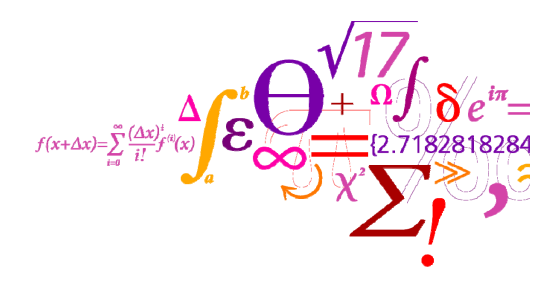


Confocal Microscopy and Atomic Force Microscopy (AFM) of biofilms

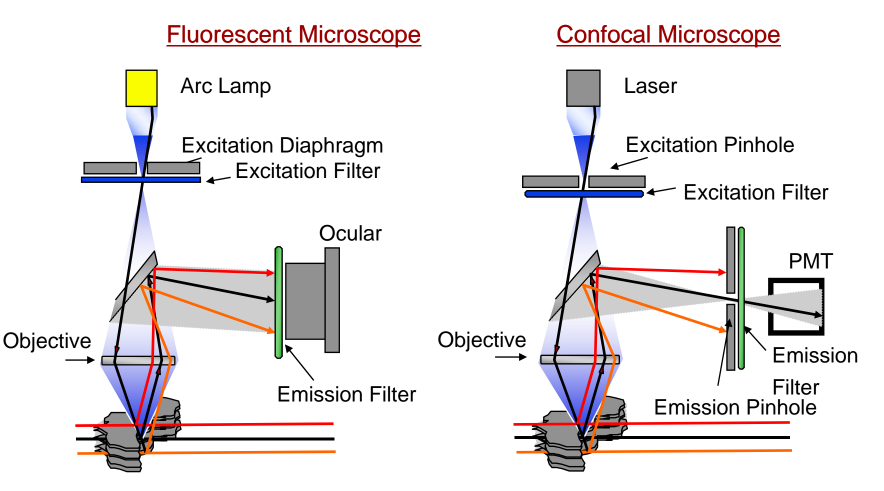
A very brief primer...



DTU Systems Biology Department of Systems Biology

Fundamentals of Confocal Microscopy

Based on a conventional fluorescence microscope

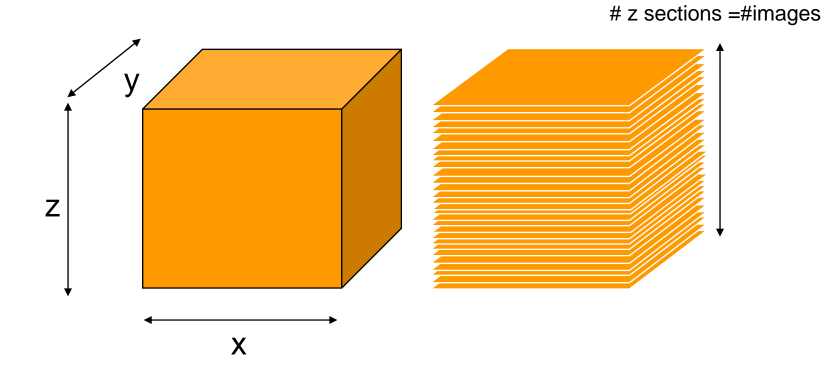


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Confocal and Atomic Force Microscopy 15 March 2010



3D reconstruction



Pseudomonas putida cells mixed with Acinetobacter cells in a microbial biofilm 20 µm

Christensen et al. 1998. Appl. Environ. Microbiol. 64: 2247-55

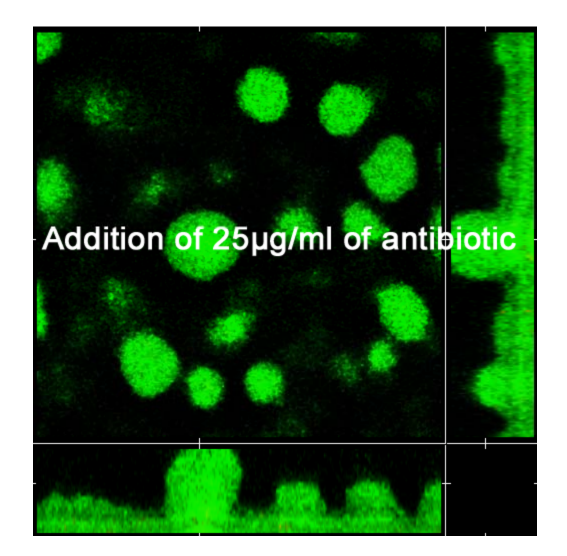
Benefits of Confocal Microscopy

- Reduced blurring of the image from light scattering
- Increased effective resolution
- Improved signal to noise ratio
- Clear examination of thick specimens
- Z-axis scanning (3D-reconstruction possible)
- Magnification can be adjusted electronically
- X-Y resolution: ~200-250 nm (Ernst Abbe)

