoresce again ("fluorescence recovery"). This effect often causes characteristic "blinking" and is used in single molecule spectroscopy (for a review see [33]). For more than 40 years "fluorescence recovery after photobleaching" (FRAP) has been applied to measure cell, membrane and, in particular, protein dynamics (for reviews see [36,37]). In this case, part of a fluorescent specimen is photobleached, and re-diffusion of molecules from outside this part is measured. However, this method should be applied with care since high light exposure may damage living specimens.

3.3. “Big Data”

3D live cell imaging, especially light sheet microscopy of larger specimens, generates large datasets that need to be stored and processed. Multimodal configurations, e.g., time-lapse or multispectral devices, or high-throughput/high-content setups add even more data leading to multidimensional datasets in the gigabyte or even terabyte range [38,39]. High-end computer hardware and central networks for efficient storage and retrieval of data as well as for fast processing of huge datasets and appropriate data management are needed. Open source software applications, e.g., OME Remote Objects (OMERO), enable access to and use of a wide range of biological data and provide open, flexible solutions for data management [40]. Recently, automated image processing and machine learning have become valuable tools to extract meaningful information from large datasets. As cells can be regarded as highly controlled objects, microscopy is well suited to pattern recognition tools based on neural networks and deep learning [41]. Several commercial (Imaris, Amira, Arivis) as well as non-commercial (BigDataViewer plugin for Fiji/ImageJ [42], ilastik [43]) software applications for high-performance 3D visualization and analysis are available.

4. 3D Microscopy

For many imaging applications optical microscopy and, in particular, fluorescence microscopy plays a predominant role, and, therefore, relevant techniques are summarized in this section.

4.1. Confocal Laser Scanning Microscopy

In Confocal Laser Scanning Microscopy (CLSM) samples are scanned point by point, resulting in a two-dimensional image within typically a few seconds. A spatial pinhole in a conjugate image plane is used to block out-of-focus light thus acquiring optical sections with high contrast, as depicted in Figure 3. Information (e.g., fluorescence) from individual
planes is then used for 3D image reconstruction by appropriate software. However, the whole procedure needs comparatively long scanning times and high light exposure, which may damage living specimens. One possibility to reduce light exposure is simultaneous illumination of several parts of a sample by Spinning Disk Confocal Microscopy [44].

In a classical laser scanning setup resolution is only slightly improved compared to wide-field microscopes (see below). If, however, the pinhole selects only a small part of the diffraction pattern (“Airy disk”) of individual points, the resolution may be improved up to almost a factor 2, while the detection sensitivity is reduced considerably. To circumvent this problem, the single pinhole has recently been replaced by an array of several individual detectors (Airy Scan Microscopy [45]), or by an ultrasensitive camera chip, where each pixel behaves as an “infinitely small pinhole” (Image Scan Microscopy [46]). Alternatively, resolution as well as sensitivity can be improved significantly by scanning the pinhole with another laser (Re-Scan Confocal Microscopy [47]).

Multiphoton microscopy (MPM) achieves optical sectioning by focusing picosecond or femtosecond laser pulses onto a small spot in the sample, thus exciting fluorescence by two or more photons in the small focal volume during an extremely short time without the need of any pinhole [27,48]. The scanning process and vertical displacement of the specimen is the same as for CLSM. Although multiphoton microscopes have vastly improved over the last 10 years, MPM still has some limitations in terms of the temporal resolution and long-term applicability as well as instrumentational challenges, e.g., suitable objective lenses for NIR transmission [49,50].

4.2. Widefield Microscopy

In conventional wide-field microscopy, an image is generated from the focal plane of a sample with a lateral resolution $\Delta x \geq \lambda / 2A_N$, as defined by the Abbe criterion for the case of coherent illumination. Alternatively, the Rayleigh criterion $\Delta x = 0.61 \lambda / A_N$ defines the
resolution of a luminescent spot or an object illuminated incoherently with the wavelength $\lambda$ and the numerical aperture $A_N$ of the microscope objective lens. In both cases values around 200 nm are achieved for numerical apertures $A_N \geq 1.30$ at $\lambda = 500$ nm. Often the axial resolution is related to the depth of focus $\Delta z = n \lambda / A_N^2$, resulting in $\Delta z \geq 400$ nm for the refractive index $n = 1.50$. This implies that the focal plane is very thin, and a lot of out-of-focus information blurs the image considerably. Therefore, similar to CLSM, optical sectioning is often necessary.

