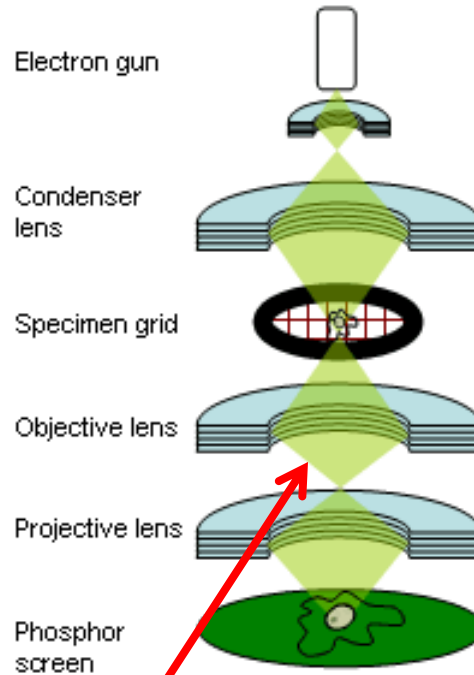
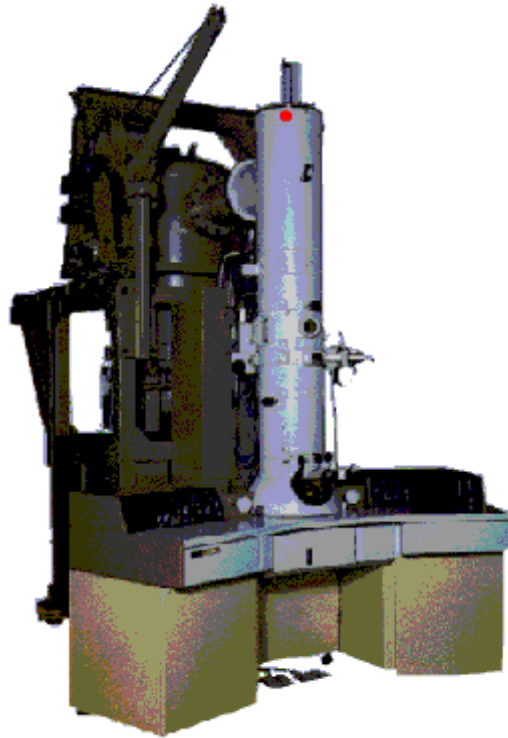


Cryo-Electron Microscopy

Pavel Plevka

Transmission Electron Microscope



Vacuum!

Electron source:

Thermal emission from heated cathode

Focussing:

Electro-magnetic Lenses

Detection:

Phosphor screen or CCD camera (former times: negative)

Pro & Con of cryo-EM

Pro

- Short wavelength => high resolution
- Strong interaction with materials => good contrast
- Electromagnetic lenses => standard optics (in contrast to X-ray crystallography)
- High intensity is easy to produce
- Inner structure of biomolecules is accessible

Con

- High vacuum requires special treatment of sample
- Sample has to be thin to avoid 100% absorption
- Electron beam damages biological samples => short measurements => low contrast of biomolecules

Microscope Optics

In order to see an “object” which is too small to be seen by our eyes, one needs to magnify the image. An example of magnifying an image by a lens is illustrated in Fig. 2.1:

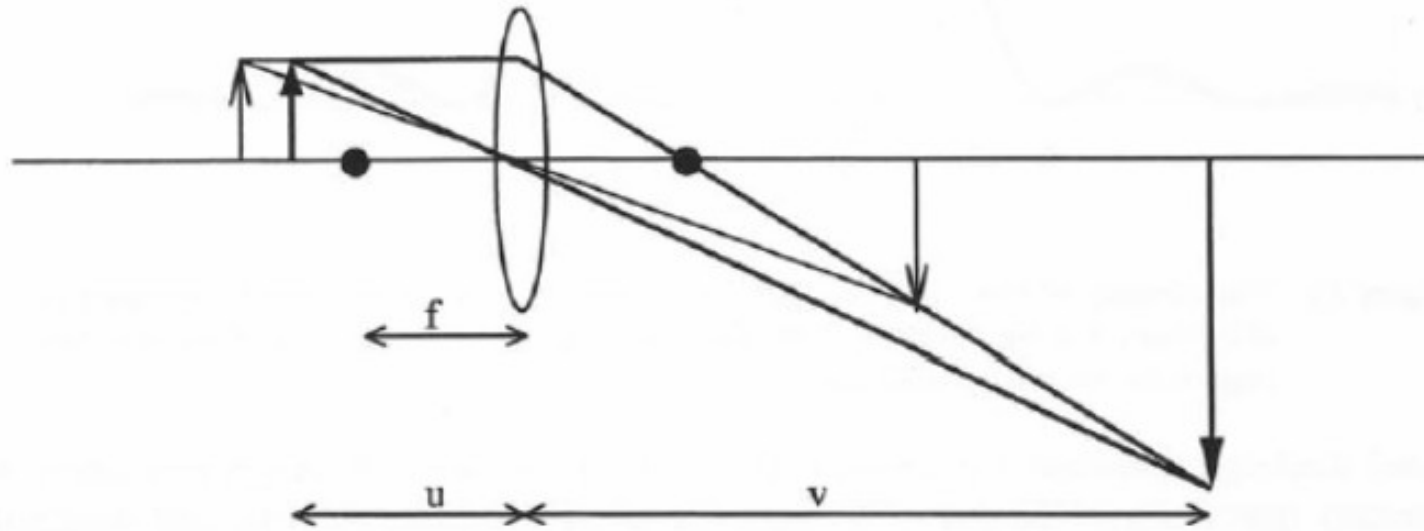
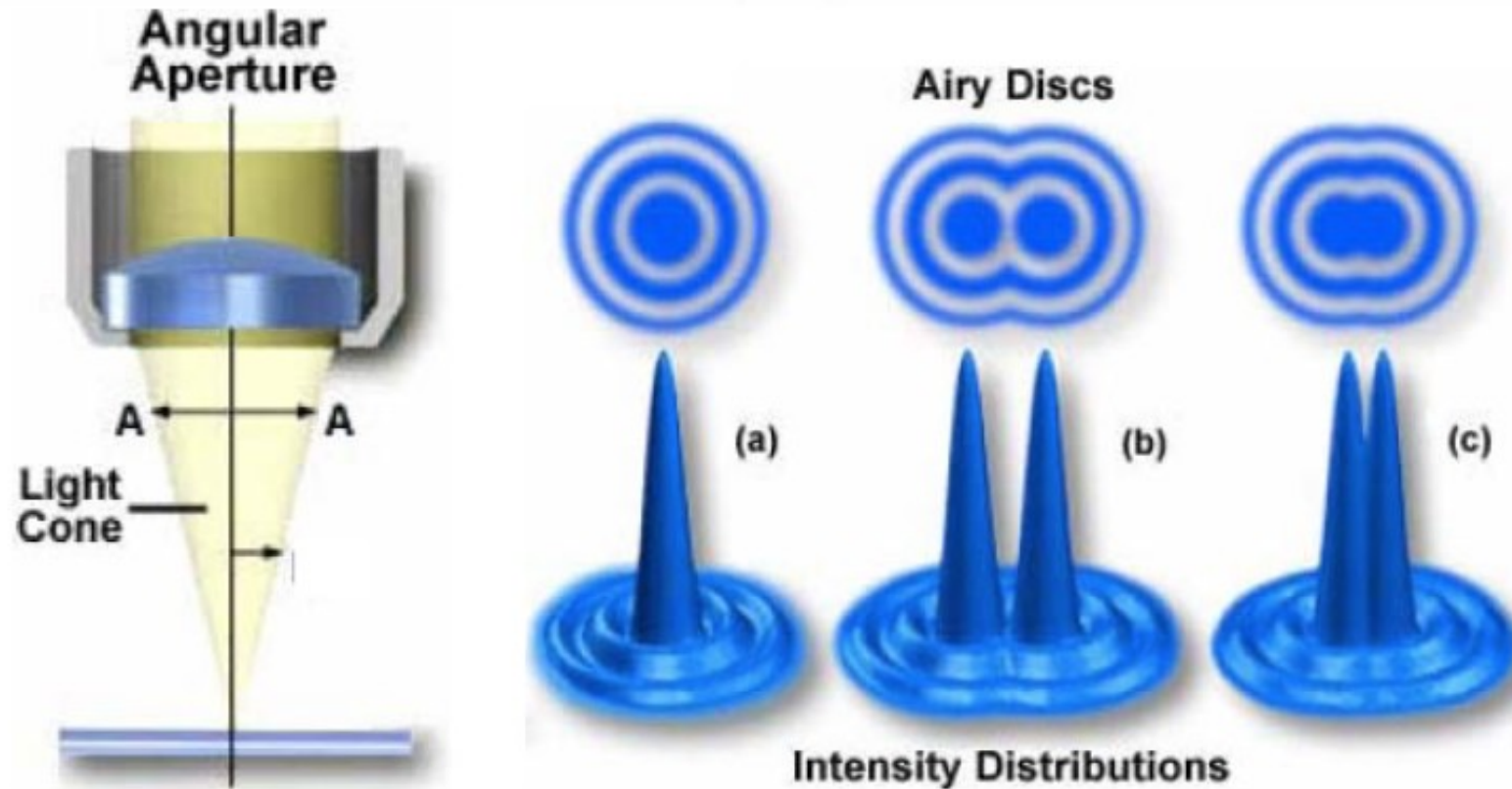


Figure 2.1 Ray diagram illustrating the formation of images by a lens. f = focal length, u = the distance between the object and the lens, v = the distance between the image and the lens.

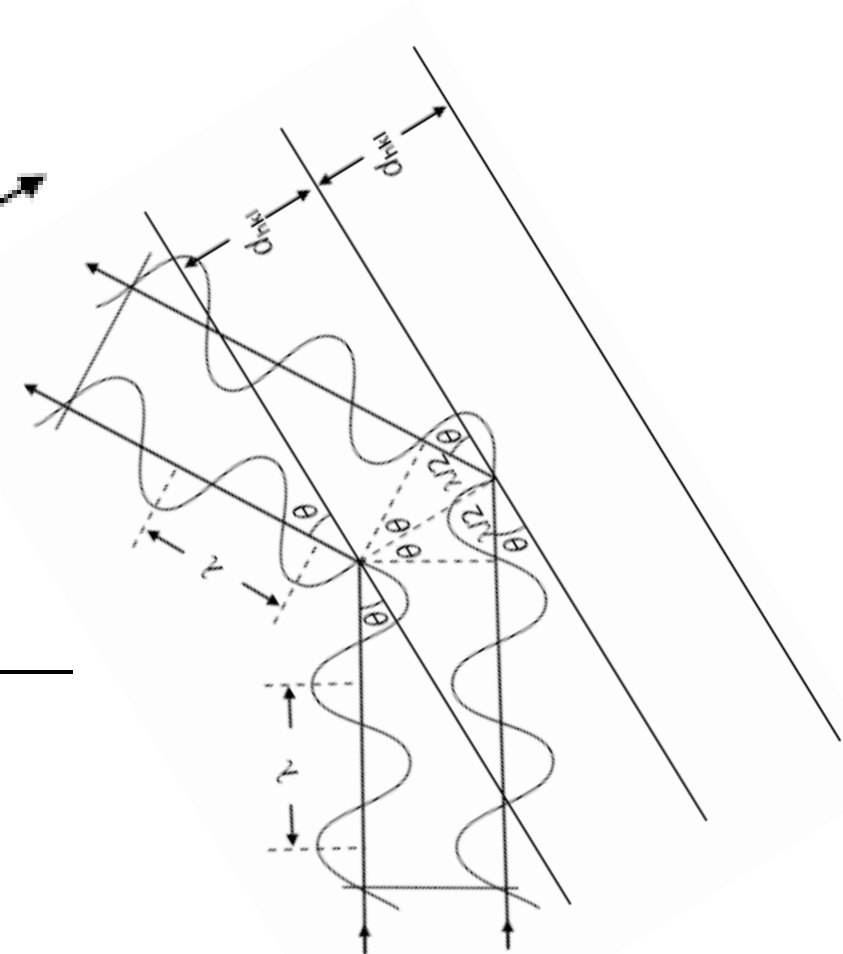
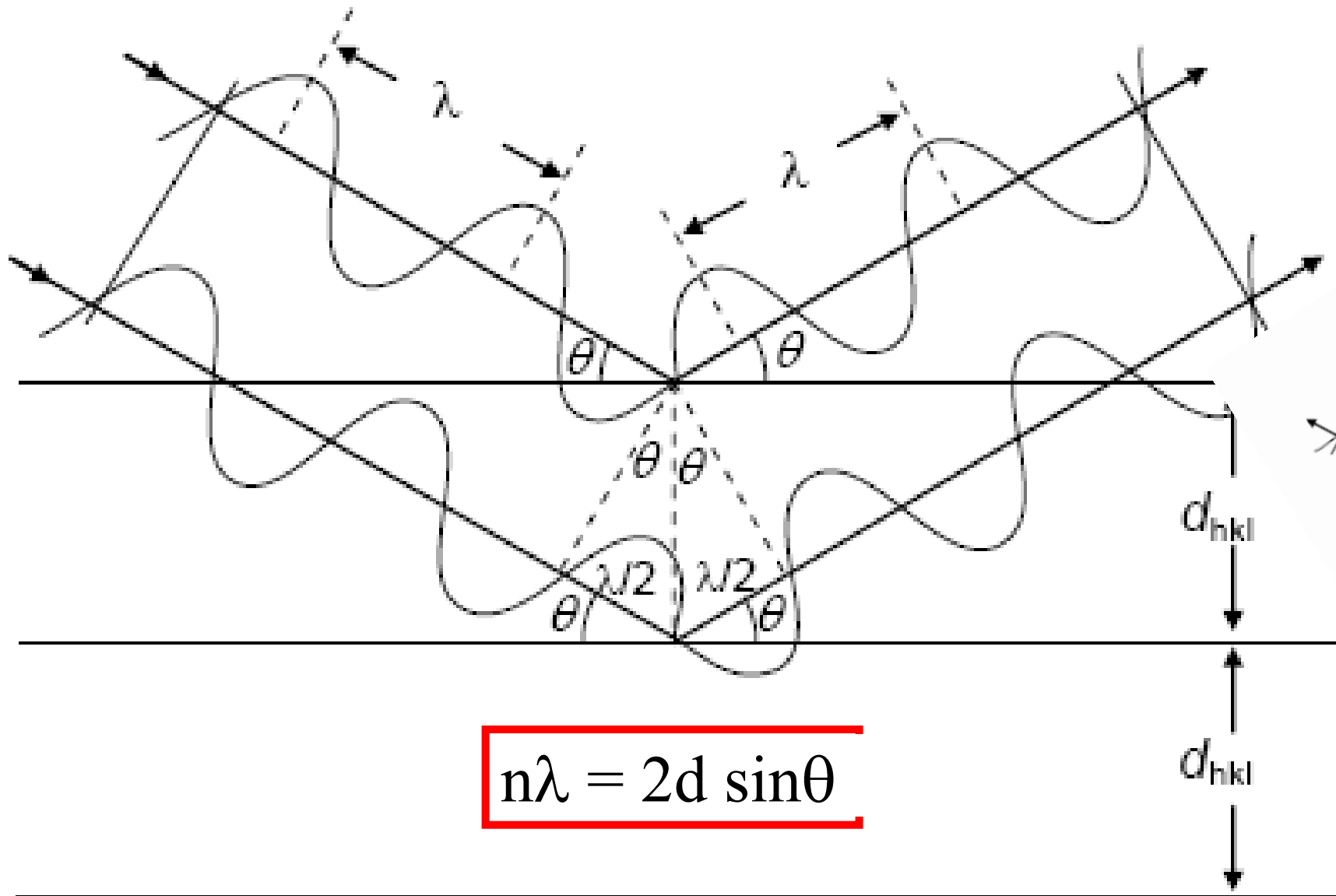
Optical Resolution

An image cannot be endlessly enlarged due to the limit of the resolution. "Resolution" is the closest distance between two points on the object which can be clearly seen through the microscope to be separate entities.



The central maximum of the Airy patterns is often referred to as an Airy disk, which is defined as the region enclosed by the first minimum of the Airy pattern and contains 84 percent of the luminous energy.

Bragg's law



Rayleigh criterion

the intensity maximum of the Airy disc from one point coincides with the first minimum of the Airy disc from the second point, then the two points can be just resolved. The Rayleigh resolution can be derived from diffraction theory to be:

$$r = \frac{d}{2} = \frac{0.61\lambda}{\mu \sin \alpha} \quad (2-2)$$

where λ is the wavelength of the light and μ is the refractive index of the medium between the object and objective lens. α is the semi-angle above which the light is stopped by the aperture, see in Fig. 2.3.