Disadvantages of Confocal Microscopy

- Requires fluorescent samples
- Uses laser illumination (expensive, few wavelengths)
- Instrument expensive to acquire and run
- Z-resolution typically >500 nm

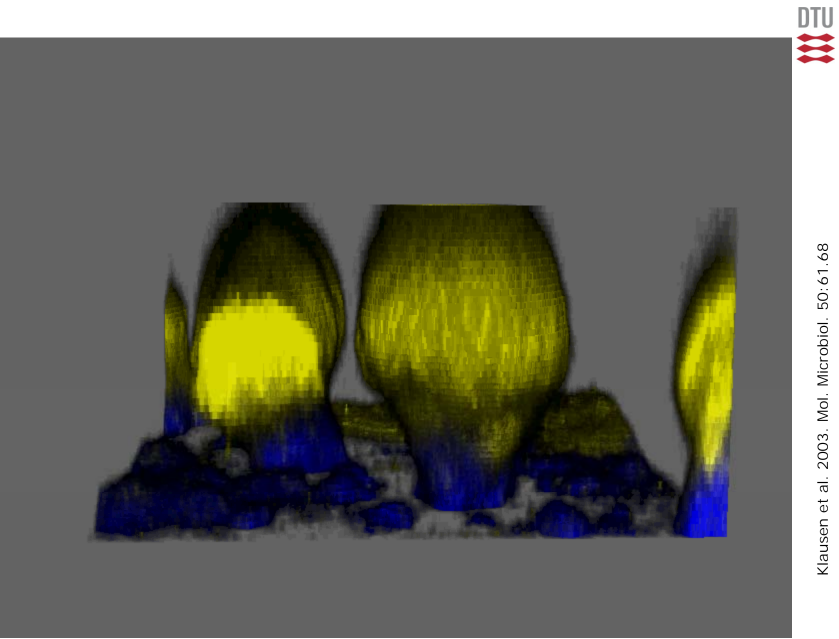


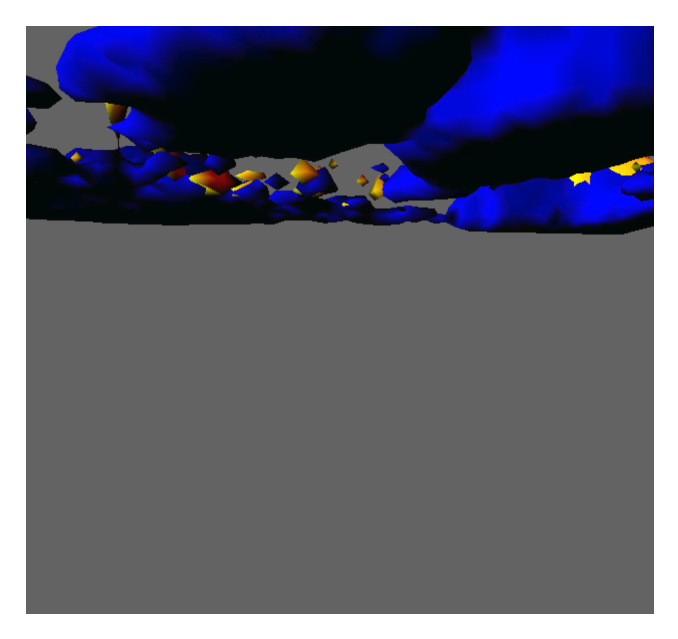


Haagensen et al. 2007. J. Bacteriol. 189:28-37

6 DTU Sytems Biology, Technical University of Denmark

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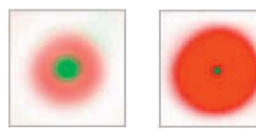


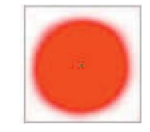
New developments in confocal microscopy

- MP (multi-photon) or two-photon confocal microscopy: Two or more photons from a long wavelength illumination at a time excites a fluorophore. High resolution in Z-axis (practally down to ~200 nm)
- White Laser (Leica): A continous wave "white" laser (tunable from 470-670 nm, with up to 8 lines simultaneously)
- **STED (Stimulated Emission Depletion) (Leica):** A new method where fluourescence is depleted around the area of interest (see next slide)
- Superresolution structured microscopy (Zeiss): A methods where images are rotated and combined to create a moire pattern which is deconvolved to create a high res image.
- **Photo-Activated localization microscopy (Zeiss):** Sequential illumination and localization of fluorophores combined with computational reconstruction of high res images.
- Raman-confocal microscopy (Leica, under development): Confocal microscope combined with raman spectroscope. Enables localized determination of [changes in] concentrations of metabolites etc.