Optical Sectioning Structured Illumination Microscopy (OS-SIM) is a wide-field technique to separate in-focus information from signals generated in out-of-focus planes. To achieve this, an optical grid is imaged in the plane of the sample in 3 phase positions $\Phi_1 = 0$, $\Phi_2 = 2\pi/3$ and $\Phi_3 = 4\pi/3$ resulting in the intensities $I_1$, $I_2$ and $I_3$. An algorithm $I = [(I_1 - I_2)^2 + (I_1 - I_3)^2 + (I_2 - I_3)^2]^{1/2}$ permits to calculate an image from the focal plane, while out-of-focus images add up to zero, as depicted in Figure 4. Upon variation of the sample’s vertical position, several focal planes can thus be detected, and a 3D image may be calculated [5]. While Figure 4 shows the principle of this method, Figure 5 shows an application to CHO cells transfected with a membrane associated Green Fluorescence Protein (GFP).

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Principle of Optical-Sectioning Structured Illumination Microscopy (OS-SIM). Reproduced from [51] with modifications.

![Figure 5](https://example.com/figure5.png)

**Figure 5.** Spheroid of CHO-pAcGFP1-Mem cells. Conventional wide-field (left) and OS-SIM images (right) at different depths within the spheroid. While conventional wide-field microscopy includes a large amount of out-of-focus information, OS-SIM selects focal planes and resolves single cells.
It should be added that Structured Illumination Microscopy can also be used for super-resolution imaging with a 2-times enhancement of resolution in comparison with the Abbe criterion. In contrast to OS-SIM, “SR-SIM” is based on a coherent superposition of two or three laser beams modulated by an optical grid at different angles and phase positions. However, this technique is commonly limited to thin samples, as out-of-focus light degrades the quality of the grid pattern in the sample plane. In addition, acquisition of multiple (typically 9–15) raw images needed to reconstruct a single super-resolution frame takes relatively long time and thus reduces acquisition speed. These limitations can be overcome by illuminating the sample with a lattice pattern rather than grid lines, which gives higher contrast and allows a more robust image reconstruction.

Light Sheet Fluorescence Microscopy (LSFM) is another wide-field method that uses optical sectioning with the sample being illuminated from the side (at 90° with respect to the microscope axis) by either a cylindrical lens or scanning of a laser beam. For 3D imaging, the light sheet and the microscope objective lens used for detection can be shifted simultaneously in axial direction, so that the illuminated part of the sample is always in the focus of the objective lens. Both shifts may be different due to the refractive index of the immersion fluid, but this can be corrected either mechanically or by software. Alternatively, the sample can be moved in axial direction through a static light sheet. Thus, z-stacks can be recorded with low fluorescence background and high contrast, as depicted in Figure 6 for spheroids of MCF-7 breast cancer cells incubated with the cytotoxic drug doxorubicin. A 3D image can be calculated offline from these images, as described elsewhere. The main advantage of Light Sheet over OS-SIM is that only those planes are illuminated, which are recorded simultaneously, so that light exposure is considerably lower than for those methods, where recording of each image requires illumination of the whole specimen. Commercial light-sheet microscopes (e.g., Carl Zeiss, Olympus, Nikon) as well as open-source solutions or add-ons for existing microscopes are presently available.

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With Axial Tomography specimens can be observed from various sides, and since the lateral resolution in microscopy is always better than the axial resolution, the vertical axis can be rotated into the horizontal position to profit from an enhanced isotropic resolution of about 200 nm for conventional wide-field microscopy or CLSM and about 100 nm for SR-SIM. However, rotation of samples needs specific sample holders and preparation techniques, e.g., cells embedded in a gel within a rotatable capillary, whose refractive index can be adapted to that of water or oil when using an immersion lens of high magnification. A particular problem for 3D specimens is precise positioning of the samples upon rotation as well as appropriate reconstruction of 3D images.

