### Microscopy

Jan Kybic<sup>1</sup>

2011-2015

<sup>&</sup>lt;sup>1</sup>Using material from Davidson and Abramowitz: Optical Microscopy

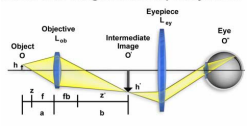
## Microscopy

Optical microscopy – since 17th century; Jensen, van Leeuwenhoek, Galilei, . . .



### Finite-Tube Length Microscope

Finite-Tube Length Microscope Ray Paths

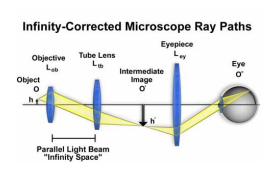


- magnification of the objective  $\frac{b}{a}$
- ► magnification of the eyepiece  $\frac{25 \text{ cm}}{f_{\text{eyepiece}}}$
- ▶ thin-lens equation

$$\frac{1}{a} + \frac{1}{b} = \frac{1}{f}$$

- narrow range of image distances
- specifically corrected optical systems with matching eyepieces

### Infinite-Tube Length Microscope

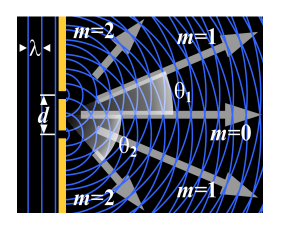


- Modern design (since 1980s)
- ▶ Objective magnification determined by  $\frac{f_{\text{tb}}}{f_{\text{ob}}}$
- Infinity space to add polarizers, prisms, retardation plates. . .
- Independently changeable objective and eyepiece

#### **Image Formation**

- Direct/undeviated light
- Deviated/diffracted light, out of phase
- Constructive/destructive interference

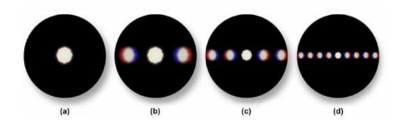
#### Diffraction



Position of maxima:

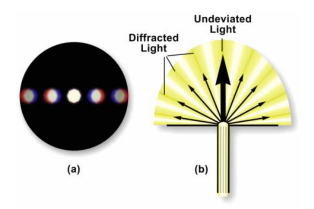
$$d\sin\theta = n\lambda, \quad n \in \mathbf{Z}$$

### Line Grating Diffraction Patterns



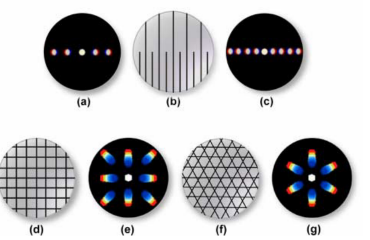
- ▶ line phantom
- close diaphragm
- ▶ telescope, observe the rear focal plane of the objective
- ▶ (a) no phantom, (b)  $10\times$ , (b)  $40\times$  (higher NA), (c)  $60\times$  (highest NA)
- ▶ 0<sup>th</sup> order, 1<sup>st</sup> order image

#### Diffraction

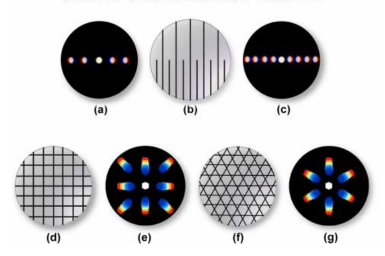


- constructive/desctructive interference
- specimen = superposition of complex gratings (Ernst Abbe)
- ▶ to resolve image, at least 0<sup>th</sup> order and 1<sup>st</sup> order images must be captured
- ▶ more orders captured → better accuracy

#### Slit and Grid Diffraction Patterns

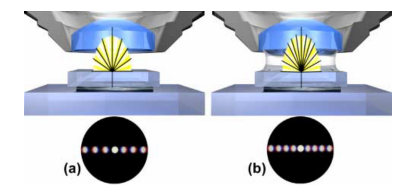


#### Slit and Grid Diffraction Patterns



- ▶ Diffraction patterns behave like Fourier transforms of the sample
- Fourier optics

