

Optical methods in chemistry
or
Photon tools for chemical sciences

Session 7:

Course layout – contents overview and general structure

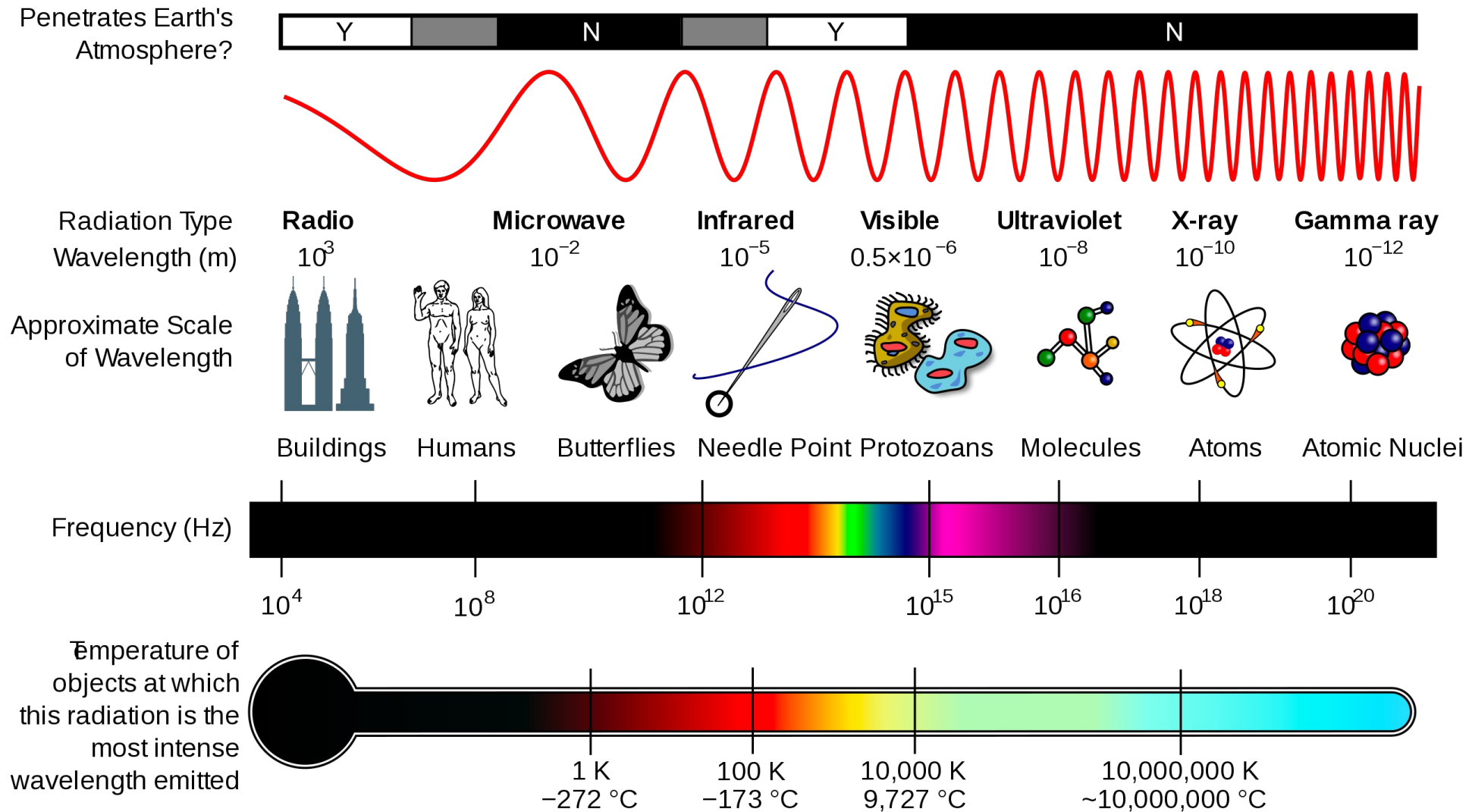
- Introduction and ray optics
- Wave optics
- Beams
- From cavities to lasers
- More lasers and optical tweezers
- From diffraction and Fourier optics
- **Microscopy**
- Spectroscopy
- Electromagnetic optics
- Absorption, dispersion, and non-linear optics
- Ultrafast lasers
- Introduction to x-rays
- X-ray diffraction and spectroscopy
- Summary

Going towards applications:

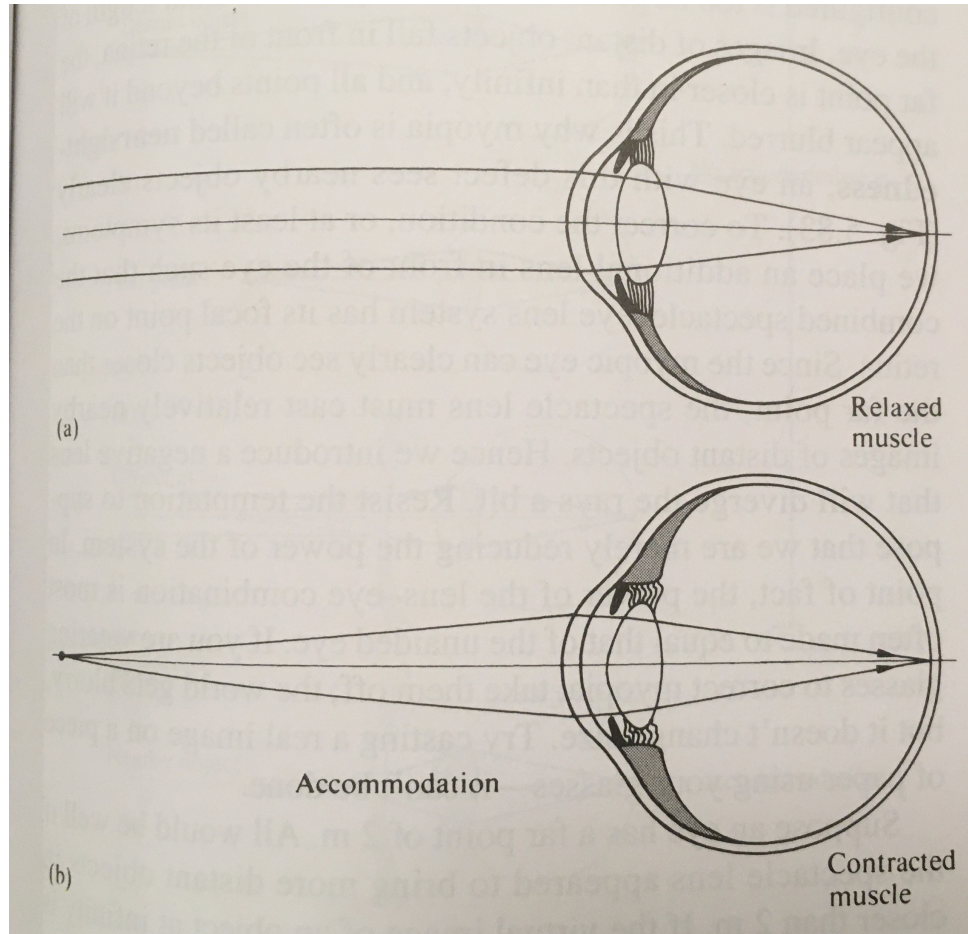
However, still focus on some concepts

Microscopy – Manipulation - Spectroscopy

Remember: Energy – wavelength – size relationship



Reminder: Our eye as an optical system

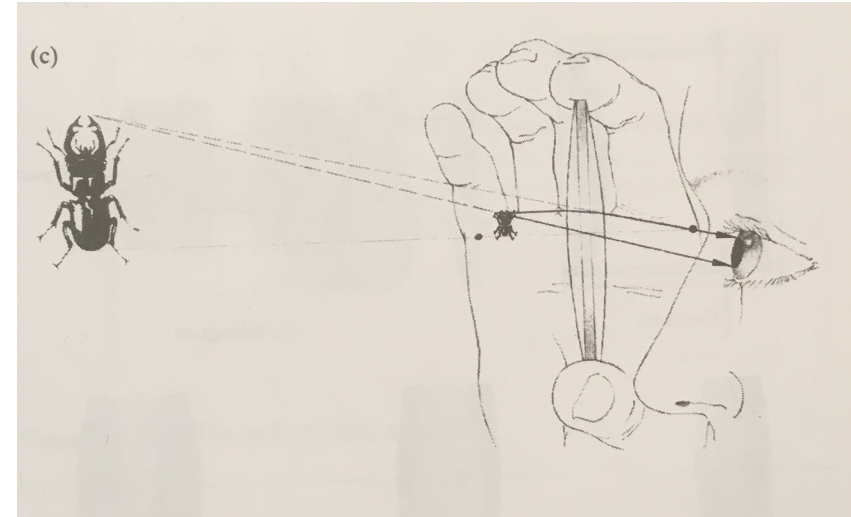
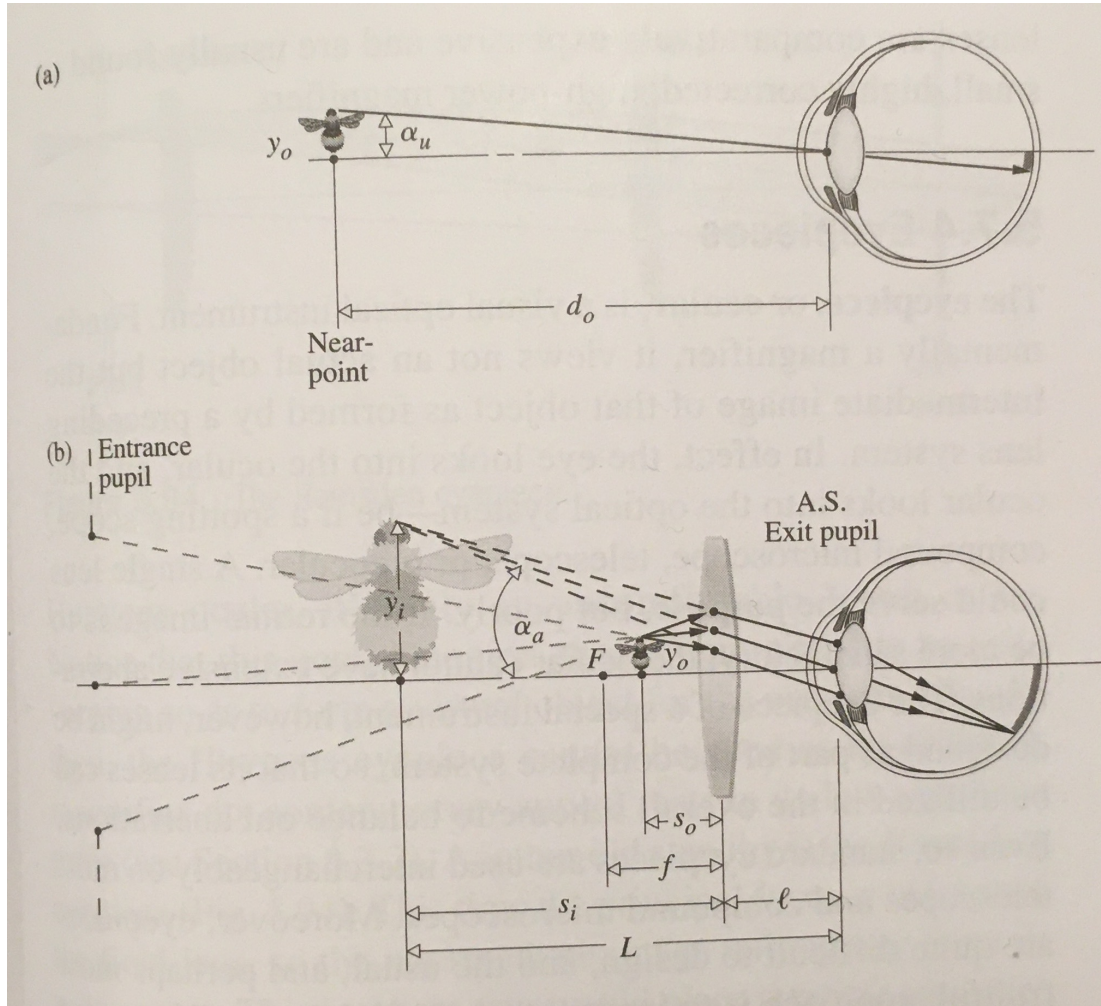


This makes laser radiation so dangerous to eye: parallel beams give perfect focus

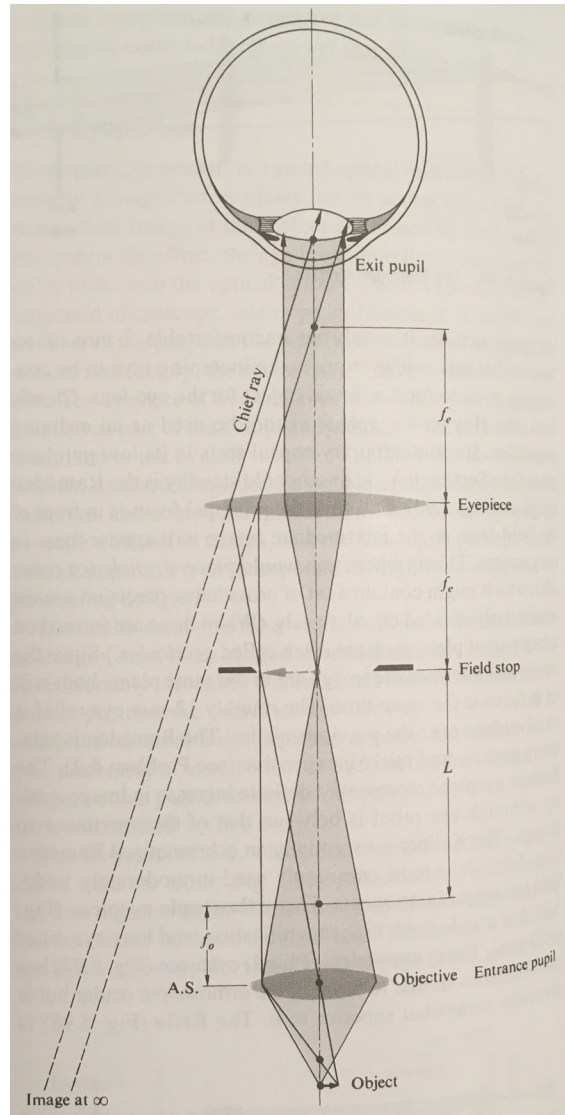
When designing optical systems (microscopes) need to consider eye as optical element.

Nowadays eye is typically replaced by camera which is also an imaging system.

Magnifying glass: a simple optical element



The microscope – a classical view



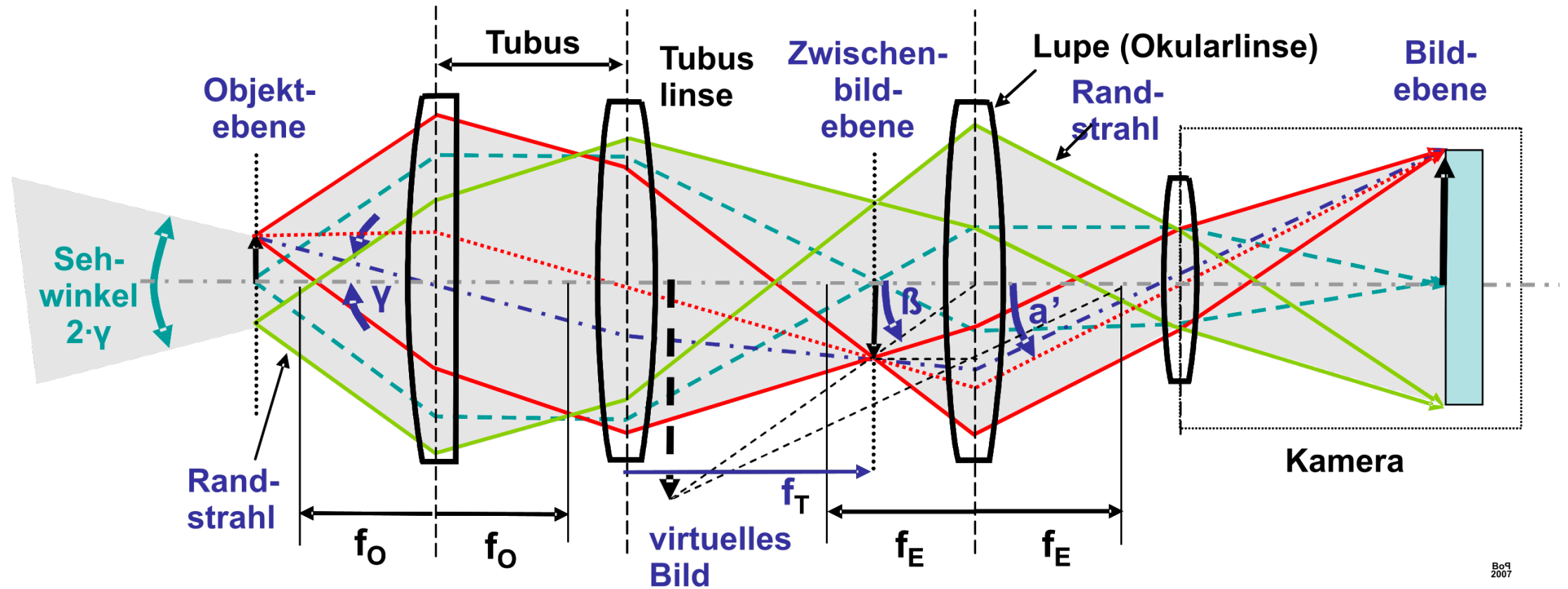
Principle:

Objective forms intermediate image

Eyepiece looks at and magnifies intermediate image

Magnification:

More extensive and more realistic schematic:



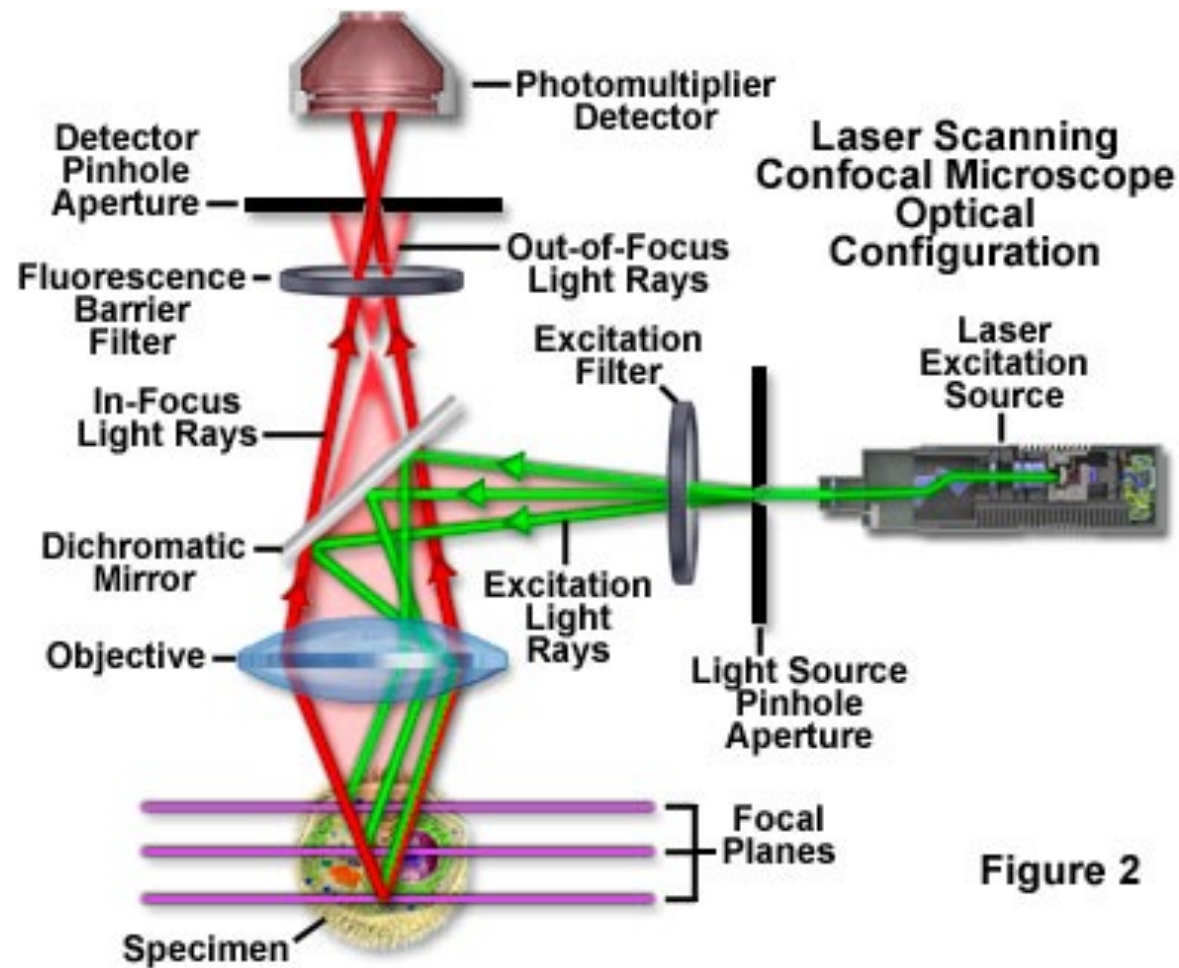
Microscope variations:

There are many variations and combinations with other techniques for microscopes in terms of

- Illumination
- Magnification
- Detectors
- Staining for improving contrast
- Fluorescence detection
- Etc

Lets look at some conceptual variations

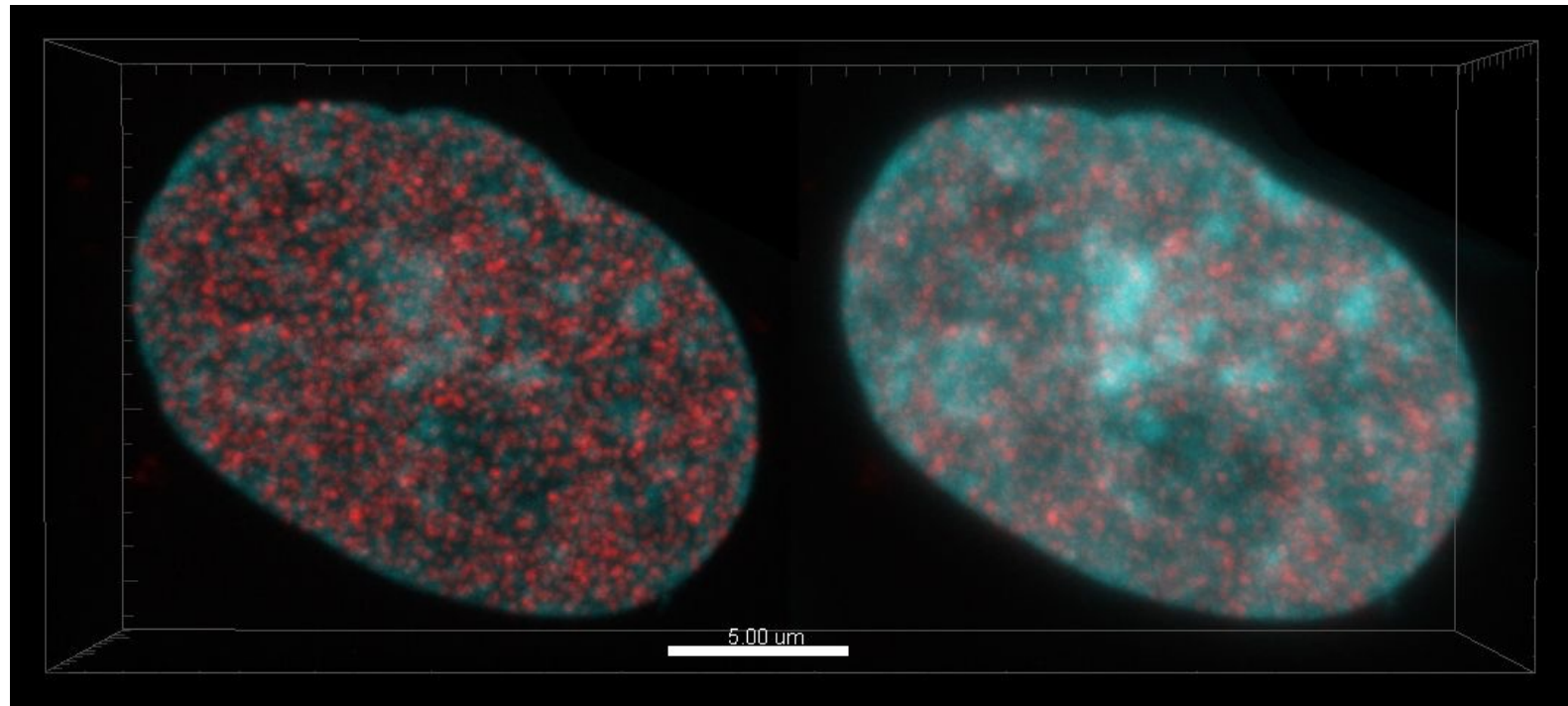
Microscope variations: Confocal microscope



Source: Olympus

Confocal microscopy: Example

- Optical cuts through cell nucleus with two different pinhole / aperture sizes
- Computer can generate 3D models. Fluorescence excitation can be used to highlight specific units
- <https://upload.wikimedia.org/wikipedia/commons/9/92/Combined-confocal-unconfocal.gif>