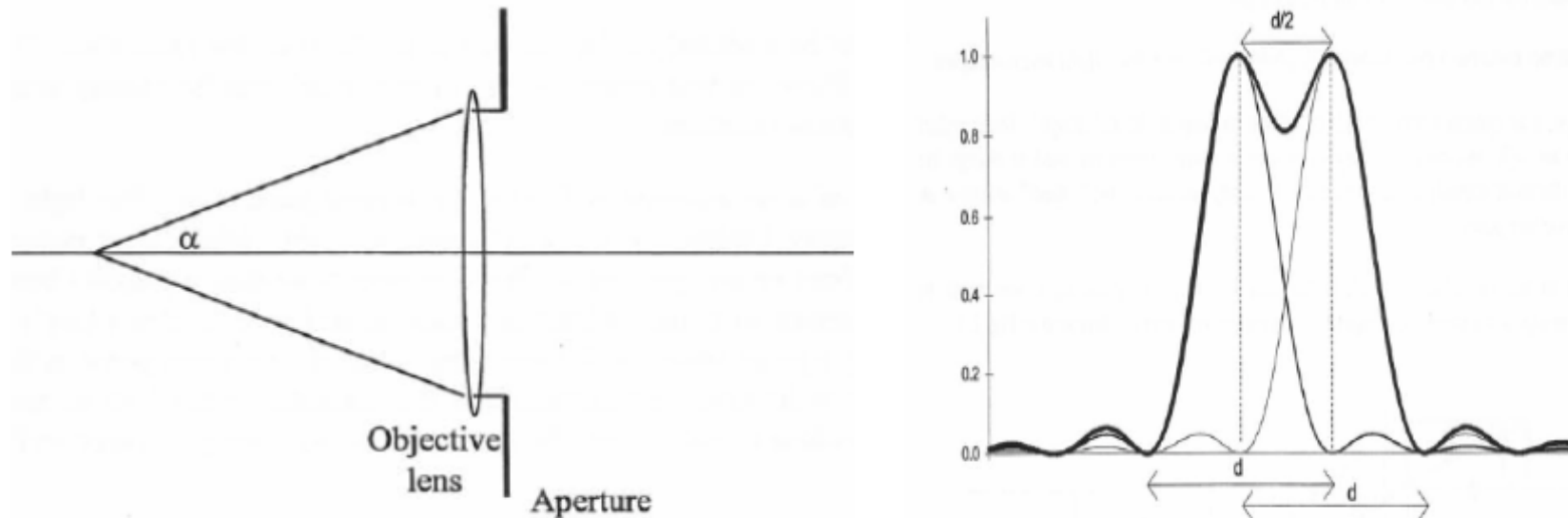
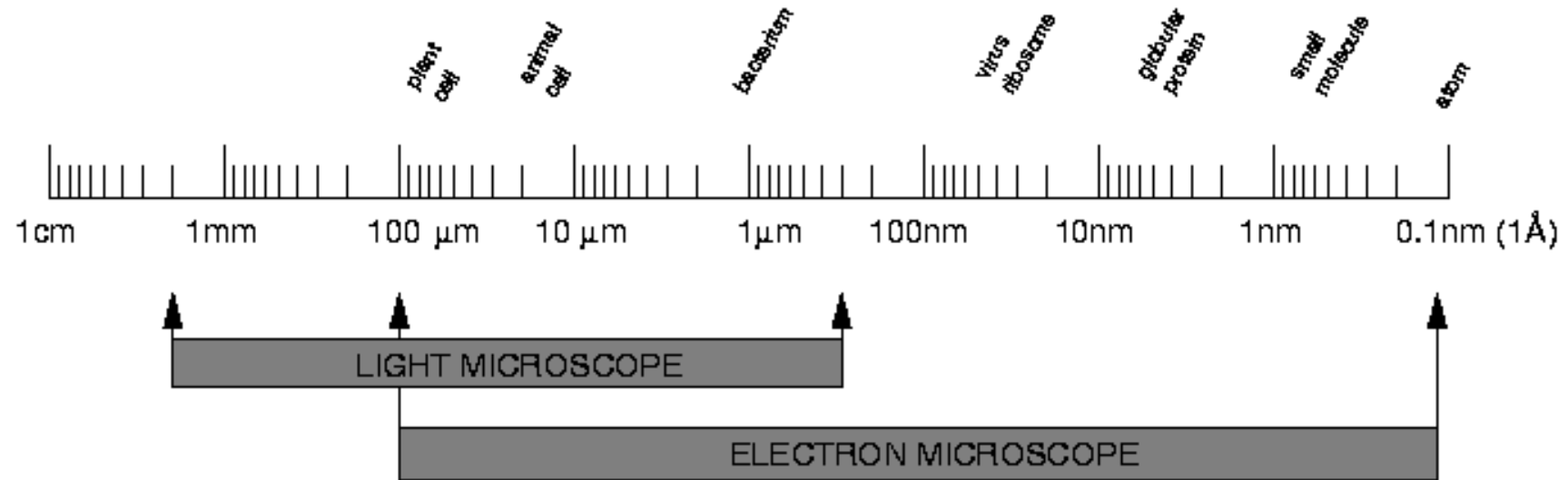


Figure 2.3 The definition of the semi-angle, α .

Why electrons?



Visible Light:

$$\lambda = 400 - 600 \text{ nm}$$

Electrons:

$$\lambda = 0.002 - 0.004 \text{ nm}$$

Sample Preparation - Staining

⇒ To increase contrast: heavy atoms interact with electrons stronger than biomolecules (C, N, O, S, P)

- **Positive Staining**

treat sample with solution of salt like uranyl acetate, lead citrate, osmium tetroxide – object is black on light background

- **Negative Staining**

place sample on dried film of heavy metal salt – object is light spot on black background

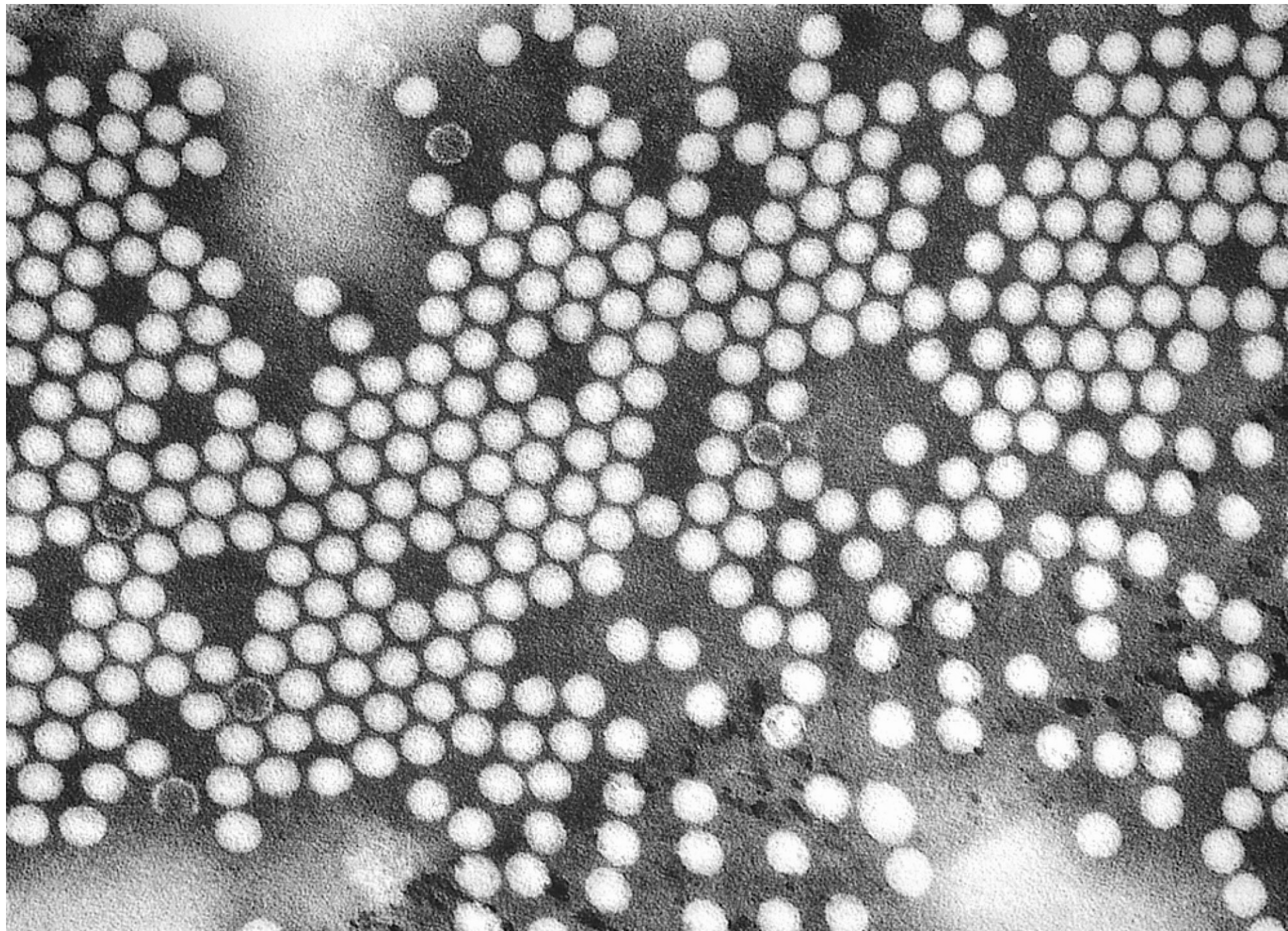
- **Shadowing**

spray thin layer of heavy metal on sample to produce a shadow

Disadvantage:

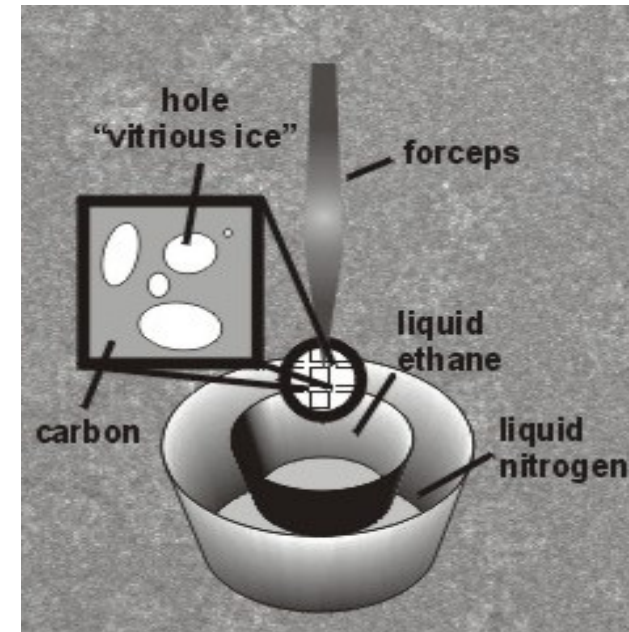
Size of stain reduces resolution to about 20-30 Å





Alternative: Cryo-EM

- to avoid harsh staining which may change the structure of your sample
- stabilization of sample by rapid freezing of sample in liquid ethane to form vitreous ice
- into electron microscope at low temperatures to keep sample stable in hydrated state in vacuum
- thickness of ice layer as small as possible!

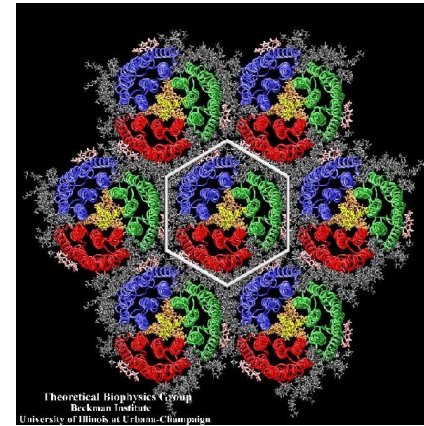


Advantage:

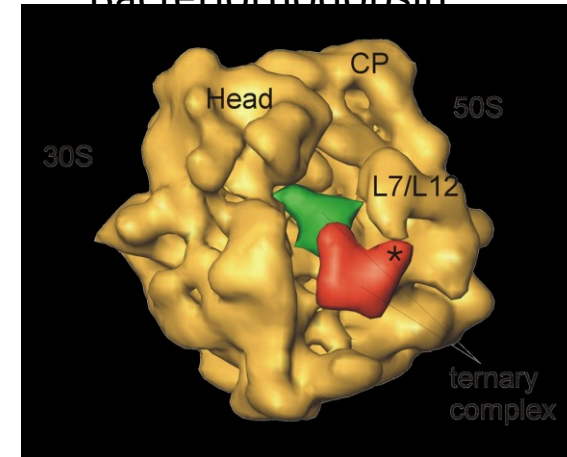
- sample structure is unchanged
- inner structures of molecules are accessible

Types of Samples

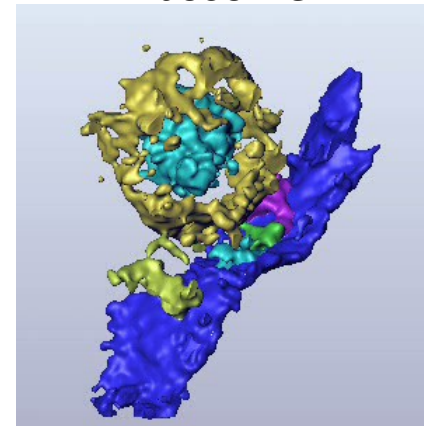
- Periodic arrangement
=> **2D electron crystallography**
small or membrane proteins < 200 kDa
resolution up to 2.5 Å
- Random arrangement
=> **single particle technique**
macromolecular complexes > 200 kDa
up to atomic resolution
- Large Organelles (Golgi, ER), whole cells
=> **tomography**
resolution > 4 Å