#### Immersion optics



- High refractive-index media (immersion oil) reduce diffraction angle
- ightharpoonup ightharpoonup More orders are captured
- ightharpoonup ightharpoonup Better image

#### Resolution limit

Rayleigh equation:

$$d \approx 1.22 \frac{\lambda}{2 \text{ NA}}$$

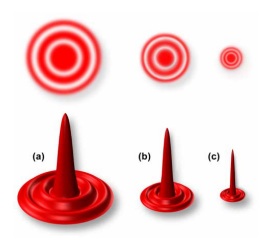
To improve resolution, use:

- Big lenses (big NA)
- Short wavelength (blue)

#### Numerical aperture:

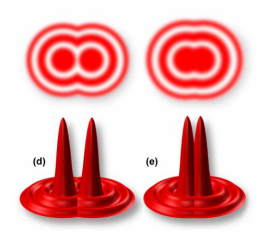
- ▶  $NA = n \sin \theta$ , with half-cone angle  $\theta$
- ▶ f-number  $N = f/D \approx 1/(2NA)$ , written as f/N

# Airy disks



- ▶ NA increases left to right.
- ► Impulse response (PSF)

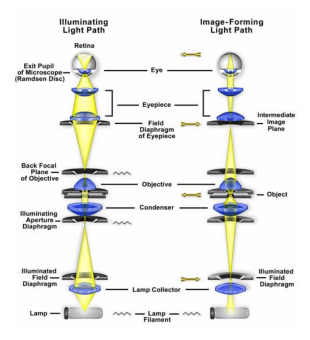
# Airy disks (2)



Resolution limit.

#### Köhler illumination

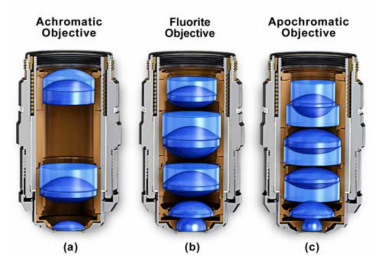
- Focused lamp image is projected to the diaphraghm of a condenser.
- Field diaphraghm controls width of the light bundle.
- Apperture diaphraghm controls the light intensity. Trade-off between diffraction artifacts and glare.
- Light is not focused on the specimen, illumination is homogeneous.
- The focal point of image-forming rays is at the level of the specimen.



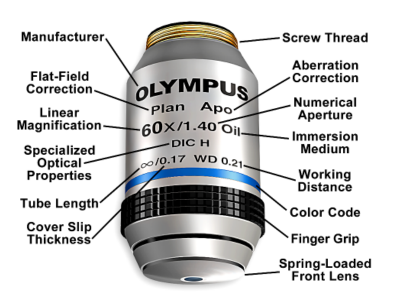
#### **Optical Aberrations**

- Geometric aberrations
  - Spherical rays on axis and far from the axis do not converge to the same point. Blurred images.
  - ► Flat-field because lenses are curved, the image is curved. Center and off-center not simultaneously in focuss.
- Chromatic aberrations rays of different color do not converge to the same point

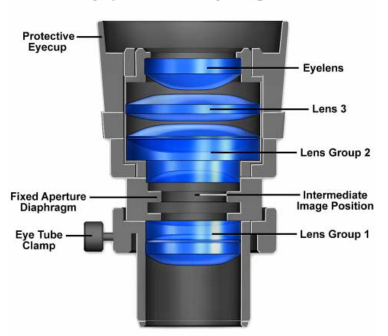
#### **Optical Correction in Objectives**