Phase contrast microscopy

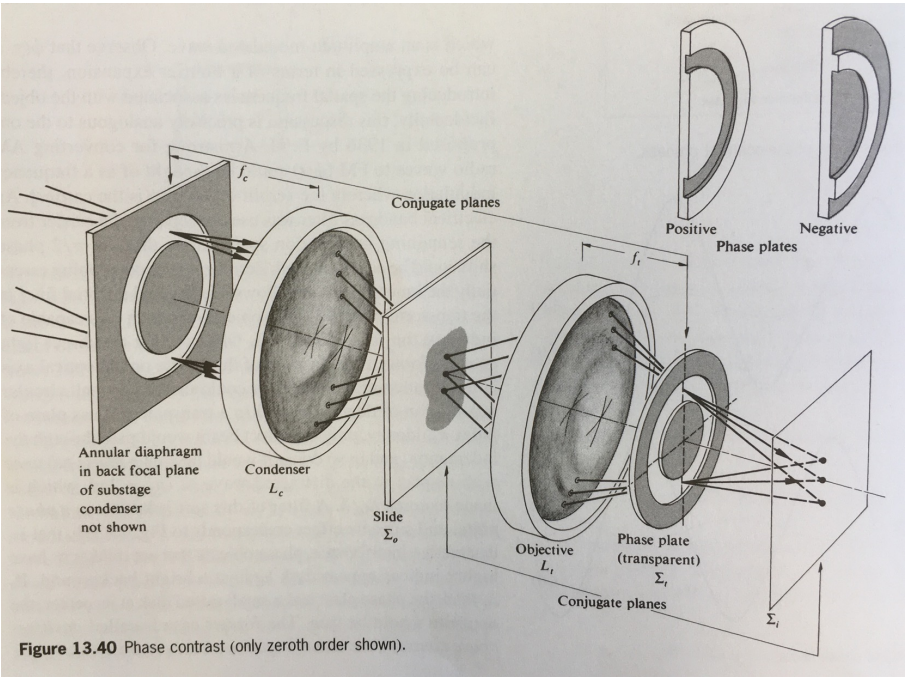
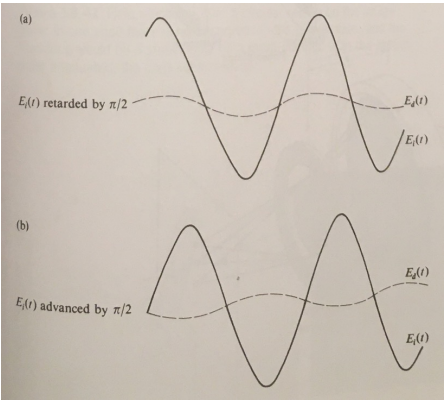
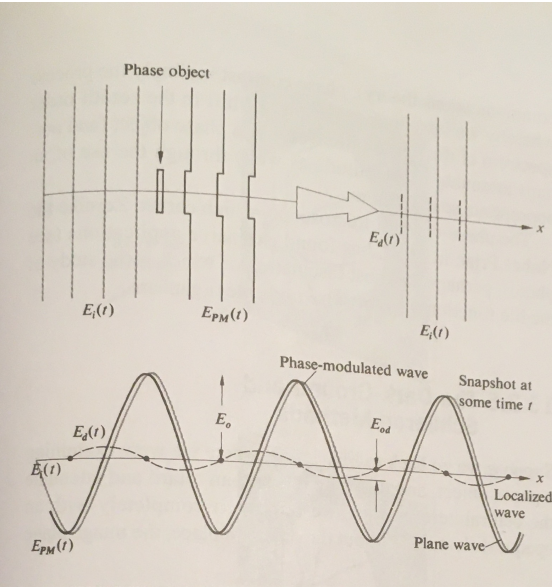


Photo from the Nobel Foundation archive.
Frits Zernike
Prize share: 1/1



Phase contrast microscopy

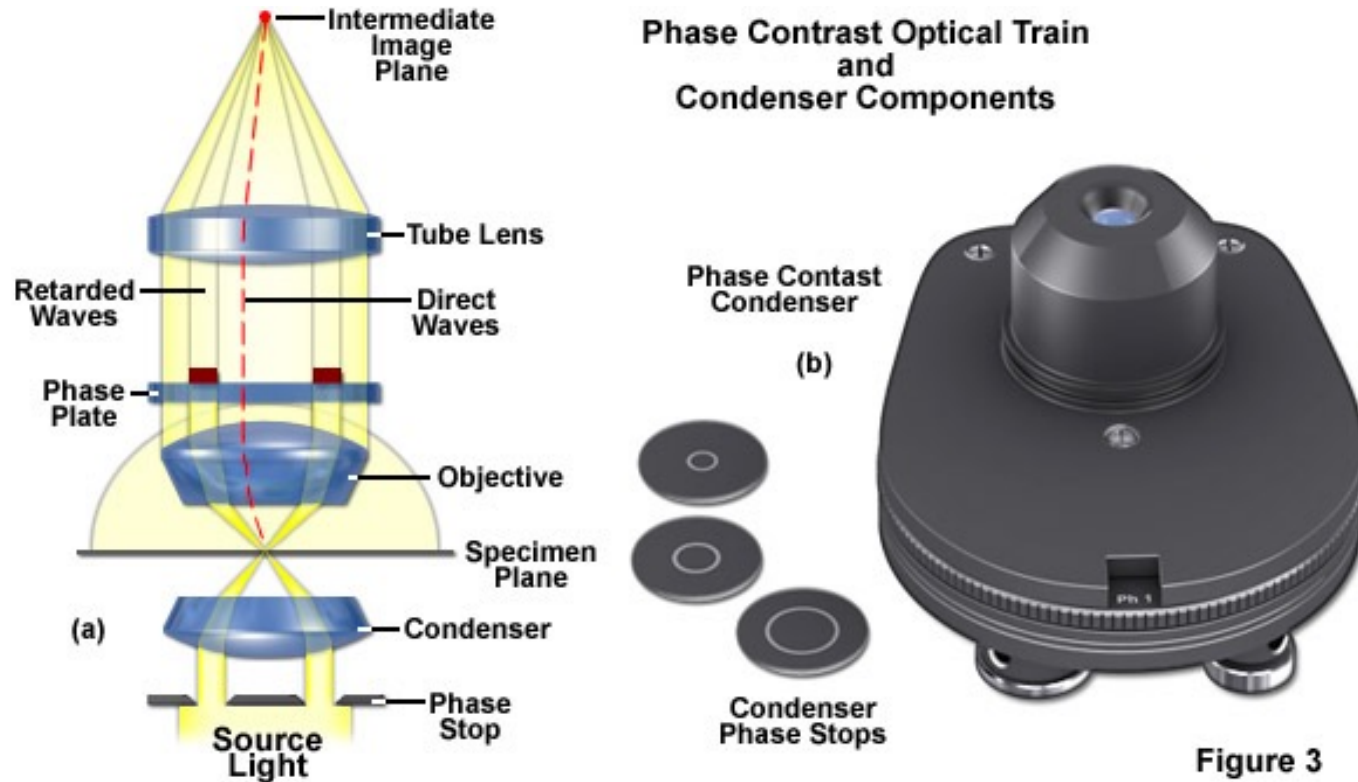
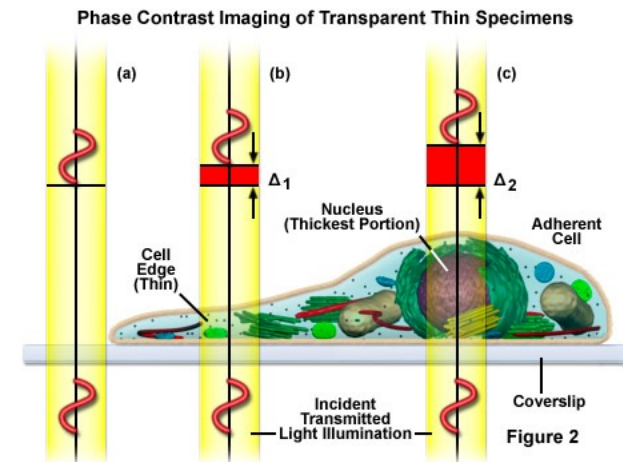
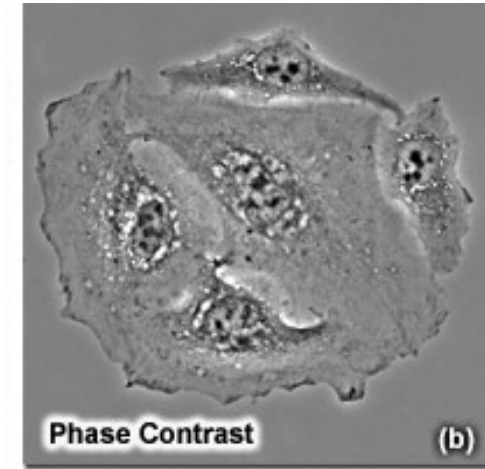
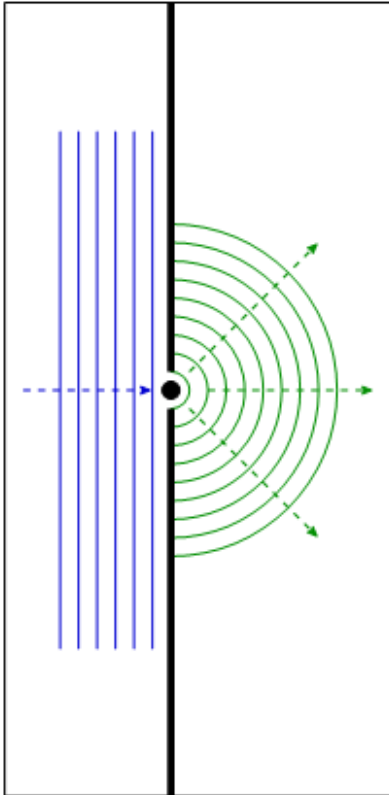