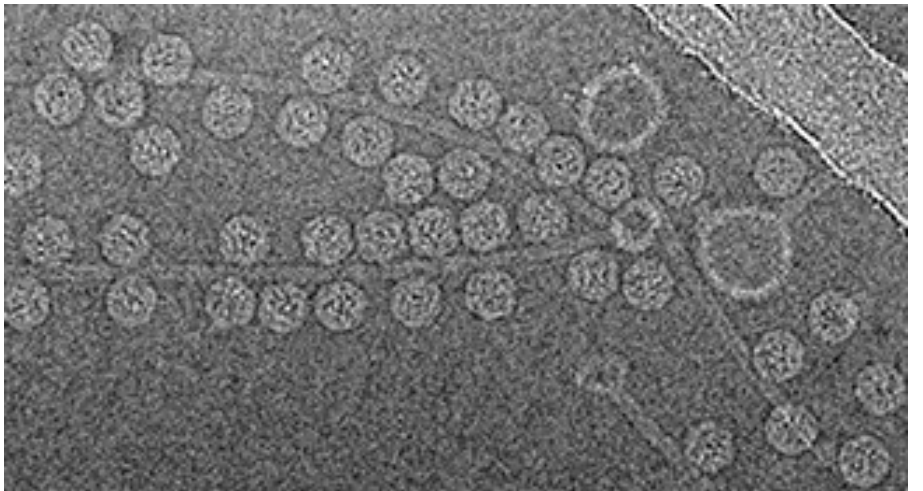
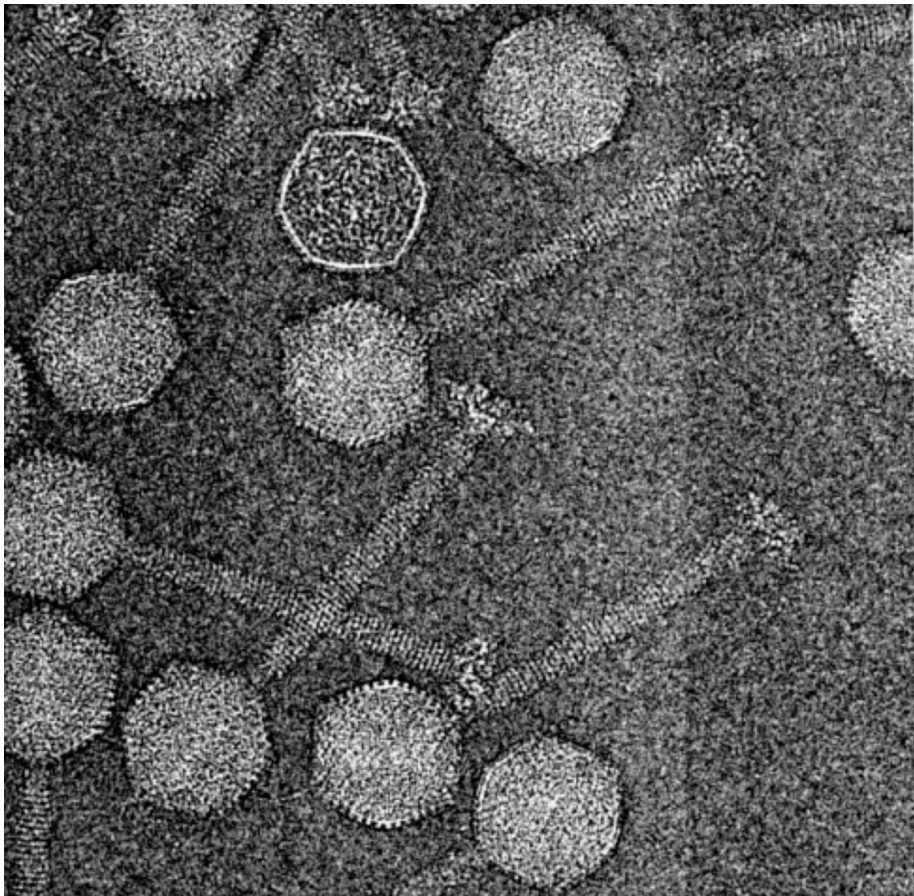
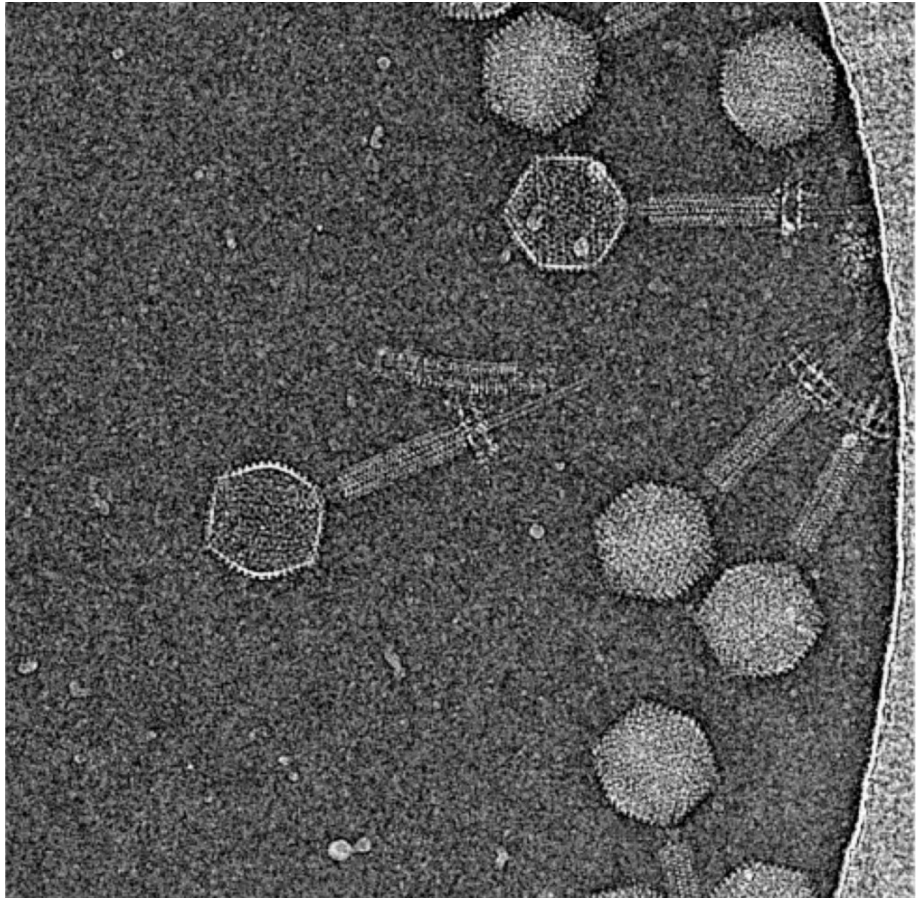
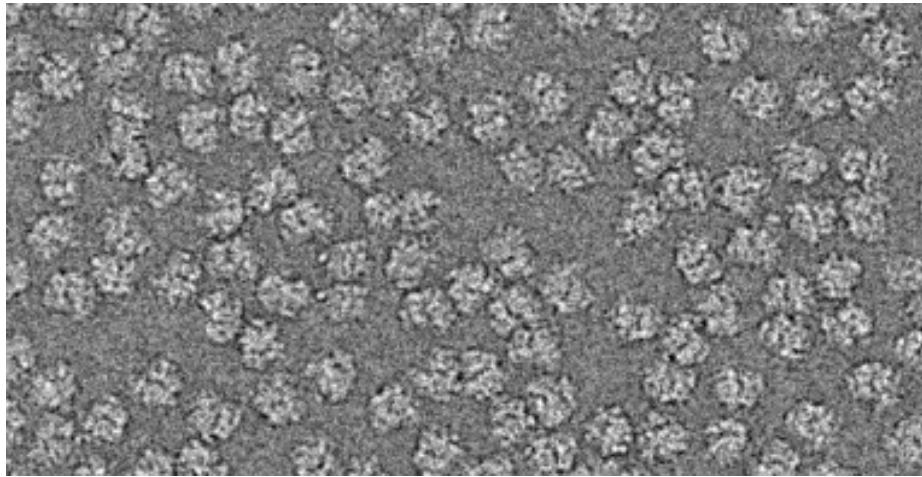


Bacteriorhodopsin

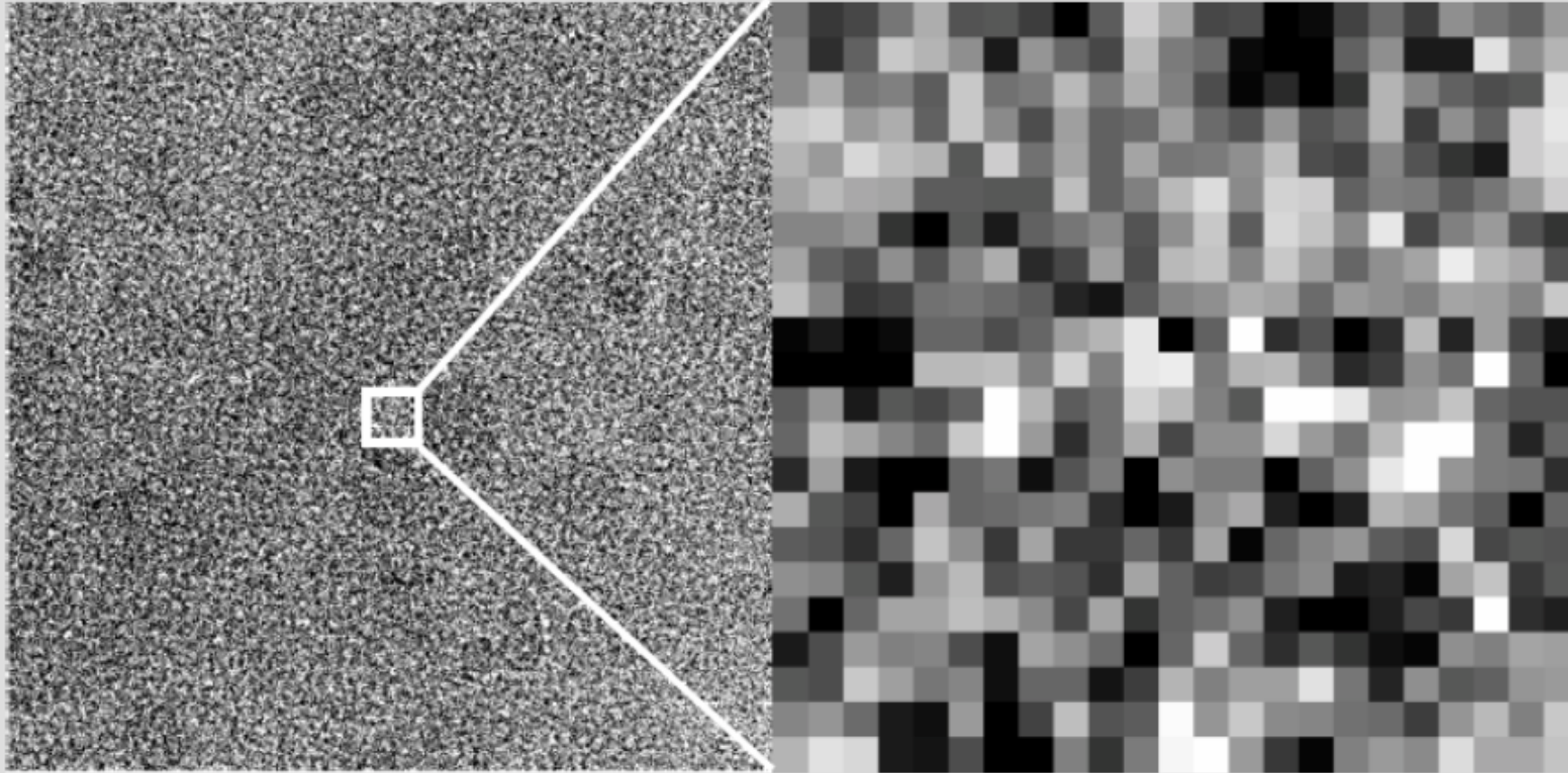


Ribosome



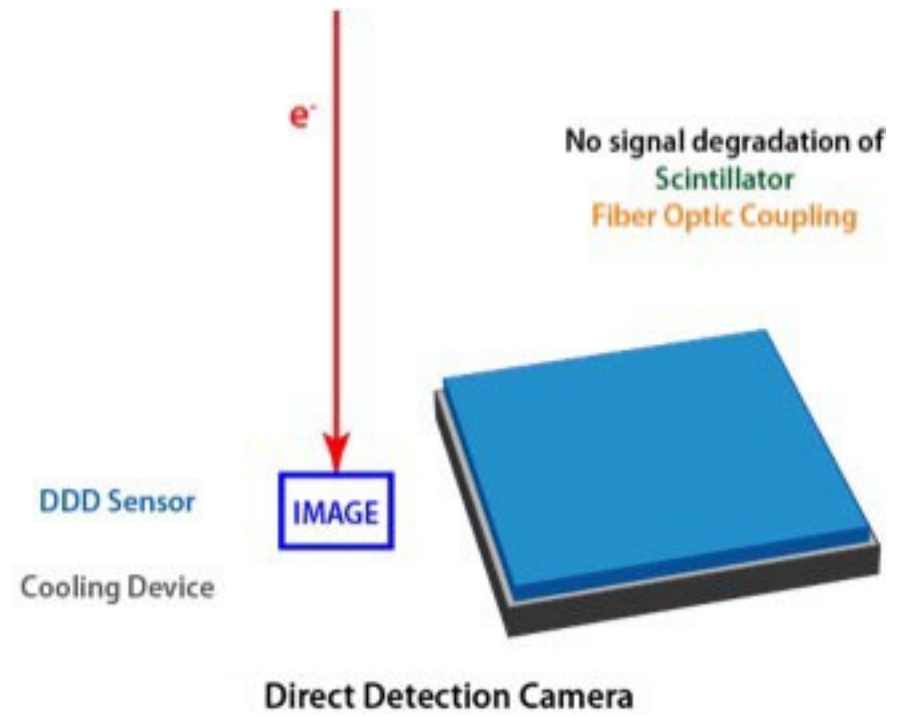
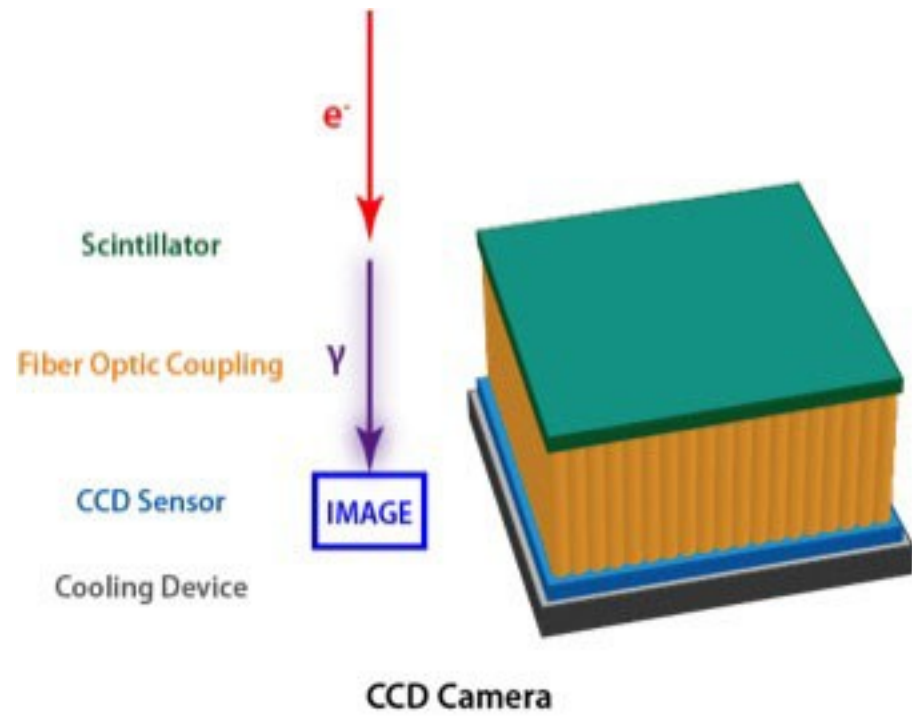


Digitization of Recorded Image



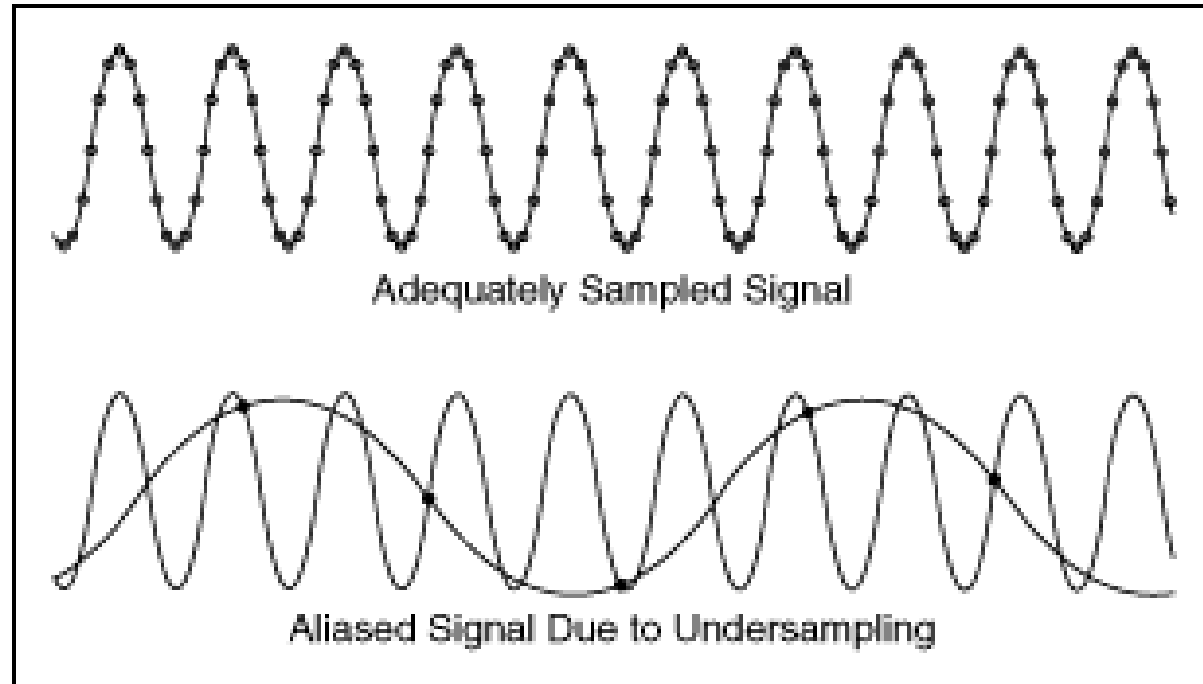
image

magnified sample of image showing
how the image is formed by a grid of
 $10 \times 10 \mu\text{m}$ pixel