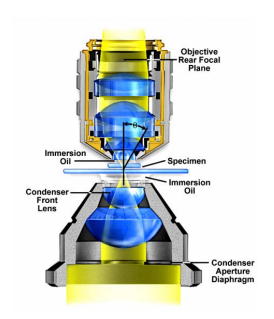
## Objective Specifications



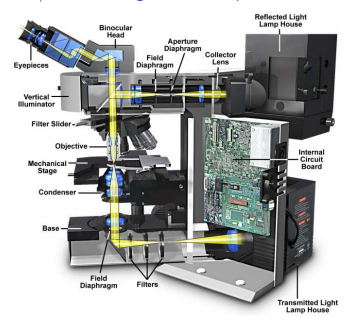
#### **Eyepiece Cutaway Diagram**



#### Condenser

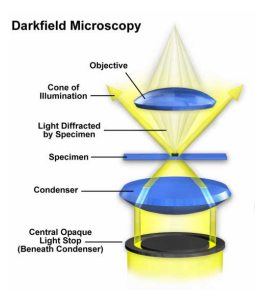


#### Transmitted/Reflected light microscope



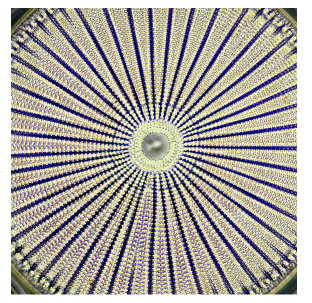
## Contrast enhancing techniques

- Dark field microscopy
- Rheinberg illumination
- Phase contrast microscopy
- Polarized light
- Hoffman modulation
- ▶ Differential interfence contrast



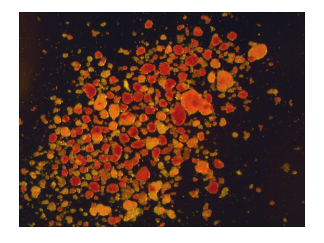
For unstained objects. Appear bright on dark background.

# Darkfield microscopy (2)

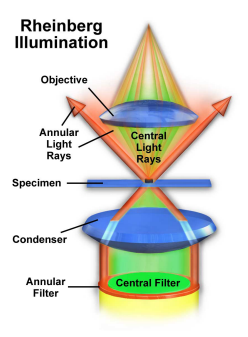


Arachnoidiscus ehrenbergi

## Darkfield microscopy (3)

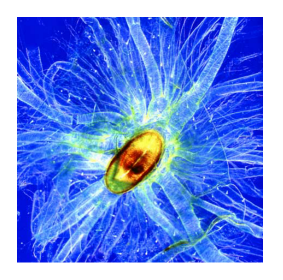


Langerhans islets

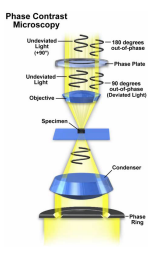


Color annular filters instead of the darkfield stop.

# Rheinberg illumination (2)



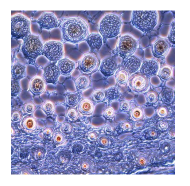
silkworm larva



Frits Zernike (1930s, Nobel price 1953). Show differences in phase/refractive index.

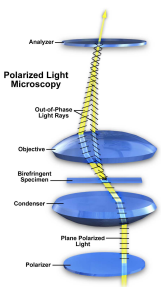
Interference. Slow down/Speed up. direct light  $\rightarrow$  bright/dark contrast

## Phase contrast microscopy (2)



mouse hair cross-section

## Polarized light microscopy

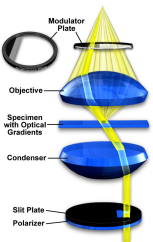


different refractive indices for different polarizations

# Polarized light microscopy (2)



#### Hoffman Modulation Contrast Microscopy

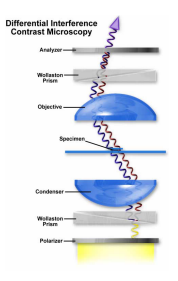


Robert Hoffman (1975). For living and unstained specimens. Detects optical gradients. Image intensity proportional to the derivative of the optical intensity of the specimen.