STED (Stimulated Emission Depletion)

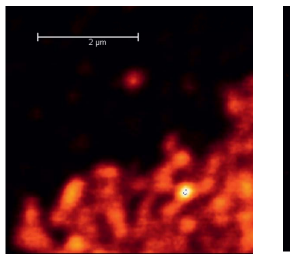
- In a Leica TCS STED microscope the sample is illuminated by two pulsed laser beams, tightly synchronized.
- The 635 nm wavelength excites the fluorophores of the sample the same way a conventional confocal system does. The excitation laser pulses are directly followed by a ring shaped illumination of a Ti: Sapphire Infrared laser 730-780 nm).
- This pulse inhibits/depletes the fluorescence in the outer regions of the illuminated spot.
- The result: A smaller fluorescence spot that allows much more accurate scanning than with other methods using focused light. X-Y res: <90 nm

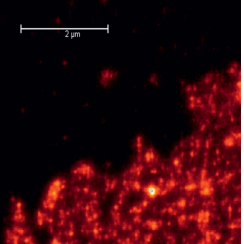




10 DTU Sytems Biology, Technical University of Denmark

Sample STED image





Confocal image

STED image

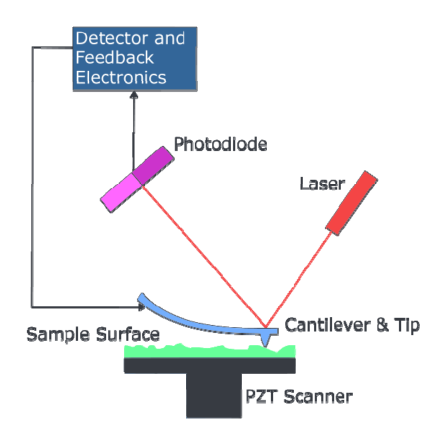
Images: Prof. Dr. T. Lang, Univ. Of Bonn, germany and Leica Microsystems

11 DTU Sytems Biology, Technical University of Denmark

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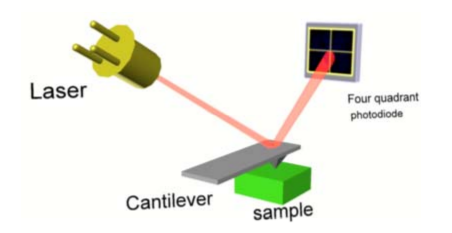
Atomic Force Microscopy (AFM)

- A method to record the topographic property of a surface
- Physical interaction with the surface is necessary

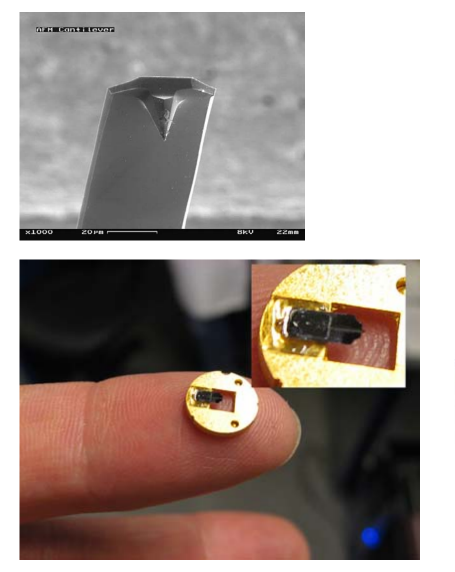




The principle



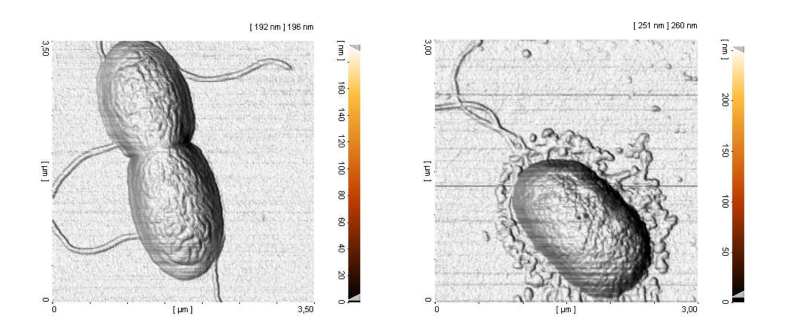






14 DTU Sytems Biology, Technical University of Denmark





Anne Louise Frost m.fl. 2007 (unpublished)

Benefits of AFM

- Extemely high resolution (down to atomic level few Å), but typically 10-20 nm)
- No staining required
- Possible to measure attractive/replusive forces

Disadvantages of AFM

- Extremely high resolution (very low "focal depth")
- Small image area
- Sample must be "flat" and tightly fixed
- Very difficult to work in humid or wet environments
- Need direct contact to sample
- Imaging depends on tip shape
- Slow

