Manual

An ImageJ Plugin to Measure 3D Point Spread Functions (alpha)

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1 Introduction

Subpixel-resolution image analysis often exploit knowledge about the imaging system to reconstruct the hidden state given the observations (recorded images). The mapping from *object space* to *image space* is often modeled using a convolution of the true state with the point spread function (PSF) of the imaging system:

$$I = O * PSF + \eta$$

where η is a random variable accouning for noise. Usually, the PSF is of Gaussian-blob like shape. The task of reconstructing the objects from the images using deconvolution techniques is much easier when the PSF is known.

Note that, when convolving a dirac delta function δ with a kernel K, the result is K itself since δ is the neutral element of the convolution operation. In fluorescence microscopy, imaging small fluorescent microspheres, also called *beads*, as an approximation to a dirac impulse, enables to observe the PSF of the system in the resulting image.

This ImageJ plugin creates high resolution PSF images by averaging many bead images as well as exploiting the assumption that the point spread function is rotationally symmetric with respect to the axial axis (z-direction).

Note that in 3D microscopy the PSF may differ depending on the axial position of the emmitting light source. This effect is neglected by the current implementation.

2 Installation

Copy the mosaic_plugins.jar file to the ImageJs plugins directory. Restart ImageJ. The plugin can be launched from the Mosaic submenu in ImageJs plugin menu.

The plugin handles 8-bit, 16-bit and 32-bit grayscale images or stacks.

Prerequisits: At least Java 5.0 and ImageJ 1.36.

3 How to use the plugin

Open a 3D or 4D image stack in ImageJ. Please make sure to correctly set the number of slices per frame and the number of frames correctly in the image properties dialog (Ctrl-P). Also put the correct value for the pixel size and focal plane distance (preferably in nm). In the Plugin—Mosaic menu, selecting the PSF measure 3D entry in the Mosaic submenue starts the plugin.

A maximum intensity projection of your stack pops up. Then, select ImageJs point selection tool. Clicking on a bead while holding the SHIFT button creates a PSF map with the first measurement (the first measurement might take some time). Repeat this for all the good quality beads in your image. The PSF-map (see figure 1) created after each new measurement is averaging all the measurements taken so far. To remove a measurement, shift-click on the bead again.

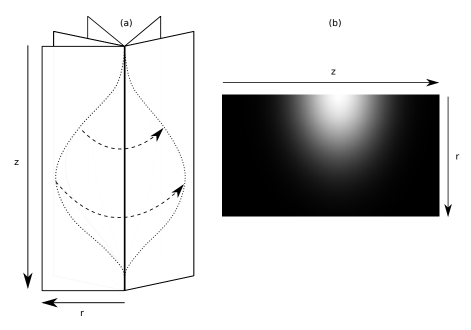


Figure 1: (a) Intensity isoline of the PSF: rotating the PSF map around the axial axis yields a 3D PSF. (b) A PSF map generated by measuring 20 beads. The beads were recorded by a spinning disk confocal microsocpe.

3.1 Parameters

- Maximum lateral sampling radius: maximal distance r_{max} (in nm) from the centroid for sampling the image.
- Maximum axial sampling radius: maximal distance r_{max} (in nm) from the centroid for sampling the image.
- Final lateral PSF map size: The final PSF map size (in pixel) in r-direction.
- Final axial PSF map size: The final PSF map size (in pixel) in z-direction.

4 Description of the measurement

The first step is a centroid detection [1] of the bead close to the position where the user clicked on the image. The image is then sampled around the centroid, the corresponding value is then put into the psf map. After having sampled all the pixel around the centroid, the PSF-map (r, z)-entries that are not yet filled with any data are filled using bi-linear interpolation. If there are multiple entries for one (r, z)-entry, the values are averaged.

5 Disclaimer

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References

[1] I. F. Sbalzarini and P. Koumoutsakos. Feature point tracking and trajectory analysis for video imaging in cell biology. *Journal of Structural Biology*, 151(2):182–195, August 2005.