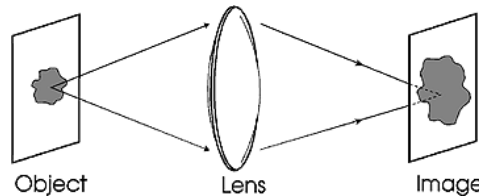
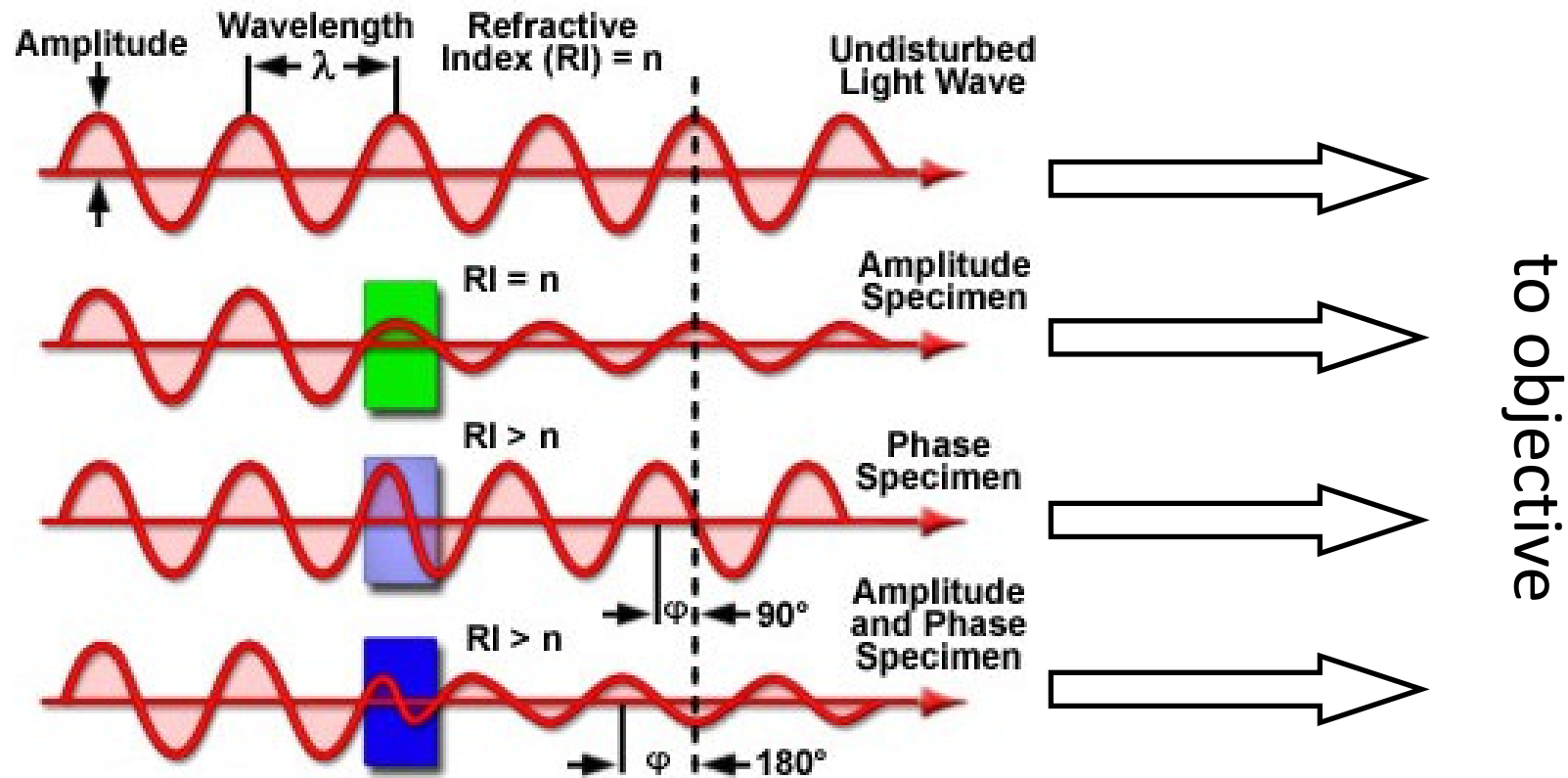
For a given sampling frequency, the maximum frequency you can accurately represent without aliasing is the Nyquist frequency, which equals one-half the sampling frequency, as shown by the following equation.

$$f_N = \frac{f_s}{2}$$

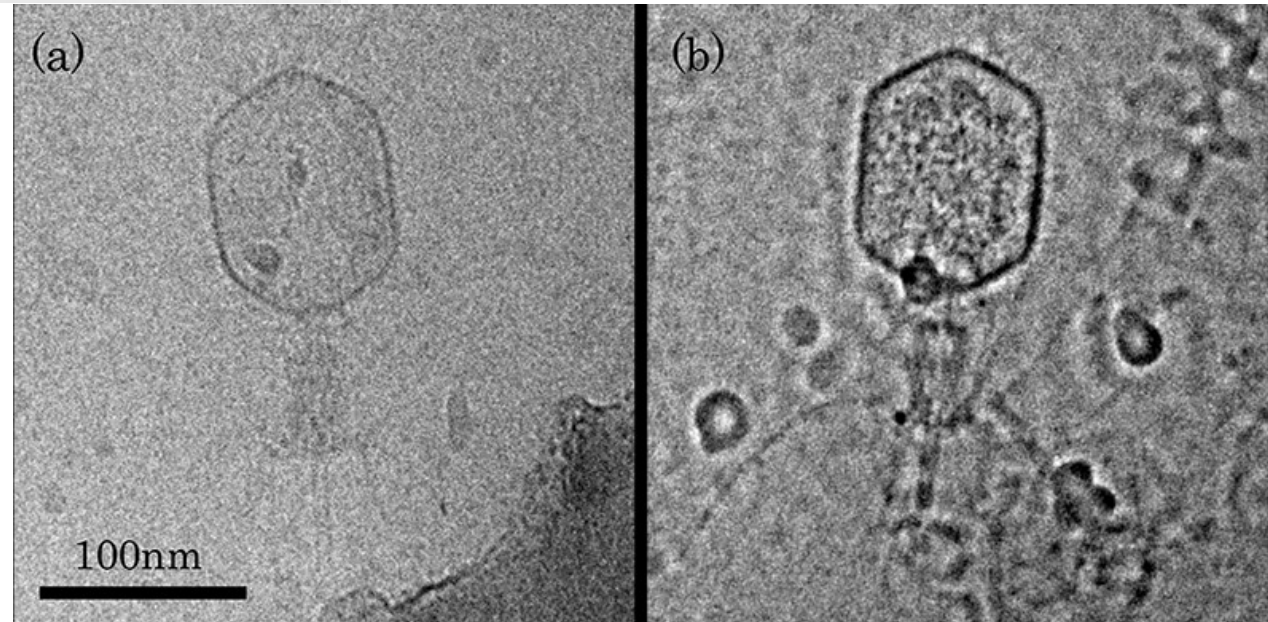
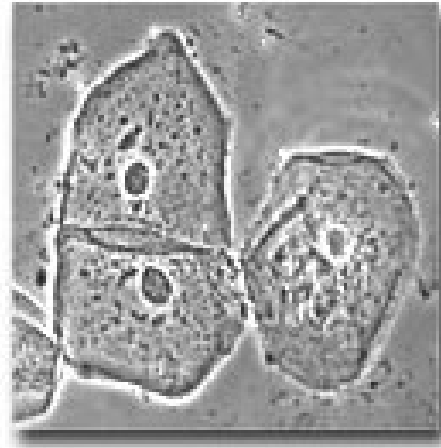
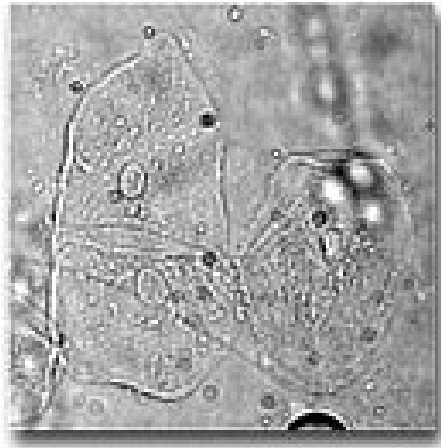




Macromolecules in water / vitreous ice are phase objects



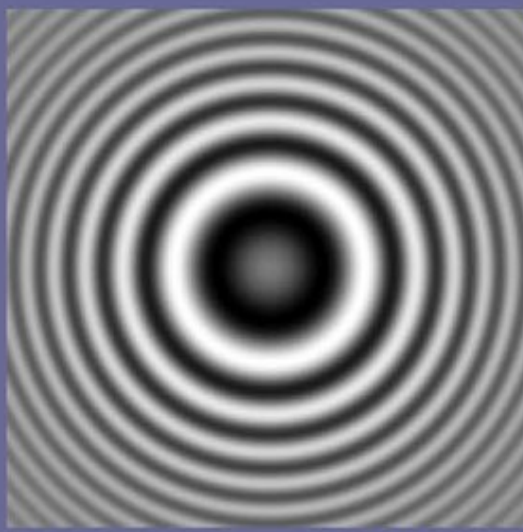
Phase Objects Require an Additional Phase Shift to be Seen





original object

\times

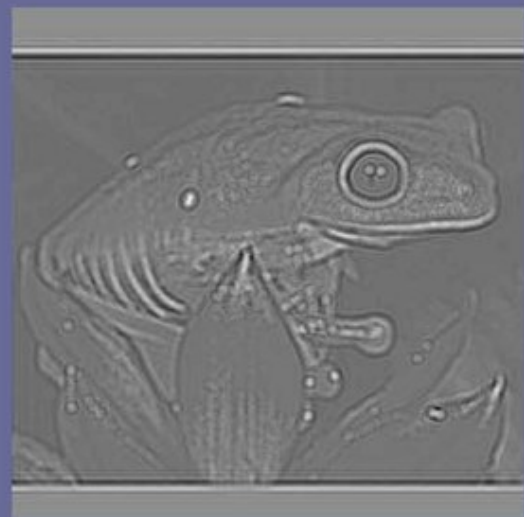
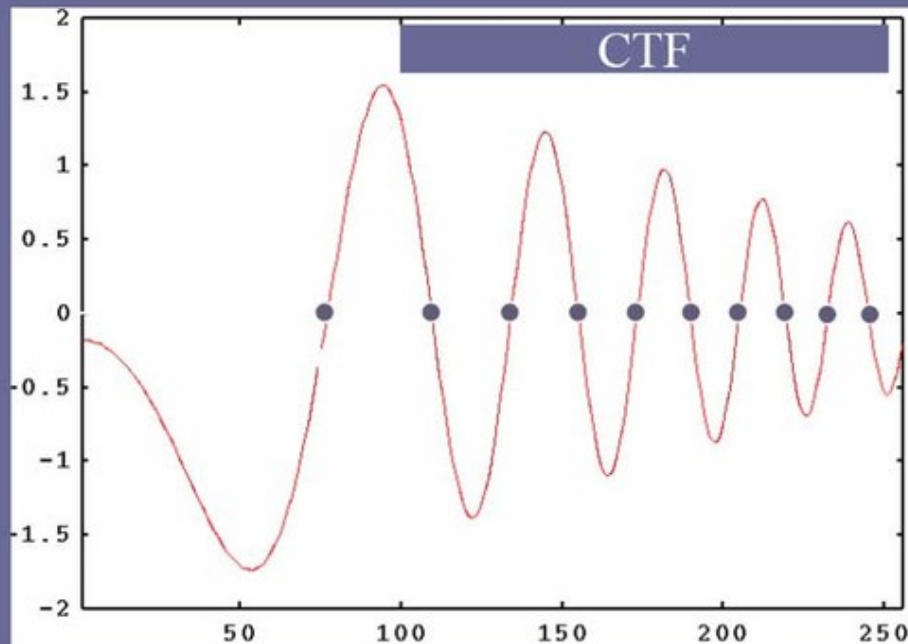


CTF for $\Delta z = 2.500 \mu\text{m}$

$=$



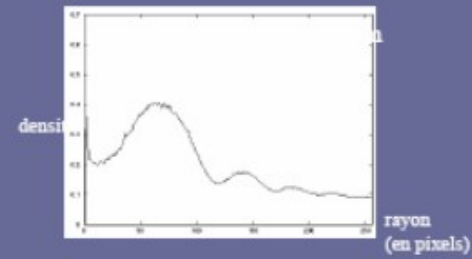
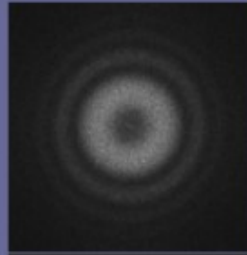
cryo-EM image



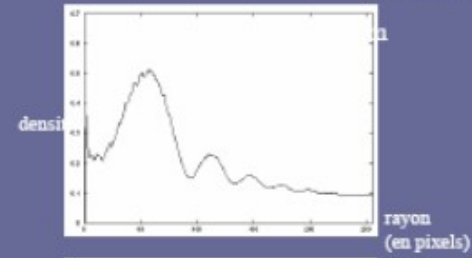
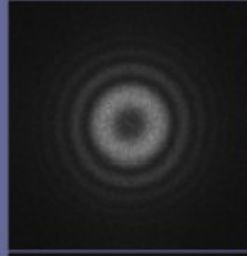
cryo-EM image,
contrast-inverted

Gallery of Power Spectra at Different Defocus

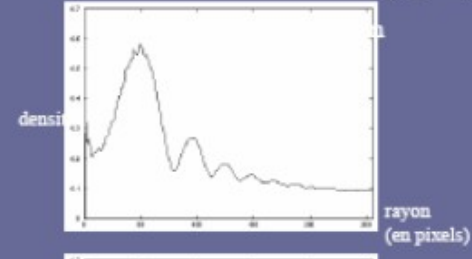
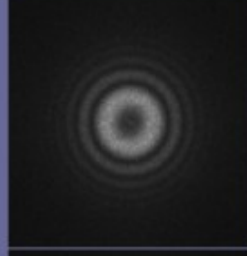
A



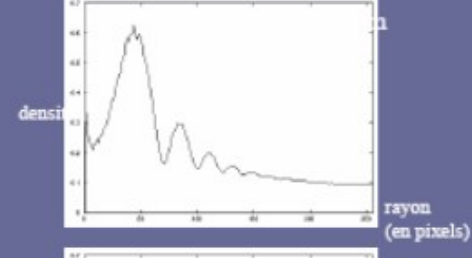
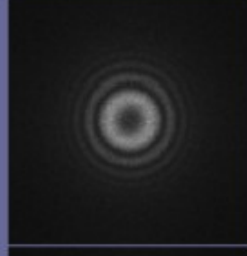
B



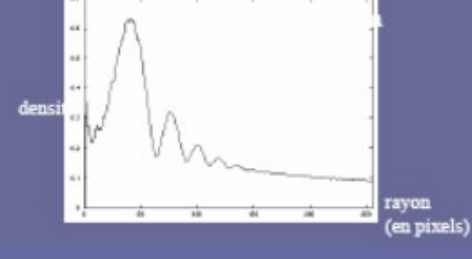
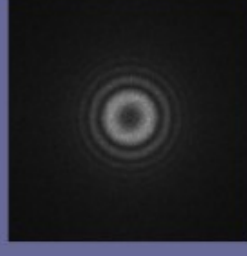
C



D



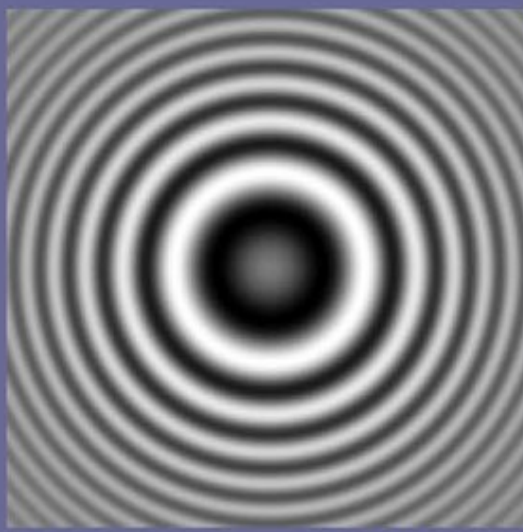
E





original object

\times

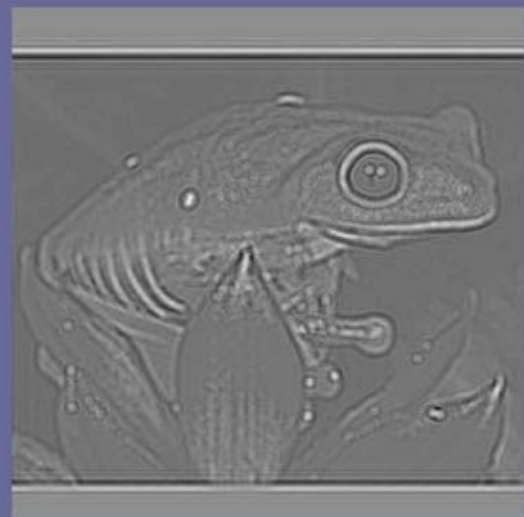
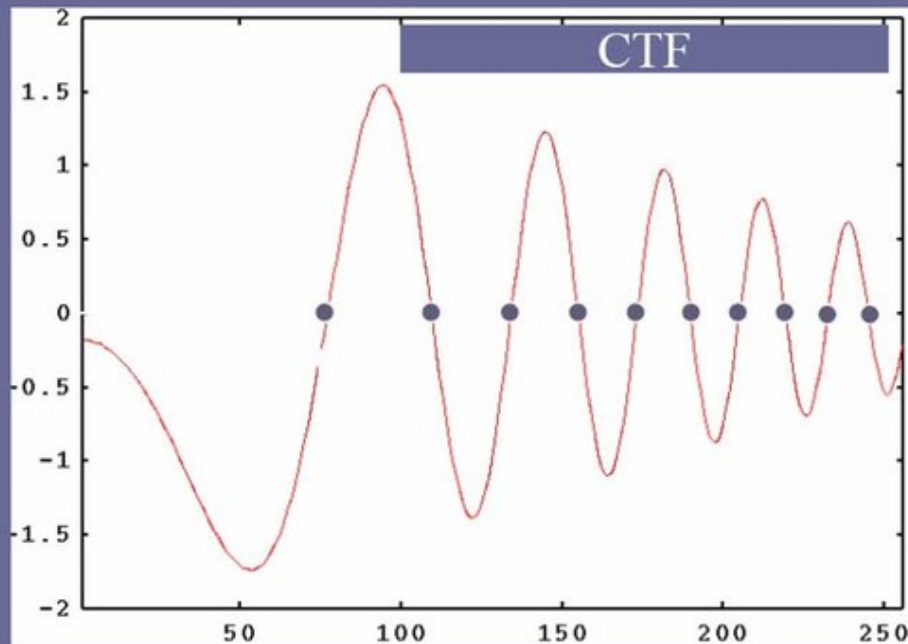