Figure 3

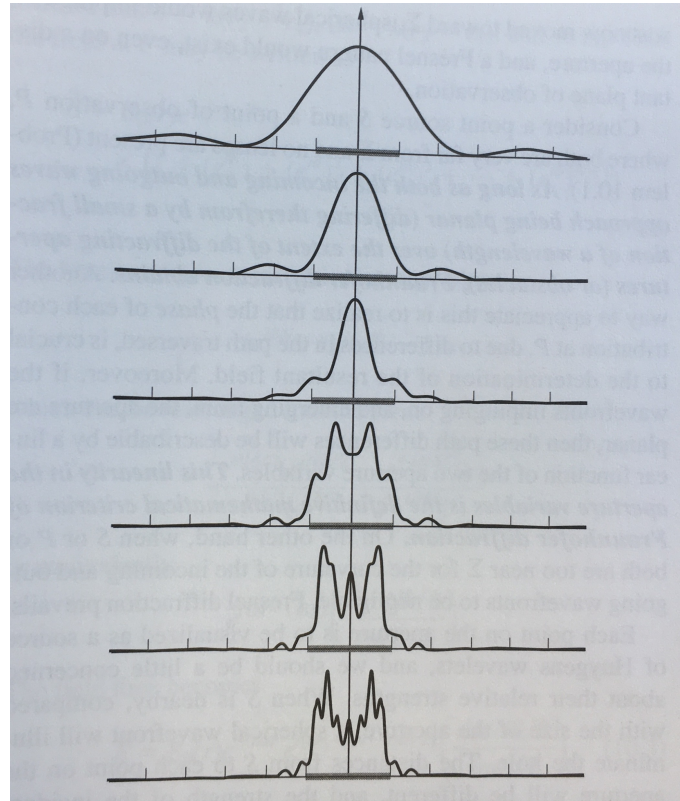


Resolution limit of a classical optical apparatus: Remember Huygens Fresnel principle, diffraction at a slit, Airy rings

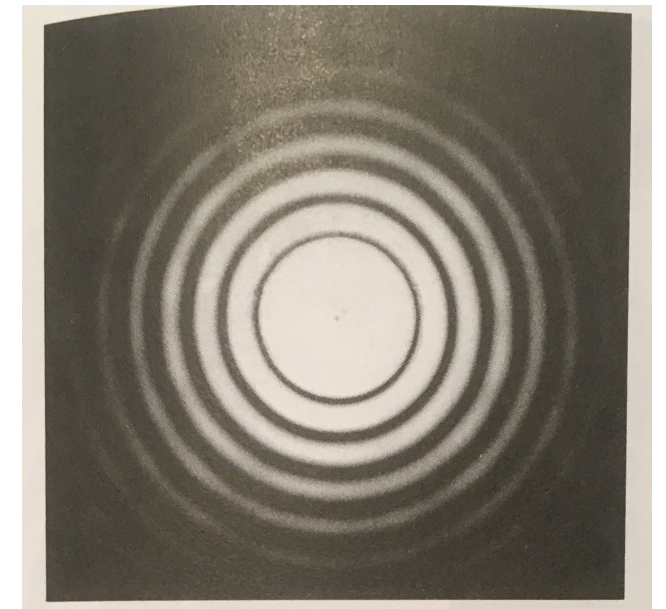
Diffraction at slit / aperture



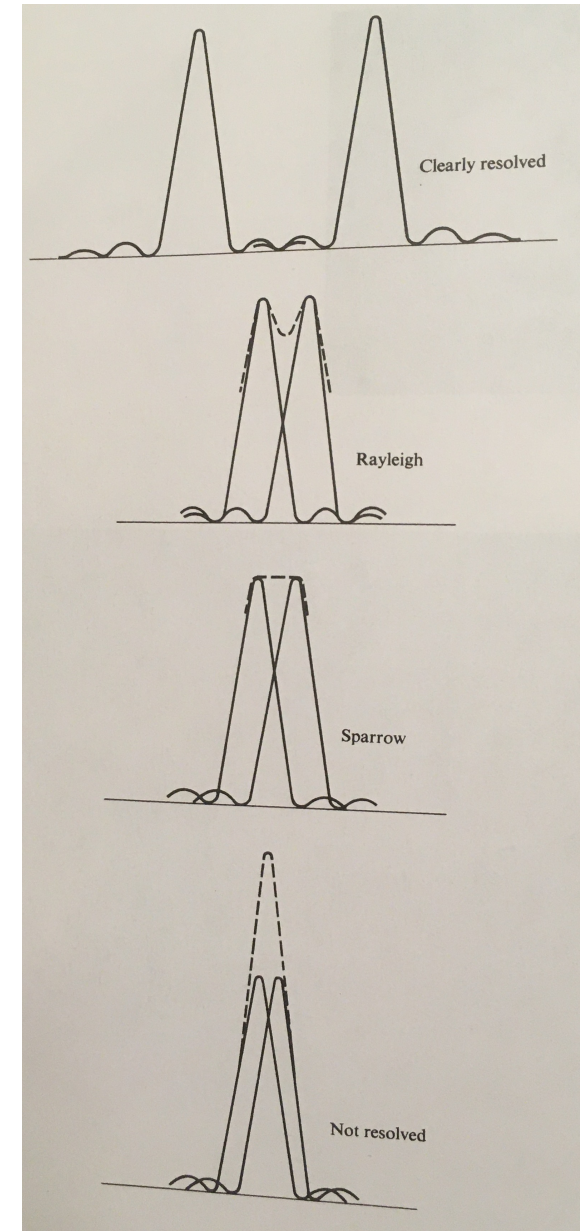
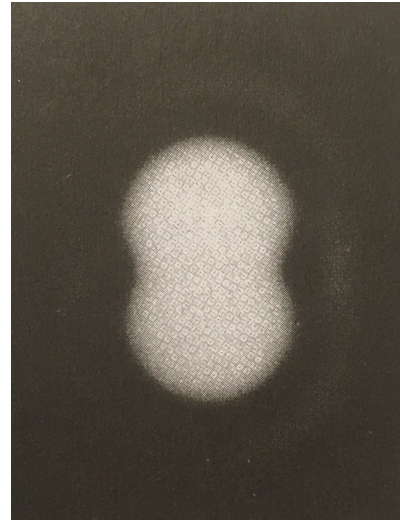
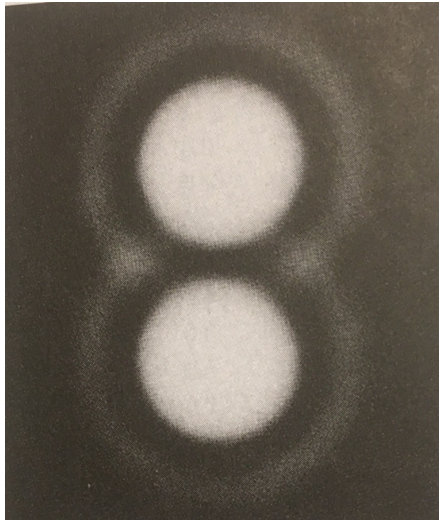
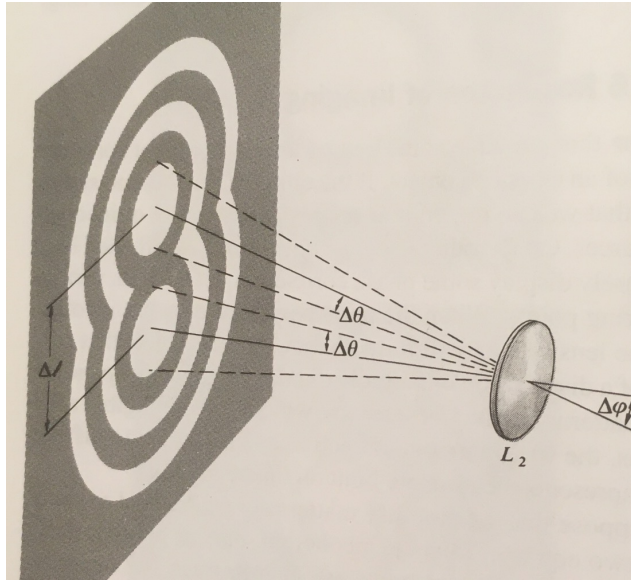
Fresnel and Fraunhofer regime