# Hoffman modulation contrast (2)

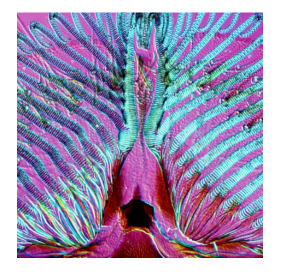


Dinosaur bone



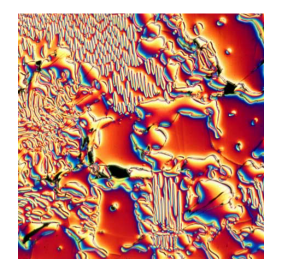
Detects differences in optical paths between two close slightly offset rays (shear).

# Differential interference contrast microscopy (2)



Mouth part of a blowfly.

# Differential interference contrast microscopy (3)

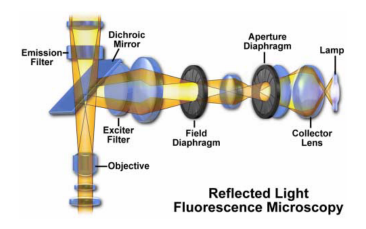


Defects in ferro-silicon alloy.

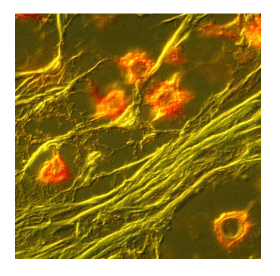
#### Fluorescence microscopy

- fluorescent dyes
- multiple sensing channels/filters
- multiple light sources visible, UV

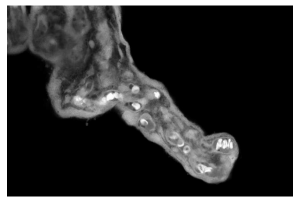
## Fluorescence microscopy (2)



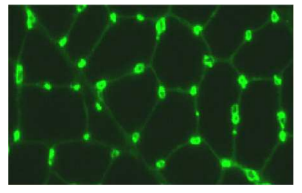
#### Fluorescence microscopy (3)



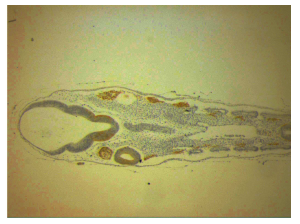
cat brain tissue infected with Cryptococcus



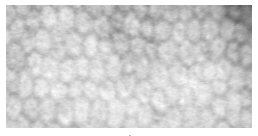
placenta cross-section



muscle capillaries



crocodile ear slice



retina

#### Advanced microscopy techniques

#### 3D microscopy

- Confocal microscopy
- Optical coherence tomography (OCT)
- Multiphoton / two-photon microscopy

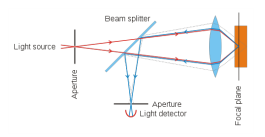
#### High resolution microscopy

- Stimulated emission depletion (STED)
- Stochastic optical reconstruction microscopy (STORM)
- Photo-activated localization microscopy (PALM)

#### Electron microscopy

- Scanning electron microscopy (SEM)
- Serial section EM (3D)
- Focused ion beam (FIB) (3D)
- Transmission electron microscopy (TEM)

#### Confocal microscopy



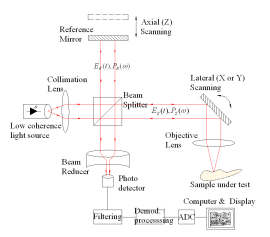
- Very good resolution
- Very thin focal plane 3D imaging
- Confocal laser scanning
- Scanning slow