CTF for $\Delta z = 2.500 \mu\text{m}$

$=$

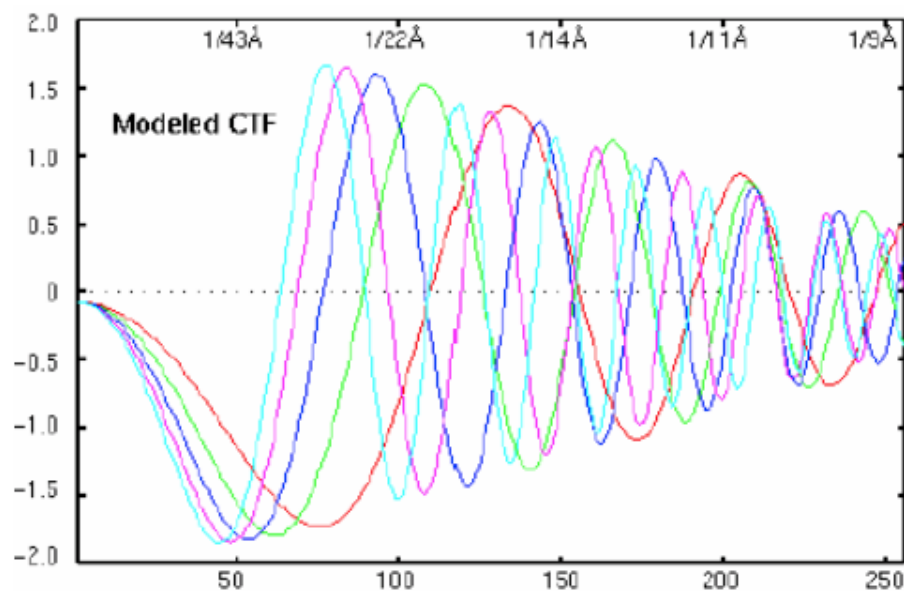
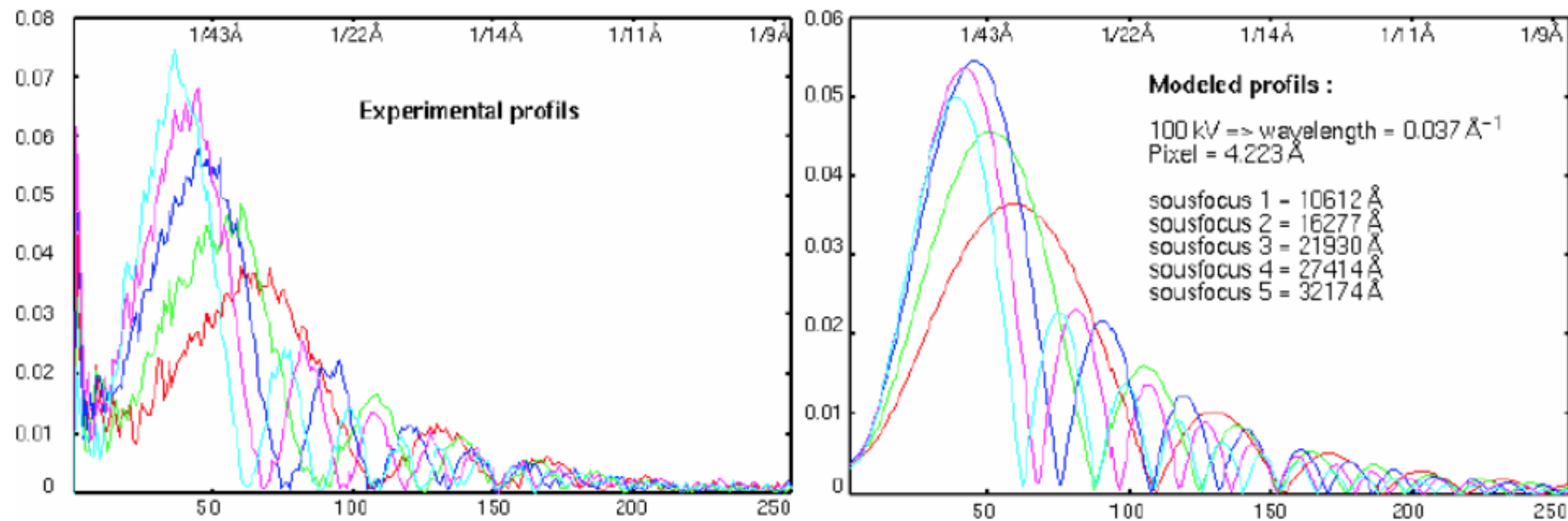


cryo-EM image

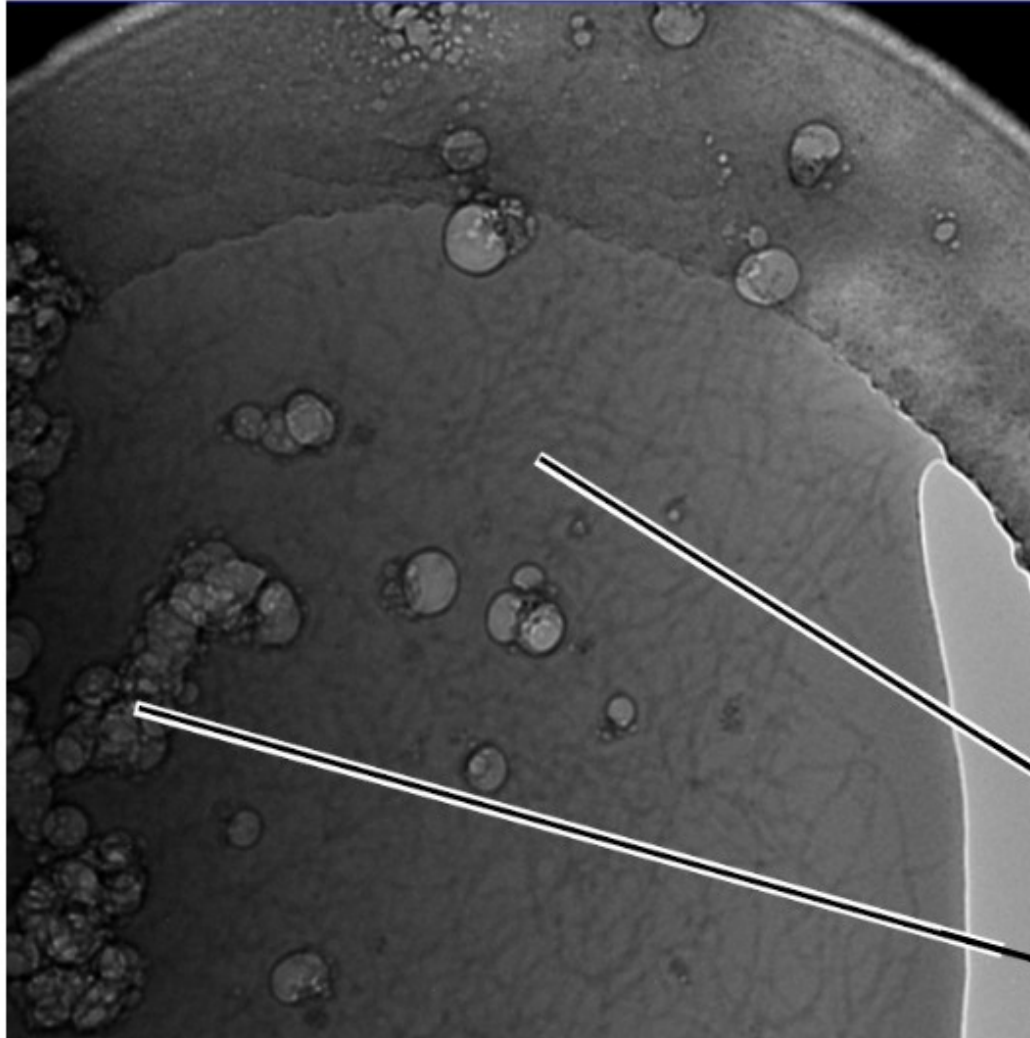


cryo-EM image,
contrast-inverted

Multiple Defocus Groups



Bubbling: A Sign of Radiation Damage



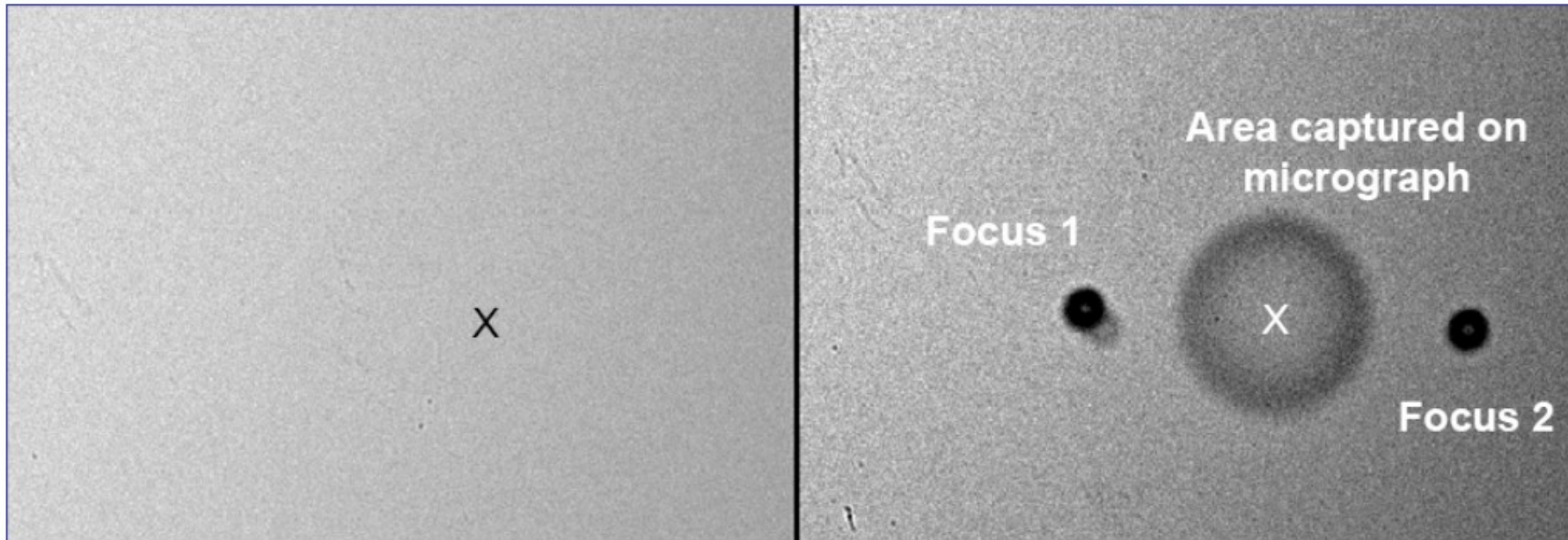
A sample of unstained amyloid materials after a few seconds of illumination with an electron beam.

While some fibers can still be detected, "bubbling" within the field of view indicates total destruction of the sample