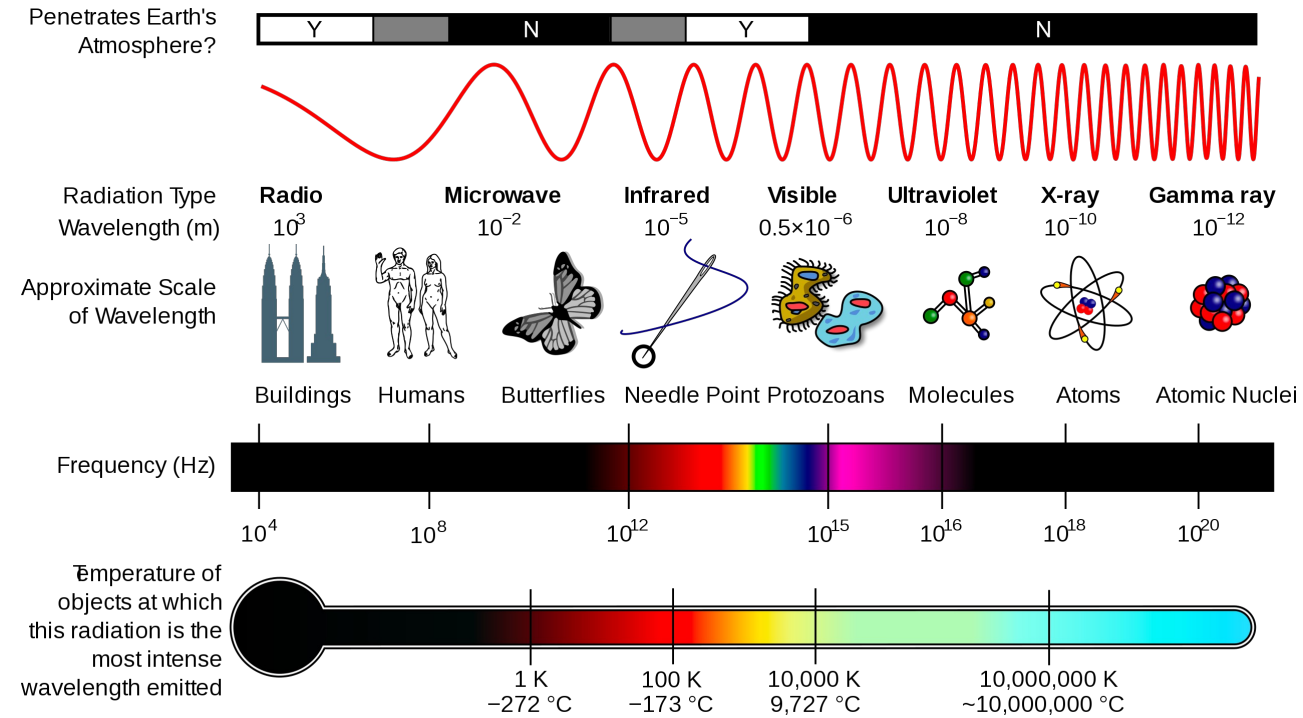
Airy rings of circular aperture



Resolution limit of a classical optical apparatus



How to beat resolution limit?



Shorter wavelength:

X-ray microscopy. No optics available. Later in course.

Electron: wave nature or particles

Smart ideas?

De Broglie: all matter has particle and wave nature!

- 1929 Nobel prize
- Example: Electrons



Electron microscopy

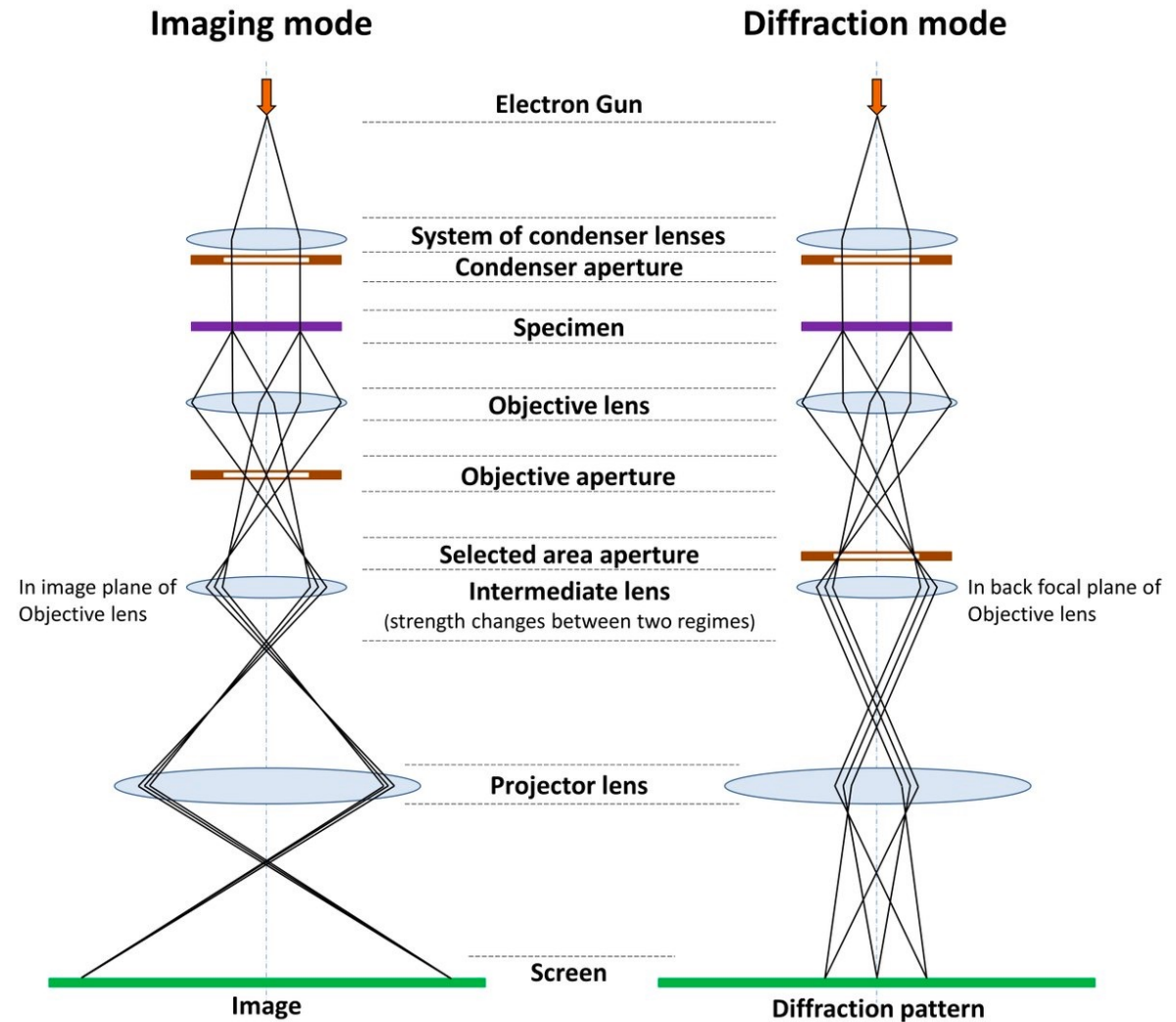
Electrostatic lenses akin optical lenses

Use short wavelength of (fast) electrons

Strong interaction with matter:
Thins samples

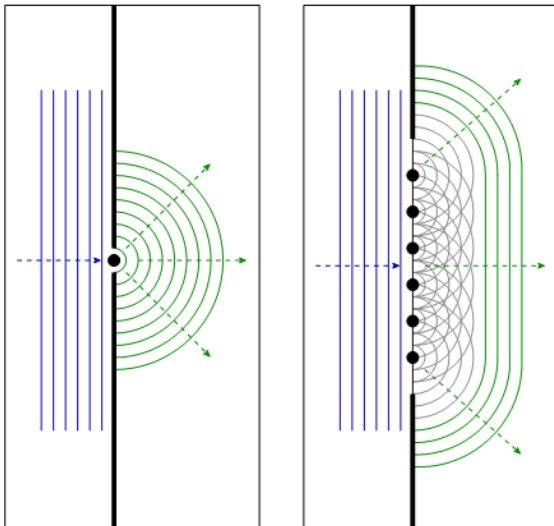
Need to use diffraction,
understand Fourier planes, etc

This is an entirely different
course but you need to know
the close relationship with all
optical concepts we have
discussed

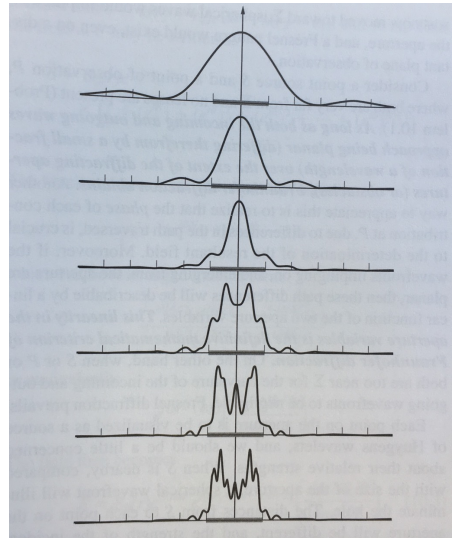


Overcoming the resolution limit: Near field approaches OR optics on the nanoscale