Figure 6. Principle of light sheet microscopy (a); image sections of MCF-7 spheroids incubated for 6 h with doxorubicin (8 µM) at a depth of 20 µm (b) or 60 µm (c) within the spheroid. Red and green-yellow fluorescence permit distinction of doxorubicin and its degradation product. Reproduced from [57] with modifications.
Recently, Light Field Microscopy became an interesting candidate for 3D imaging of fast biological processes, e.g., neural activity in whole animals [60,61]. It provides instant 3D excitation and single-shot volumetric recording that creates up to 200 3D images per second, but requires a computationally demanding image reconstruction process and offers comparably low spatial resolution. Novel approaches to image reconstruction with deep-learning algorithms aim to overcome these limitations [62].

4.3. Deconvolution

In a diffraction-limited system, image formation can be described as a convolution of light emitted from each point in the object with the point spread function (PSF) of the microscope. Deconvolution, an image processing technique used to reverse this blurring process has been introduced almost 40 years ago, and with increasing memory capacity and calculation speed of personal computers 3D-deconvolution has become a well-established lab tool for deblurring of microscopy data from various 3D techniques [63–66]. Commercial deconvolution software packages, e.g., Huygens (Scientific Volume Imaging, Hilversum, The Netherlands) or Auto Quant (Media Cybernetics, Rockville, MD, USA) and public domain tools, e.g., Deconvolution LAB2 for Fiji [67], provide tools for 3D deconvolution, which are comparably easy to use even for non-specialists in computational image processing. Deconvolution algorithms are also used during image reconstruction for super resolution methods, e.g., SR-SIM [68,69] and Image Scanning Microscopy [46].

5. Discussion

3D imaging of living specimens is a very challenging topic requiring: (1) experience in the preparation and handling of 3D samples, (2) techniques of image sectioning and reconstruction, (3) minimization of light scattering, (4) light dosimetry, as well as (5) processing strategies for large datasets. Many experimental methods are based on light microscopy, but also macroscopic or endoscopic systems as well as multi-well screening techniques may be appropriate for certain specimens and applications. Since electron microscopy is difficult to realize for living cells due to fixation of samples and high irradiation doses [70], it is not further considered in this manuscript. However, “correlative microscopy”, i.e., a combination of optical and electron microscopy presently gains considerable importance in optical nanoscopy. Often nanometer distances cannot be resolved within a microscope, but it is well known that fluorescence spectra as well as fluorescence lifetimes (for an overview see [71]) depend on the microenvironment of a relevant fluorophore, including pH, viscosity, and polarity. For example, shortening of the fluorescence lifetime of GFP in close vicinity to cell membranes or to a cell-glass interface has been reported in the literature [72,73]. Molecular sensing techniques based on fluorescence spectra or lifetimes may be applicable also to 3D cell systems. This holds in particular for studies on Förster Resonance Energy Transfer (FRET), where after excitation of a so-called donor molecule excitation energy is transferred to an acceptor molecule by interaction of optical transition dipoles, if their distance is less than about 10 nm [74]. Applications to nanometer sensing in 3D systems include studies of subdomains of proteins [75], probing of molecular cell-biomaterial interactions [76] or monitoring of apoptosis by cleaving of a specific peptide linker [77]. In case of larger molecules adjacent in cellular micro-domains (e.g., membrane domains or intracellular organelles), or for studies of more than two interacting molecular populations, their co-localization can be quantified to analyze this interaction. Co-localization methods are based on two different, complementary methods: a pixel-based approach [78,79] or an object-based approach [80,81], as well as a combination of the two approaches [82]. In the future one may expect further increase of resolution in microscopy at non-phototoxic light doses as well as novel highly specific applications of nanometer sensing in 3D cell systems.

6. Conclusions

We conclude that 3D live cell imaging is still a developing field for studying cell models with tissue-like properties, whole organs or intact organisms. Appropriate methods have
been developed, which remain to be further elaborated. Novel preparation techniques, reduction or mathematical correction of light scattering, minimization of light exposure and phototoxicity, as well as processing of large data sets remain great challenges for the future.

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