amyloid fibers

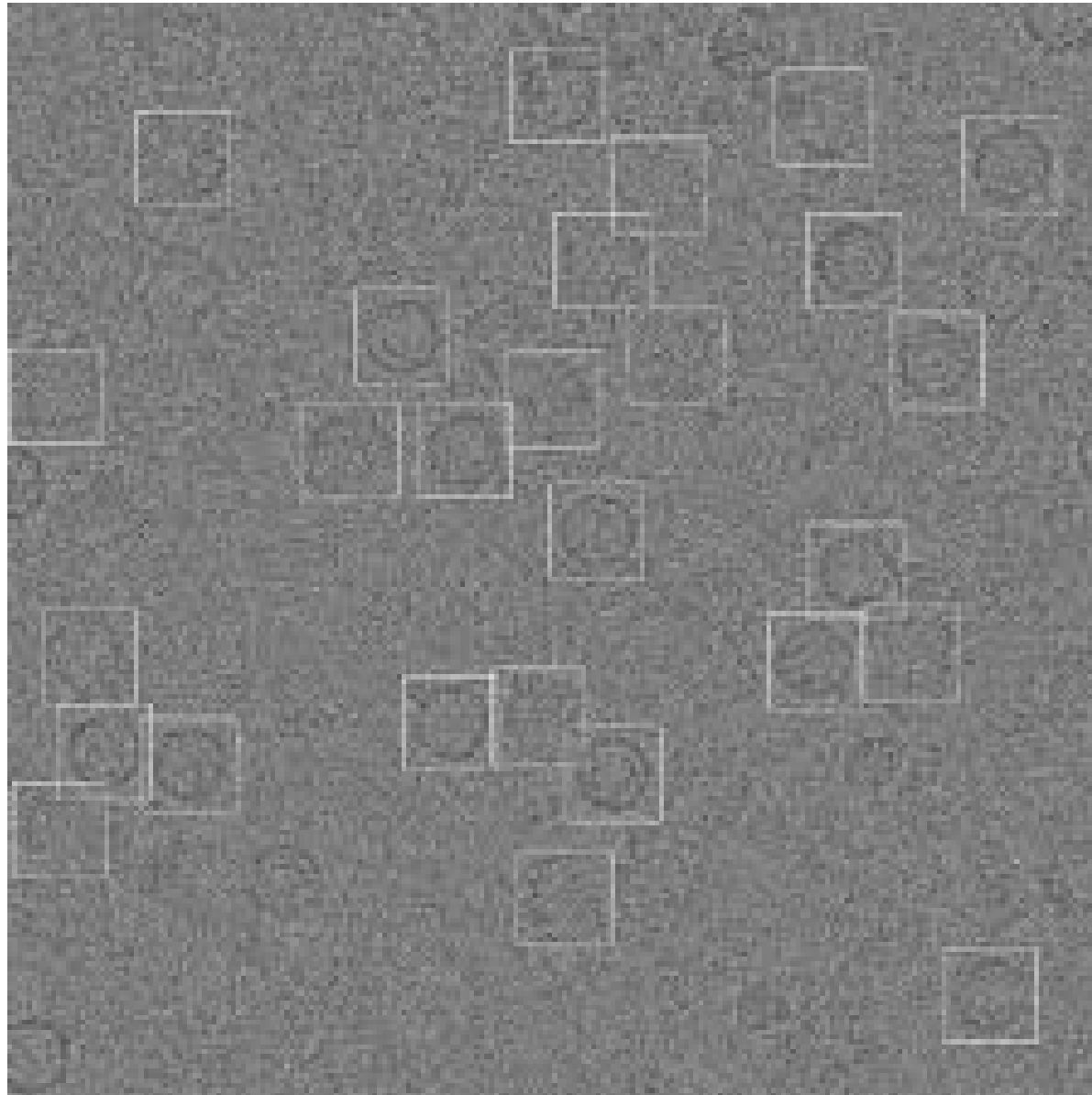
bubbles

Low-Dose Microscopy

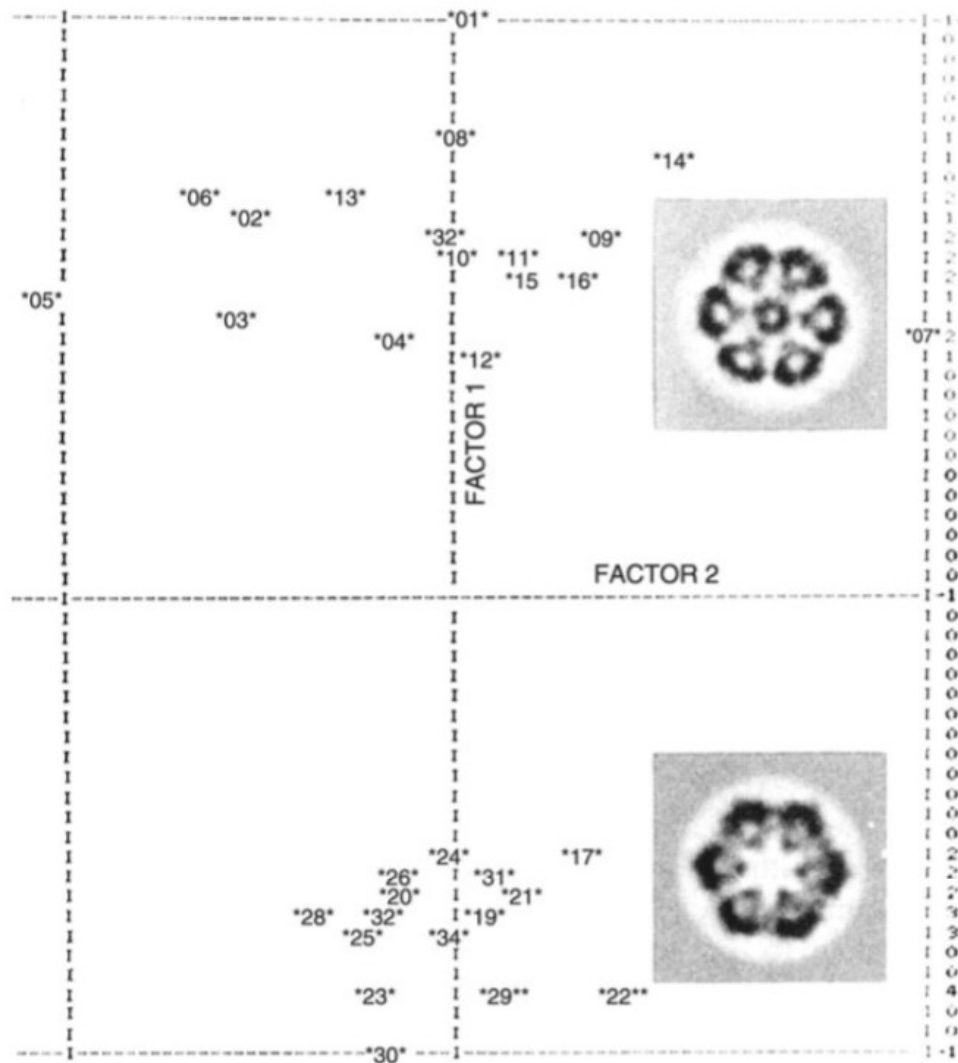
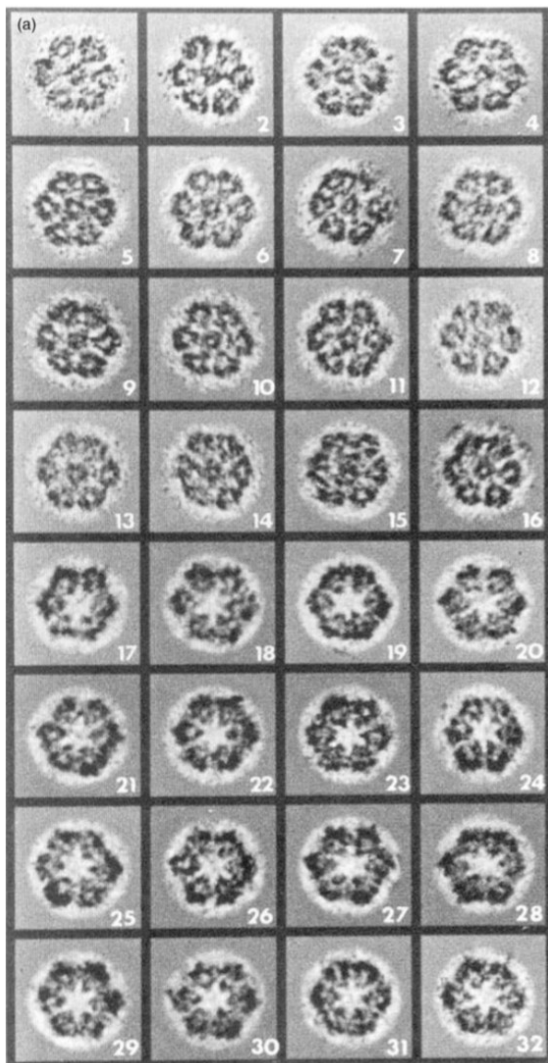


Appearance of trehalose dried down on a carbon film (left). The sugar allows to demonstrate how “low-dose” microscopy is done (right). Let X be the area of interest (for instance a crystal or virus/single particle). Prior to taking a picture some parameters such as “defocus” and “astigmatism” need to be adjusted. To avoid destruction of the specimen, any adjustments are made on small areas (Focus 1 and 2) located adjacent to the area that will be photographed. In the example, the trehalose burned as it was exposed at high magnification (220kx, Focus 1 and 2). Similarly, by exposing the area to be captured for about 30 seconds at 52,000 fold magnification.

Signal to noise ratio

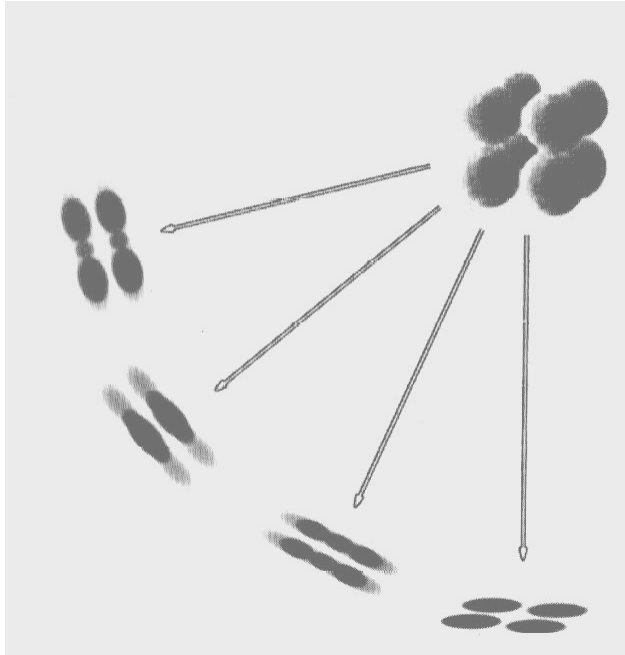


Classification and averaging (principal component analysis)



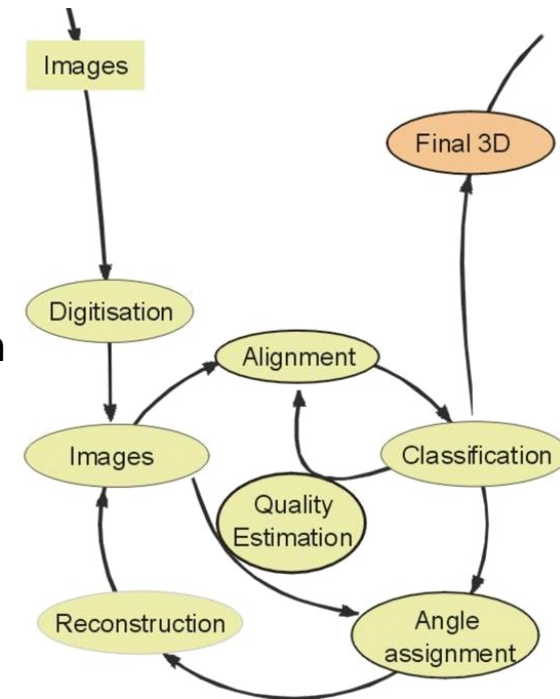
3D Reconstruction

When the angles between the different classes are known (estimated), a 3D model can be calculated.



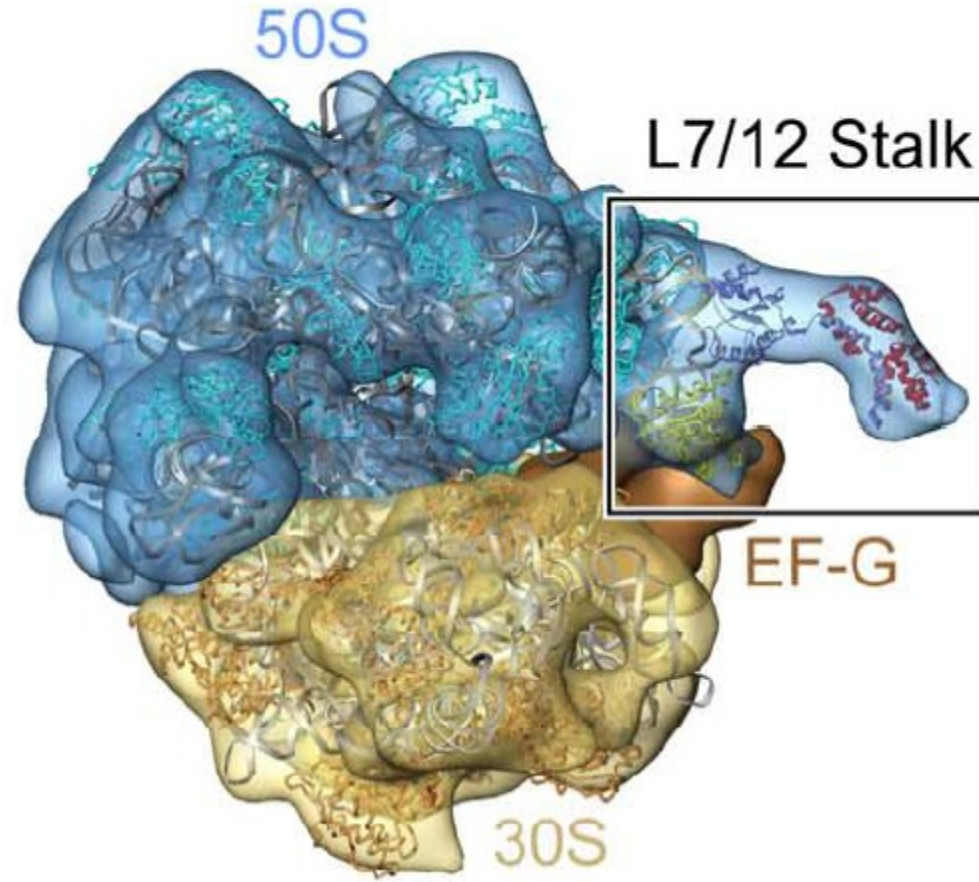
Iterative Process:

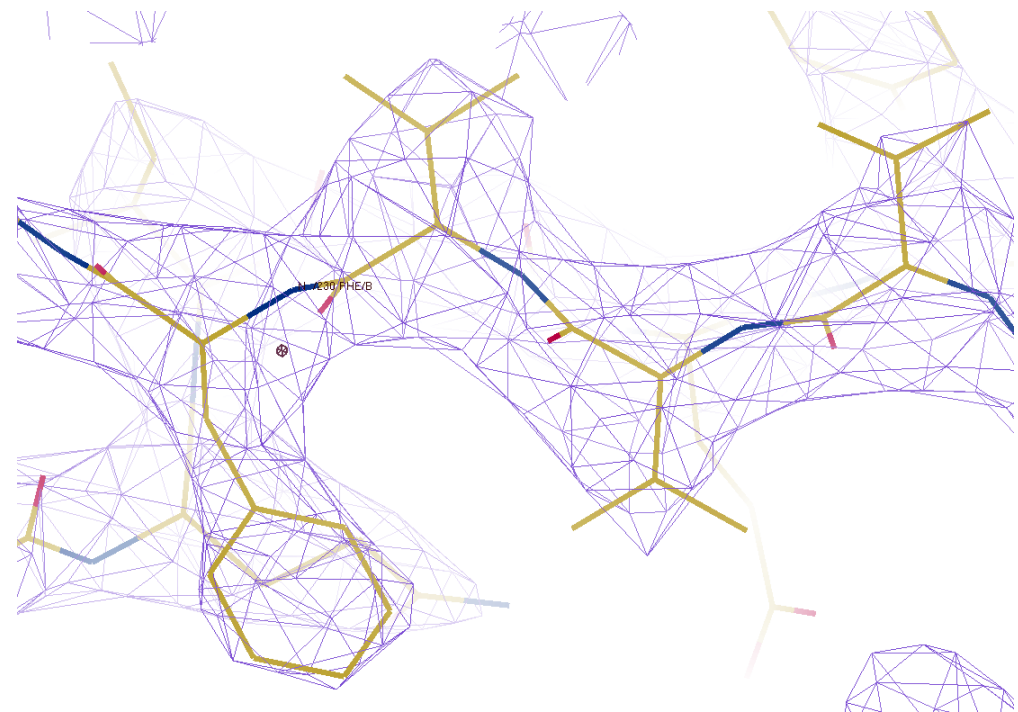
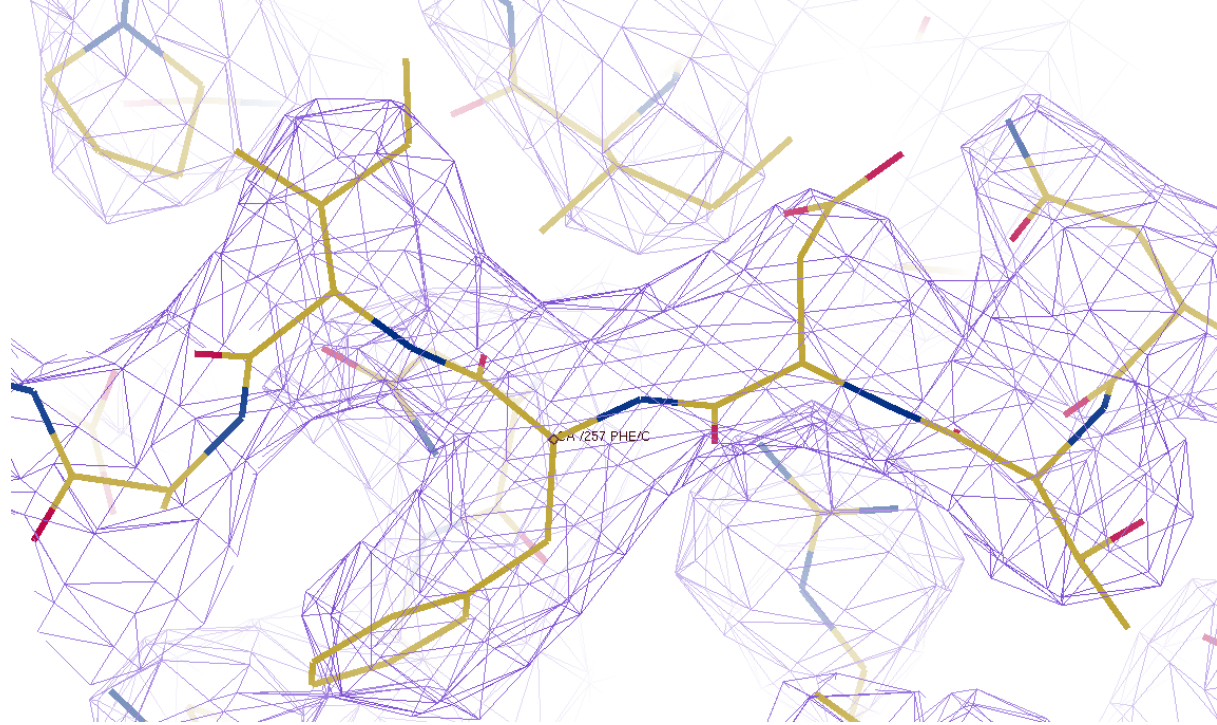
3D model is used to generate 2D images which are fed into statistical analysis of images (alignment and classification).