Small vs. real slit



Fresnel and Fraunhofer regime



Near field approach

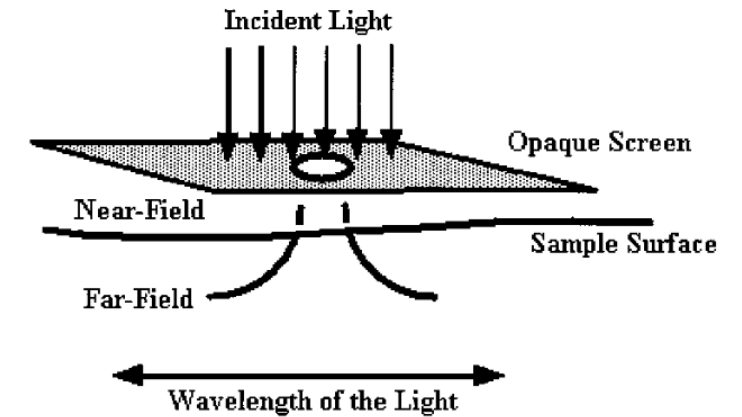
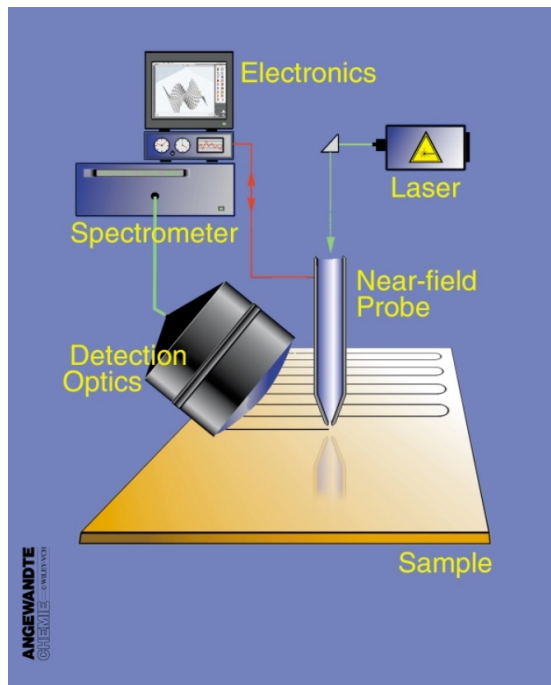


Figure 1. Schematic representation of Synge's idea for achieving subdiffraction limit spatial resolution. Incident radiation is passed through a small, subwavelength hole in an opaque screen. By the positioning of the screen close to the sample, the emerging radiation is forced to interact with the sample before diffracting out.

From: Dunn, Chem. Reviews 1999

SNOM – Scanning Near Field Optical Microscope



Zenobi and Deckert, Angew.
Chemie Review, 2000

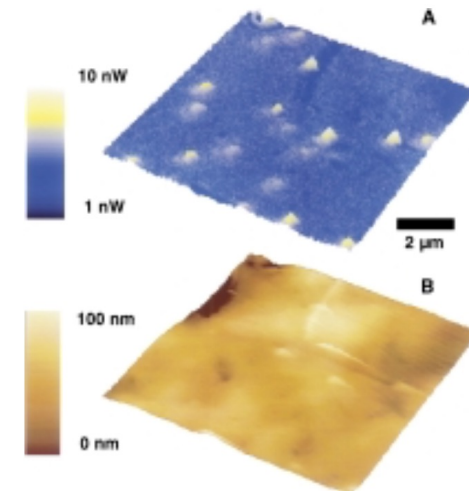
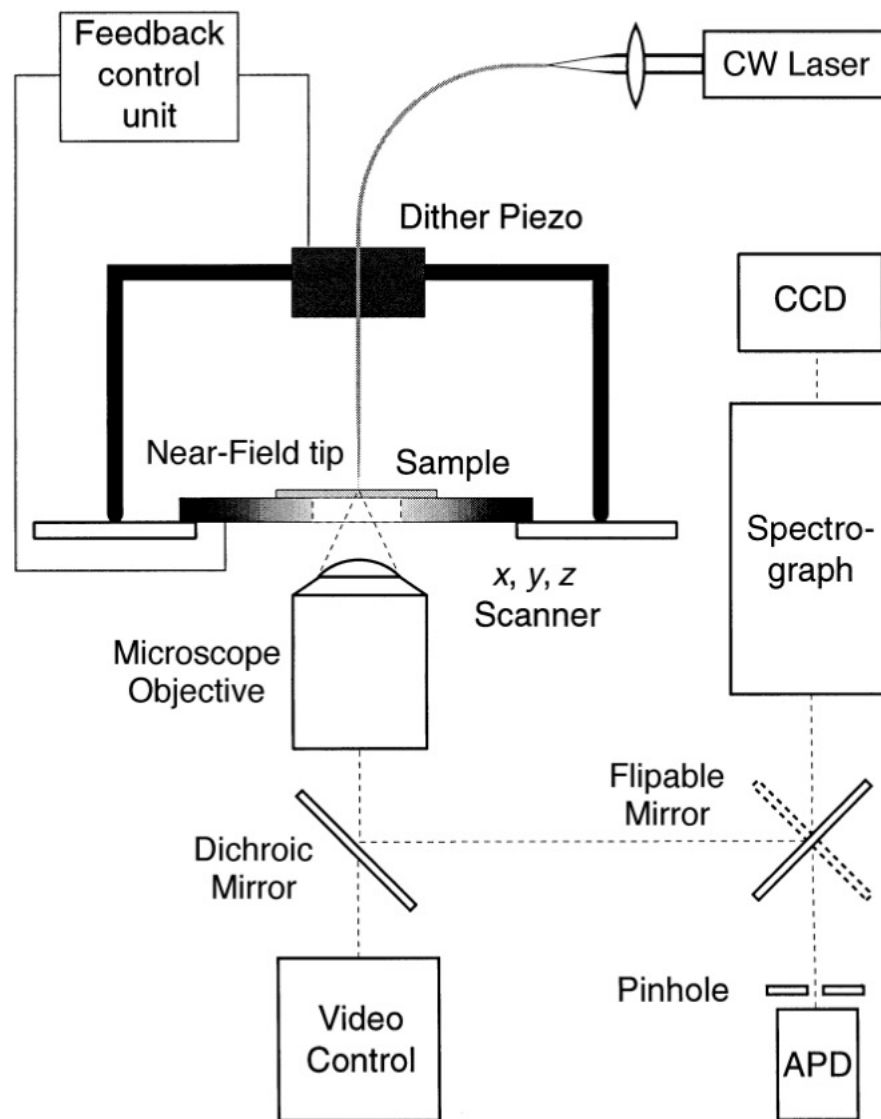


Figure 4. A) SNOM/fluorescence image and B) topographic image of a $10 \times 10 \mu\text{m}$ area of a PVB film containing fluorescent microspheres. The spheres had a diameter of 288 nm. The fluorescence was excited with the 488 nm line of an Ar-ion laser.

Increasingly fancy approaches

Reviews

A. Hartschuh

Optics on the Nanoscale

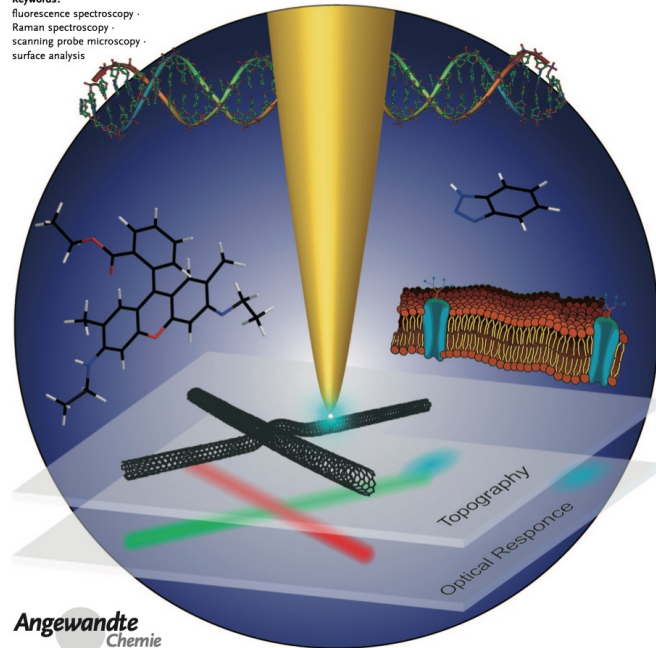
DOI: 10.1002/anie.200801605

Tip-Enhanced Near-Field Optical Microscopy

Achim Hartschuh*

Keywords:

fluorescence spectroscopy ·
Raman spectroscopy ·
scanning probe microscopy ·
surface analysis