#### Confocal microscopy example



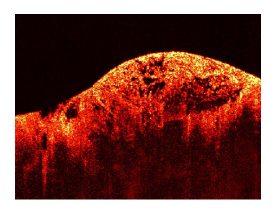
Tetrachimena

## Optical coherence tomography (OCT)



- ► 3D imaging
- ► Interferometry
- More penetration than confocal, especially near infrared

# OCT example



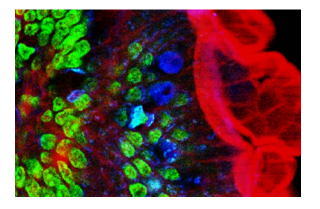
Sarcoma

#### Two-photon microscopy

- ▶ two low-energy photons → fluorescence
- high-flux laser
- better penetration
- reduced phototoxicity
- better background suppression

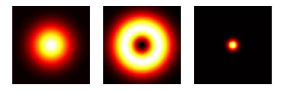
Maria Goeppert-Mayer (1931)

## Two-photon microscopy (2)



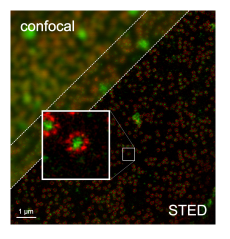
Two-photon excitation microscopy of mouse intestine. Red: actin. Green: cell nuclei. Blue: mucus of goblet cells. [Wikipedia]

#### Stimulated emission depletion (STED)



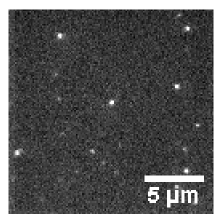
- excitation subpicosecond laser impulse
- depletion pulse around the focal spot, stimulating the emission
- fluorescence at the focal spot remains
- ▶ resolution  $2 \sim 80 \, \text{nm}$
- ► Hell and Klar, 1999. Hell awarded the Nobel Prize in Chemistry in 2014

## Stimulated emission depletion (STED) (2)



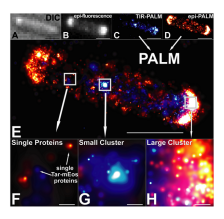
STED versus confocal

# Stochastic optical reconstruction microscopy (STORM/PALM)

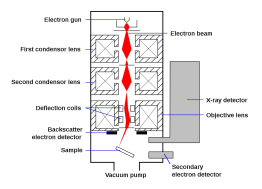


- sparse fluorophores localized by PSF fitting
- combine many images
  PALM photobleaching, STORM reversible switching

# Stochastic optical reconstruction microscopy (STORM/PALM)

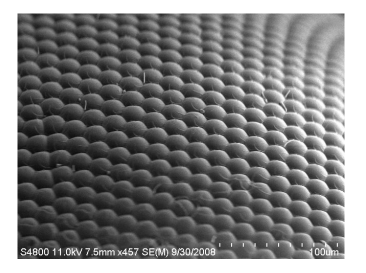


#### Scanning electron microscopy (SEM)



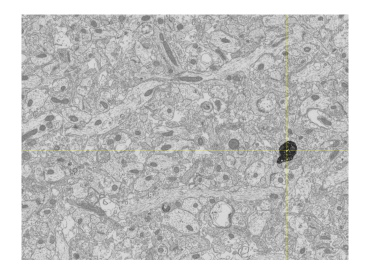
- Excellent resolution (a few nm)
- ▶ Needs vaccuum. Preparation gold coating, osmium staining, cryofixation.

#### SEM example



Fly eye

#### FIB example



▶ Focused ion beam for slice cutting. True 3D

## Microscopy — digitalization & automation

- CCD cameras
  - supercooled
  - superresolution
- Moveable specimen tray
  - Auto-focusing
  - Automated acquisition, mosaicking
- Automatic processing

#### Microscopy

- Advantages
  - ► High-spatial resolution
  - Colour and texture information
  - Affordable (optical microscopy)
  - ▶ Proven technique large body of experts available
- Disadvantages
  - Difficulties of in-vivo observations
  - Mostly 2D
  - Missing large-scale perspective