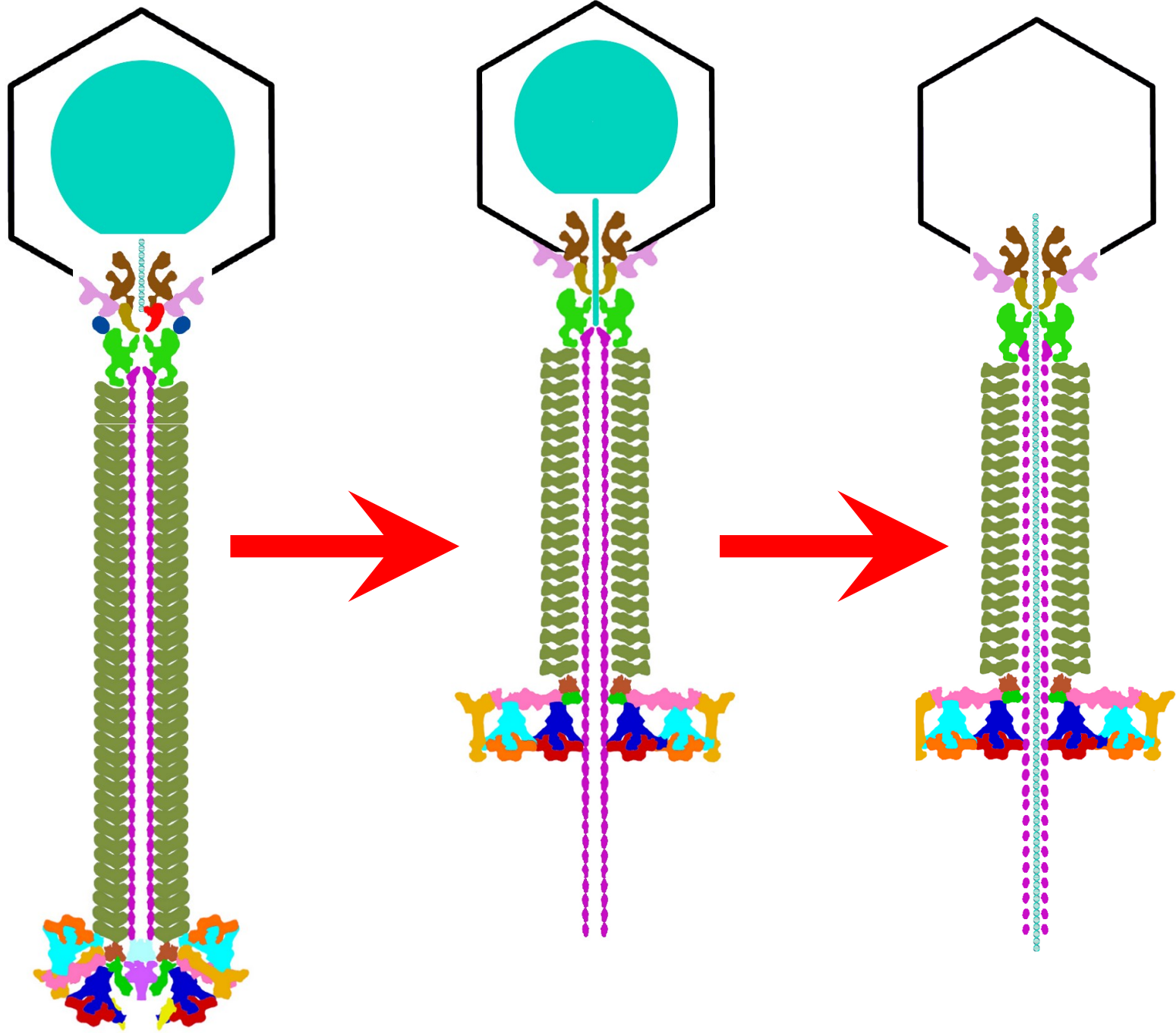


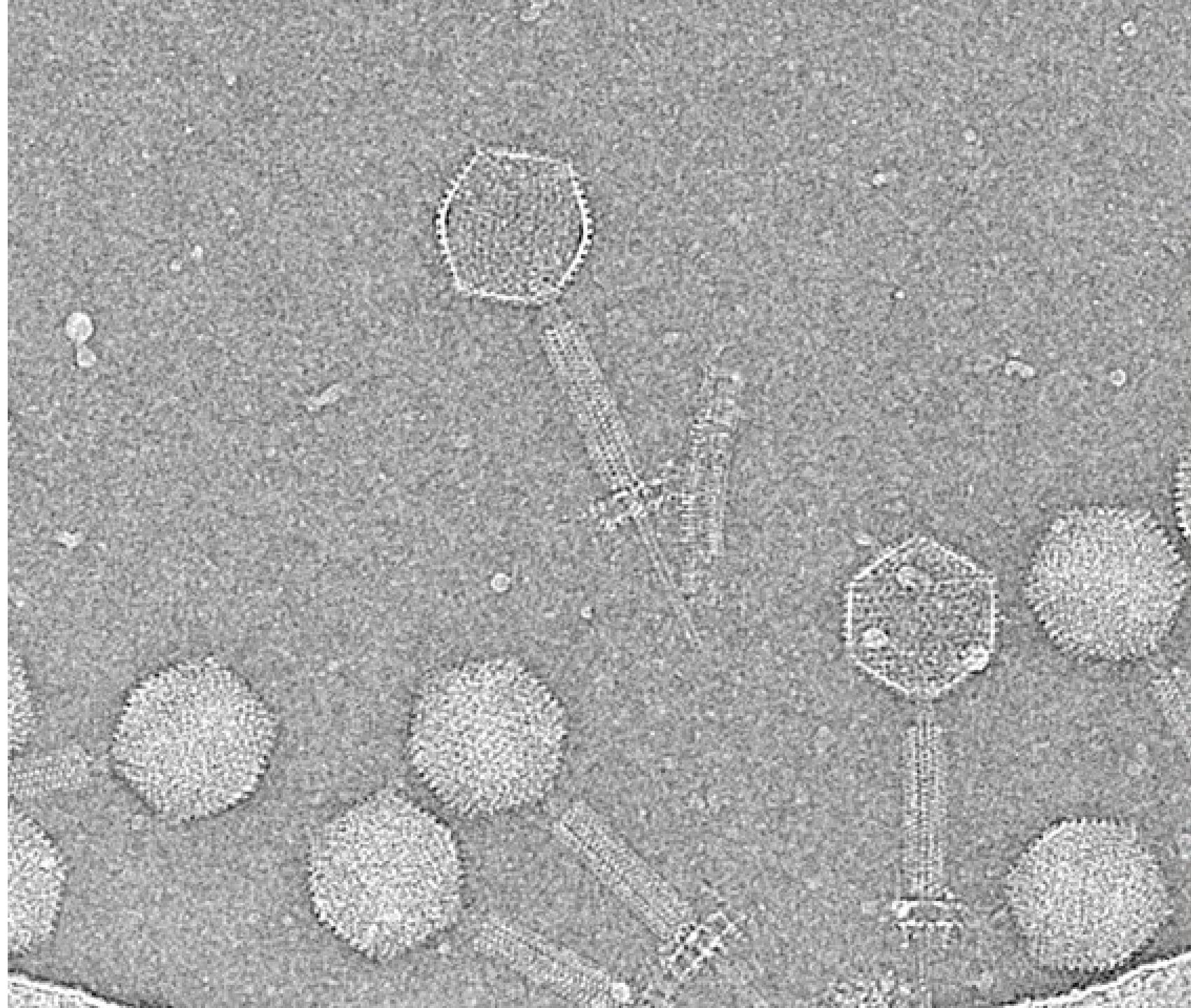
And then?

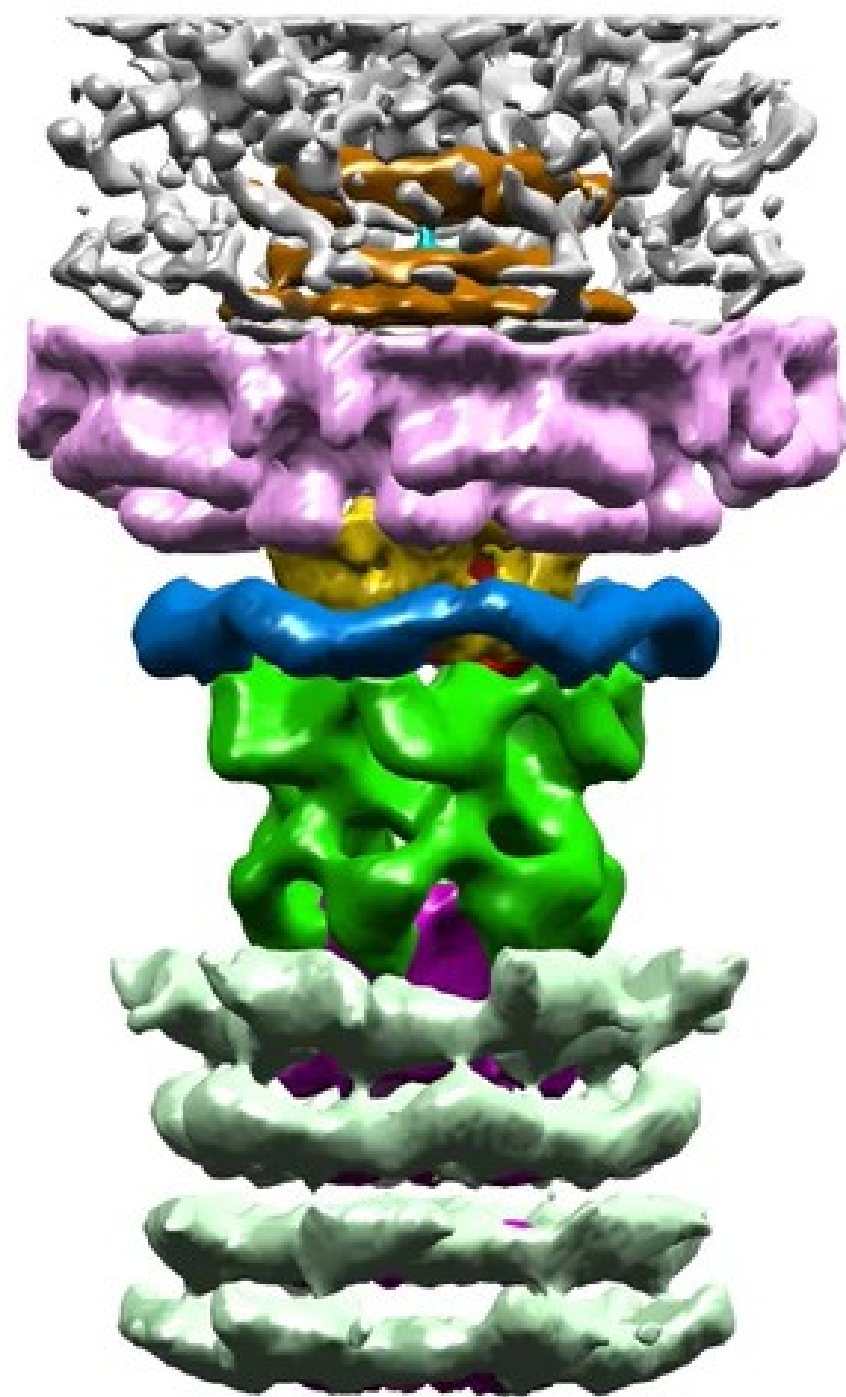
Try to interpret 3D map,
e.g. try to fit known crystal
structures into electron
density map