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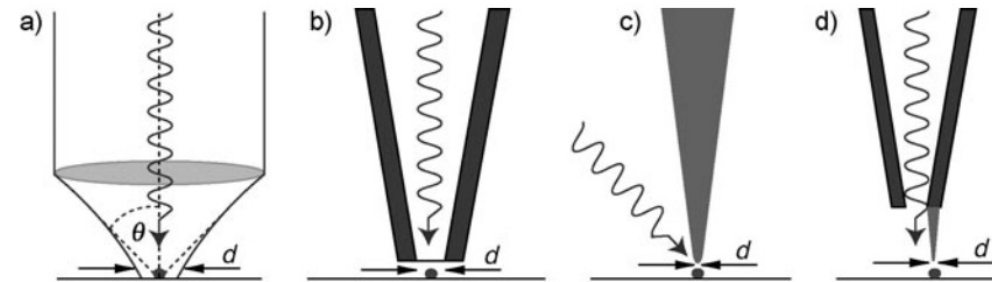
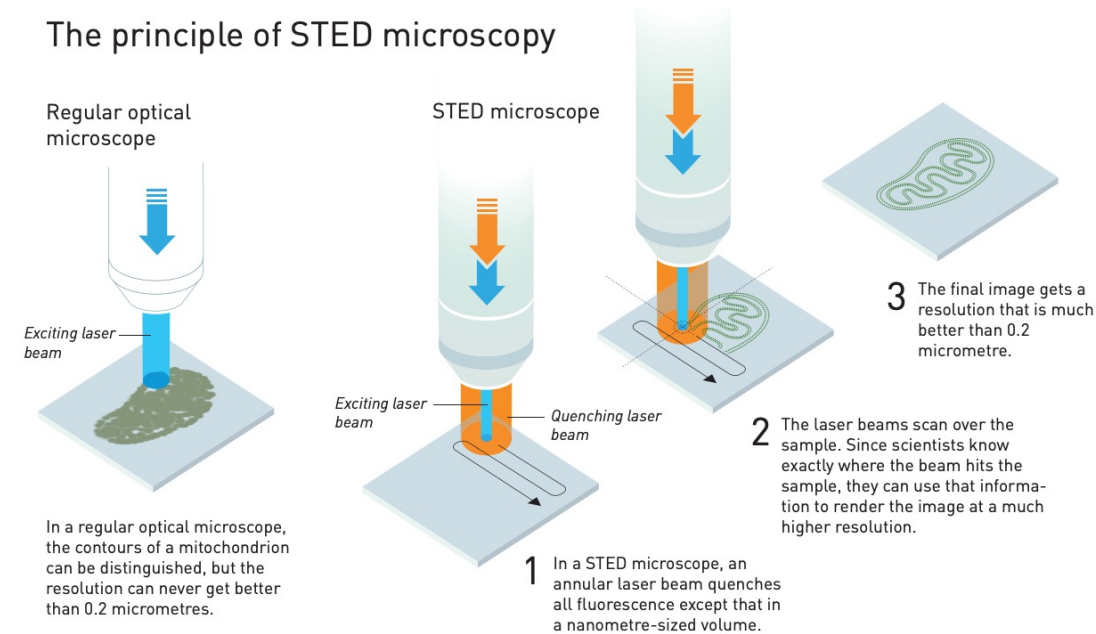


Figure 2. Focusing concepts: a) Far-field focusing using a lens. The angular frequency range of propagating waves $k_{x,\max}$ and thus the focus diameter, is limited by the aperture angle of the lens $k_{x,\max} = n \sin(\theta) 2\pi/\lambda$, with n being the refractive index and λ the wavelength of light. b) Aperture-type scanning near-field optical microscope (aperture-SNOM). c) Tip-enhanced near-field optical microscopy (TENOM). d) Tip-on-aperture (TOA) approach, which combines the advantages of (b) and (c).

Hartschuh, Ang. Chemie, Review 2008

Last comment: Remember session 1?

Recent examples of applications of light to chemistry



Eric Betzig, Stefan W. Hell and William E. Moerner are awarded the Nobel Prize in Chemistry 2014 for having bypassed a presumed scientific limitation stipulating that an optical microscope can never yield a resolution better than 0.2 micrometres. Using the fluorescence of molecules, scientists can now monitor the interplay between individual molecules inside cells; they can observe disease-related proteins aggregate and they can track cell division at the nanolevel.



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Eric Betzig
Prize share: 1/3



© Nobel Media AB. Photo: A. Mahmoud
Stefan W. Hell
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© Nobel Media AB. Photo: A. Mahmoud
William E. Moerner
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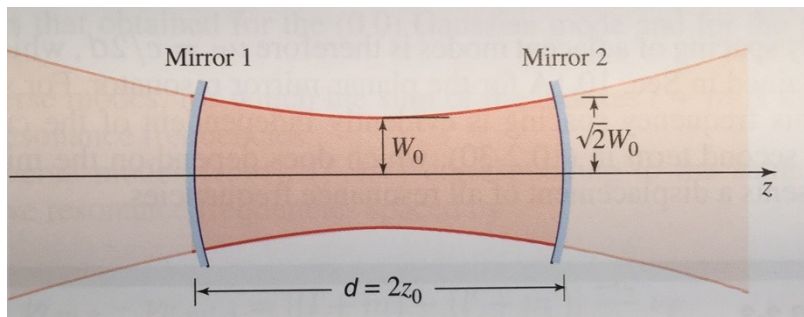
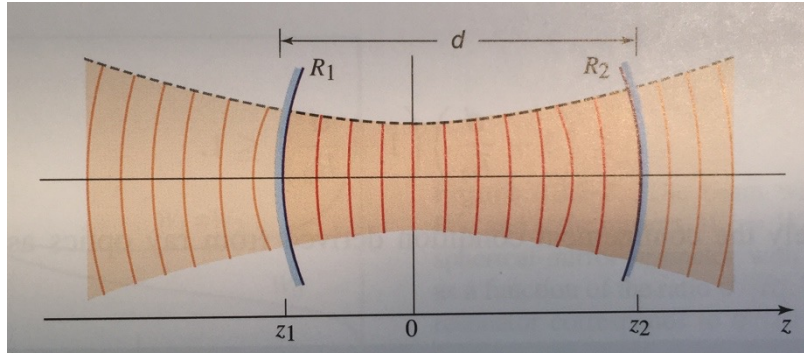
From Lecture 5: Absorption, emission, stimulated emission - overview

Spontaneous
emission

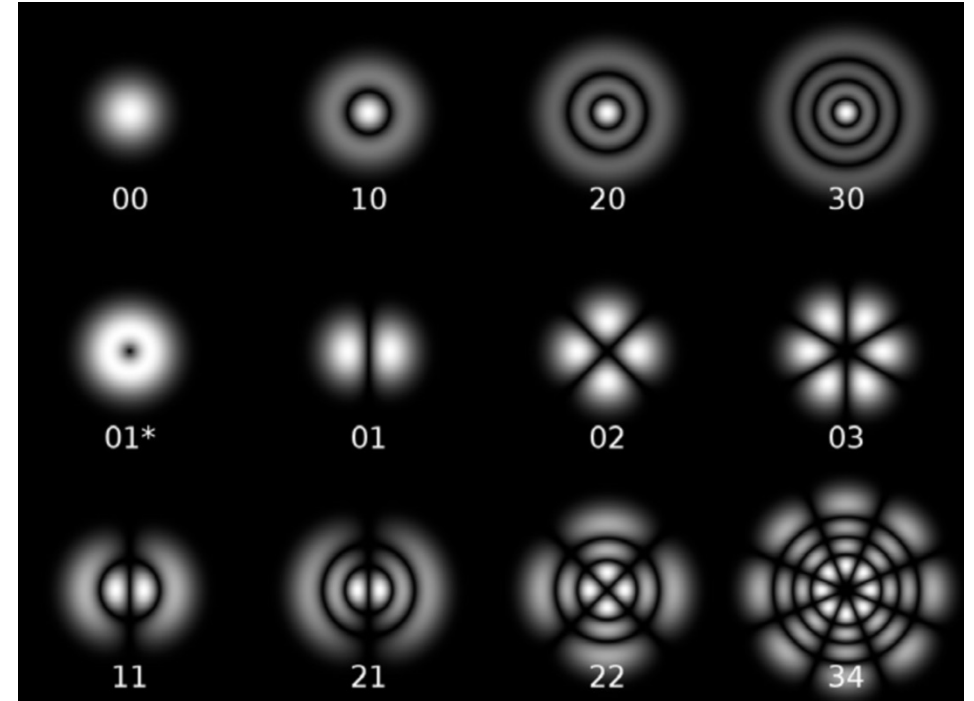
Stimulated
Emission

(Stimulated)
absorption

From Lecture 4: Gaussian Beam Resonator



Laguerre-Gaussian modes

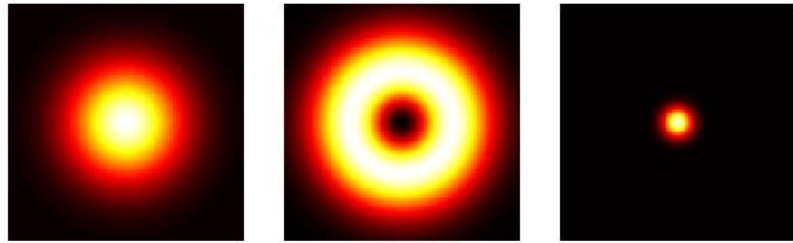


Stefan Hell: Stimulated Emission Depletion Microscope (STED)

- Based on Fluorescence Microscope
- Based on Laser Scanning Microscope

But with Important additional concepts:

Remember Gaussian, Bessel beams, focal spots



The end.