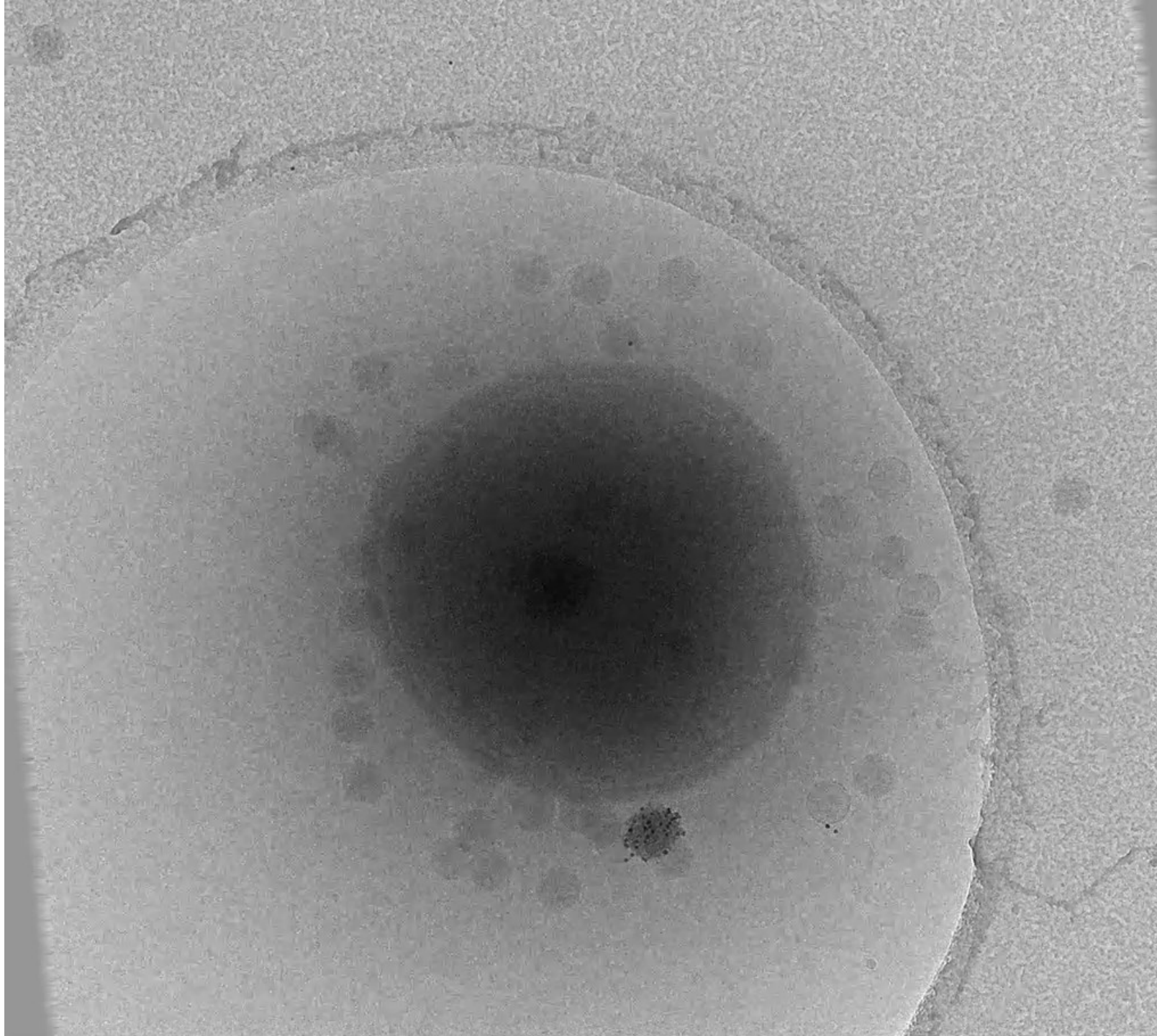






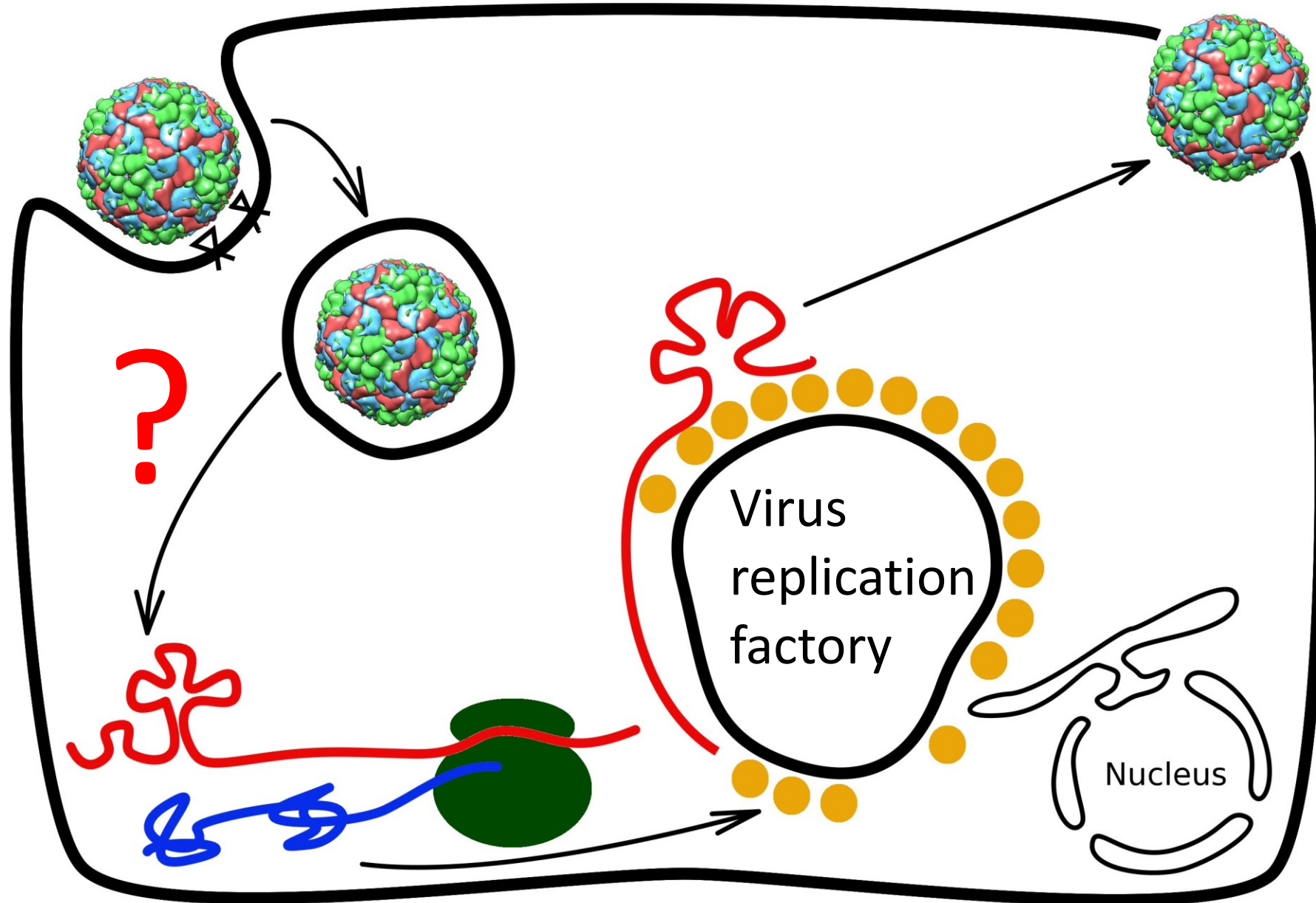




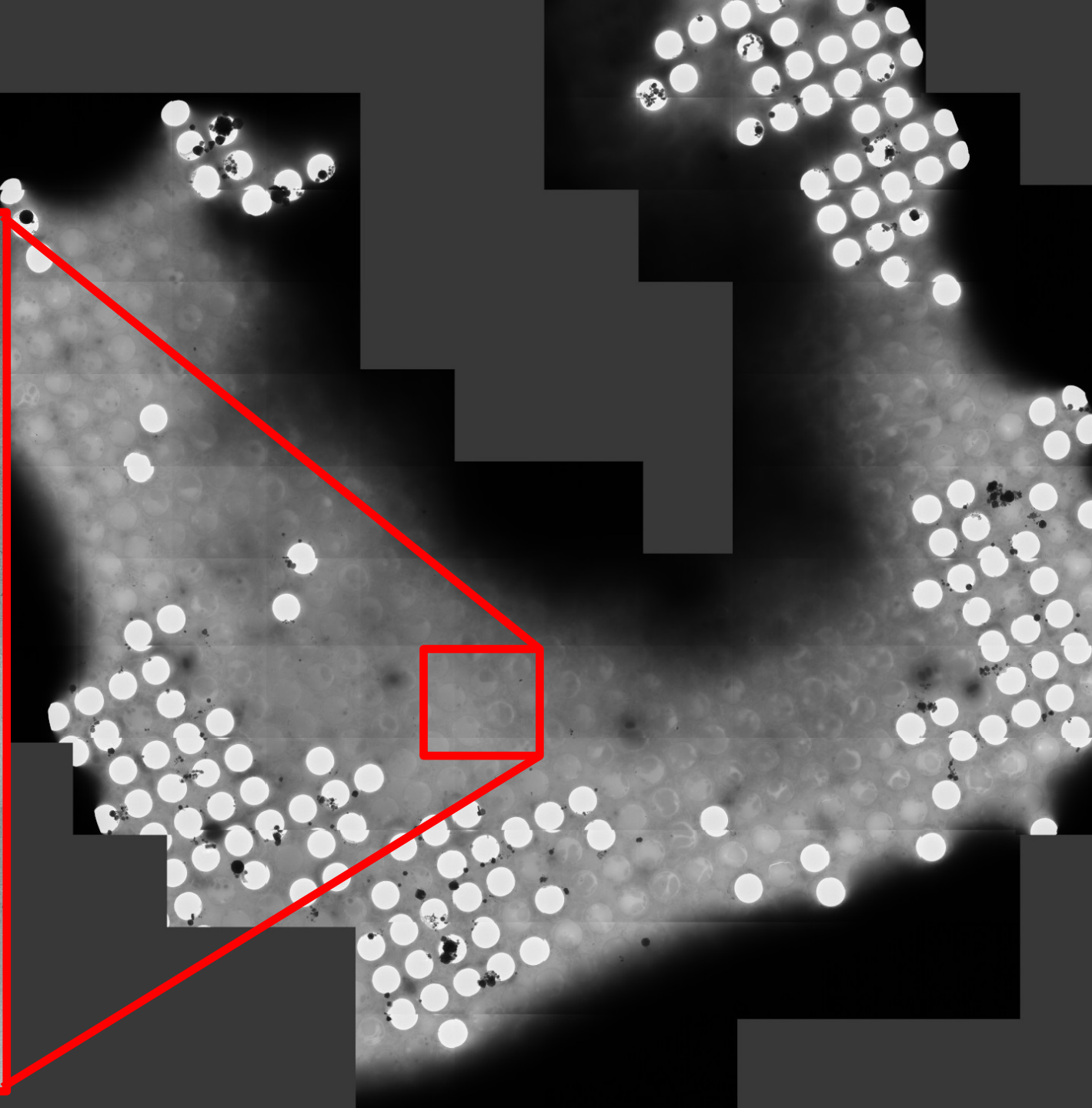
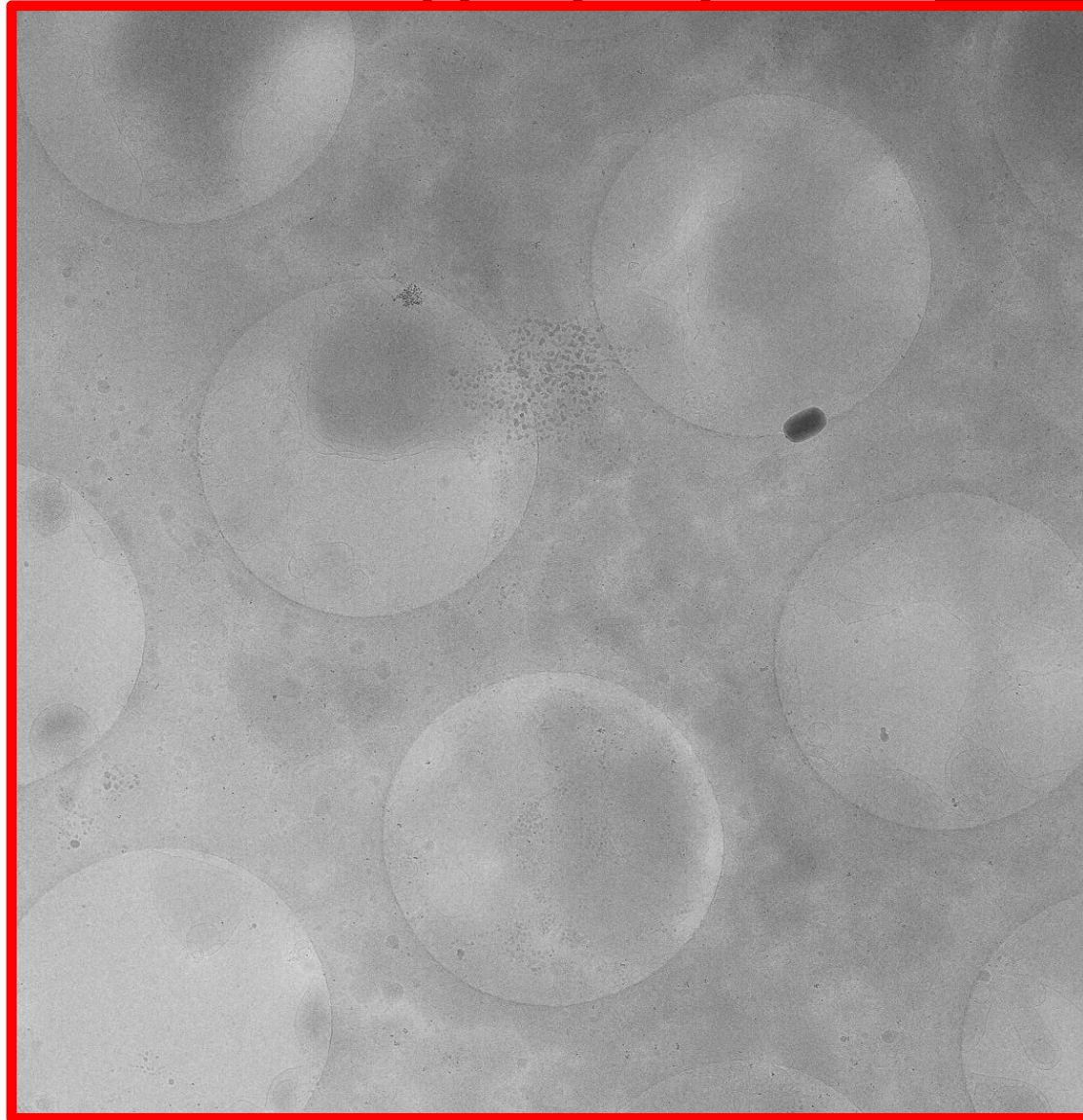




Infection cycle

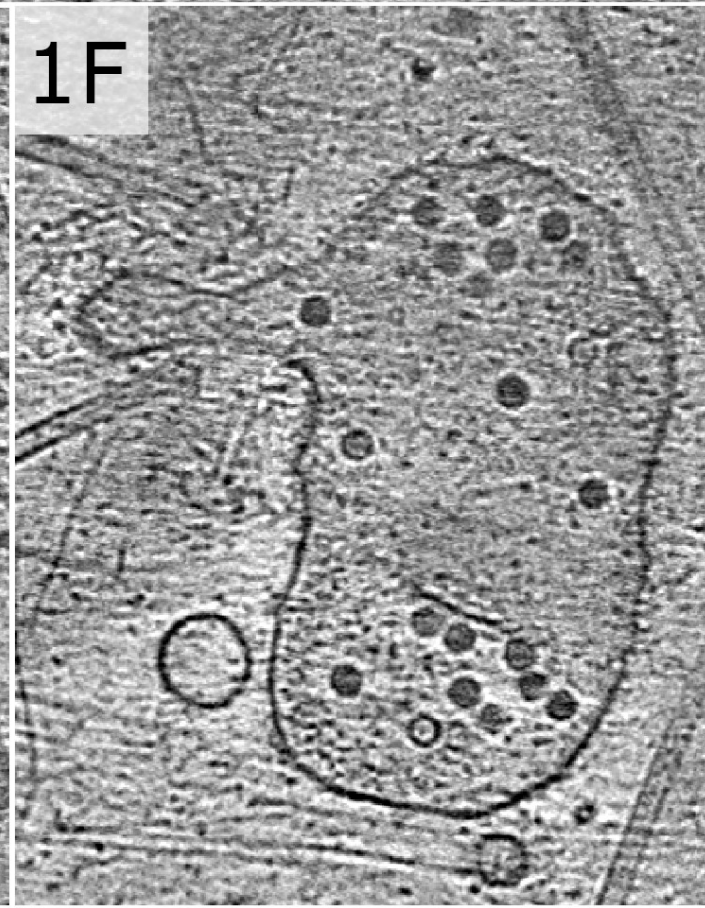
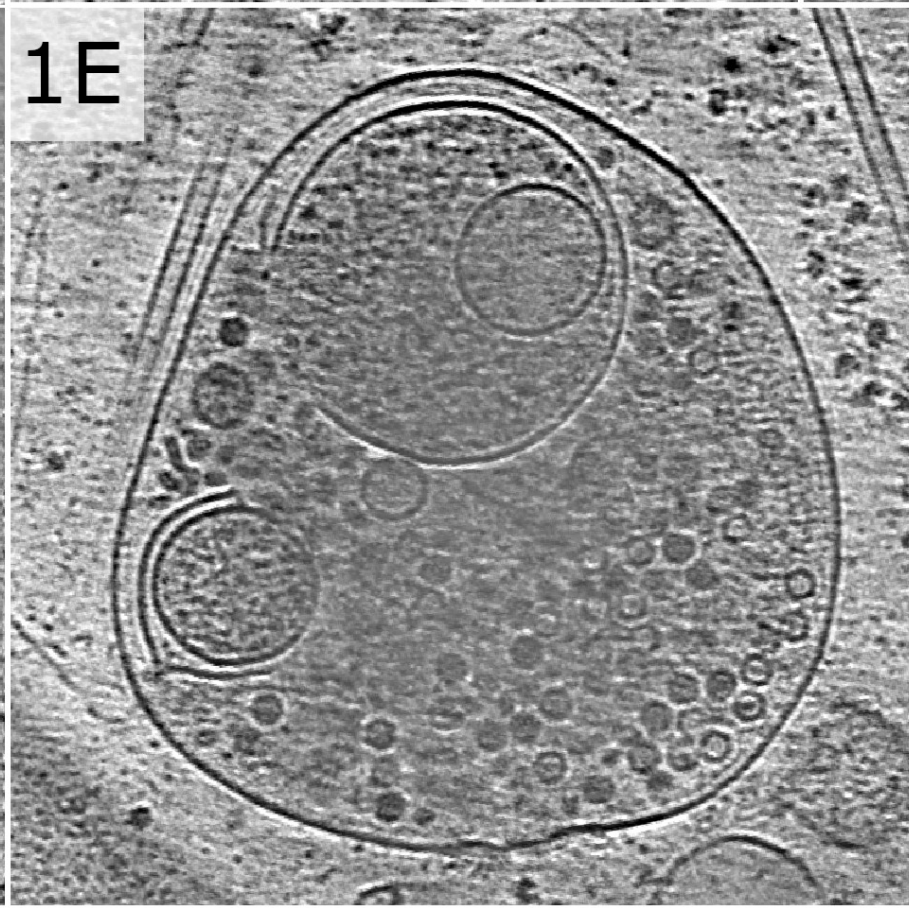
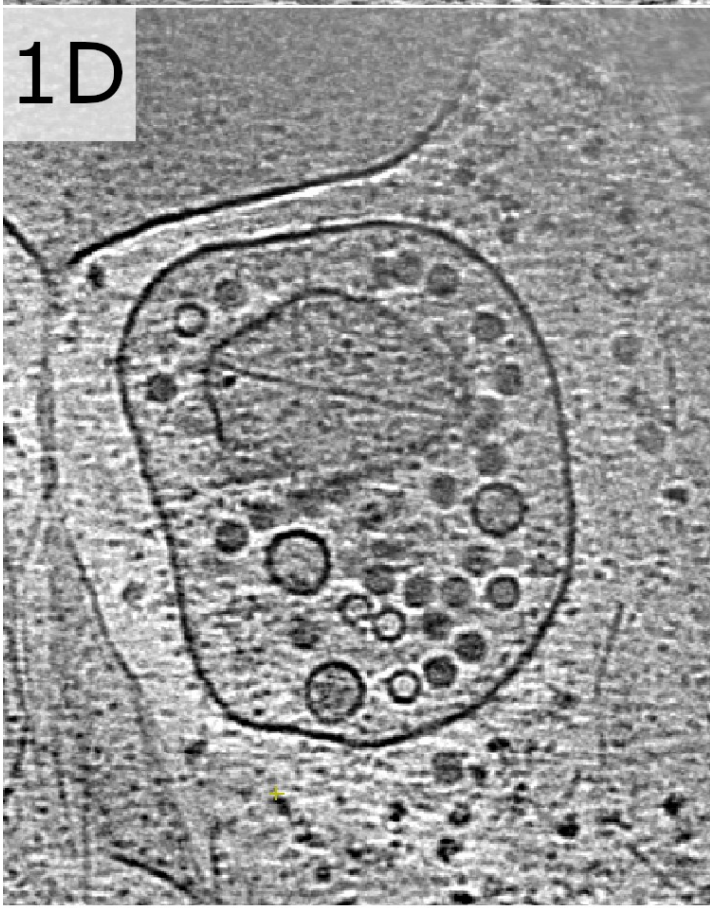
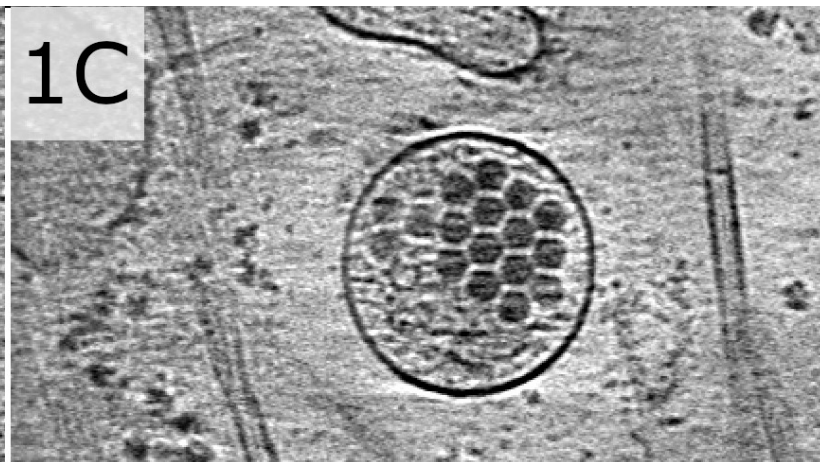
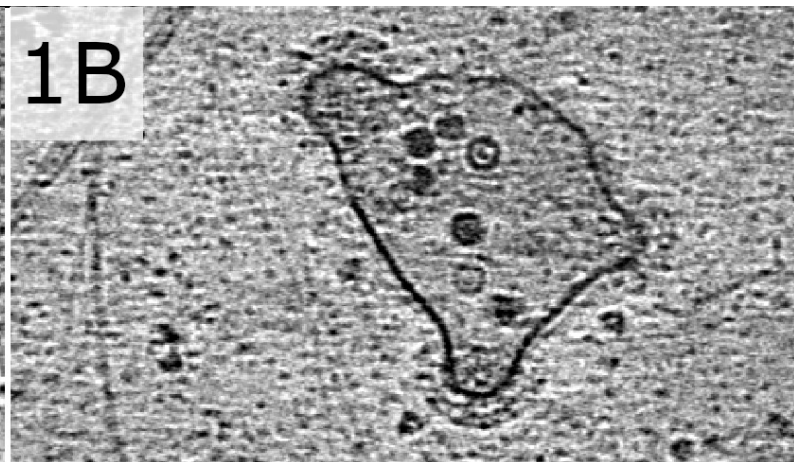
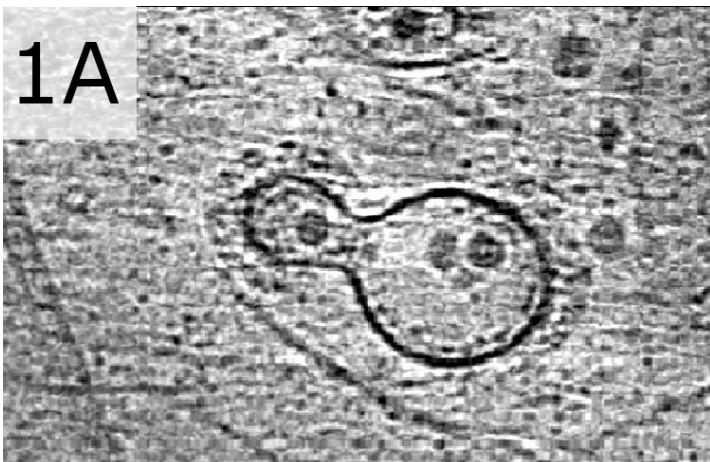


in situ cryo-electron
tomography

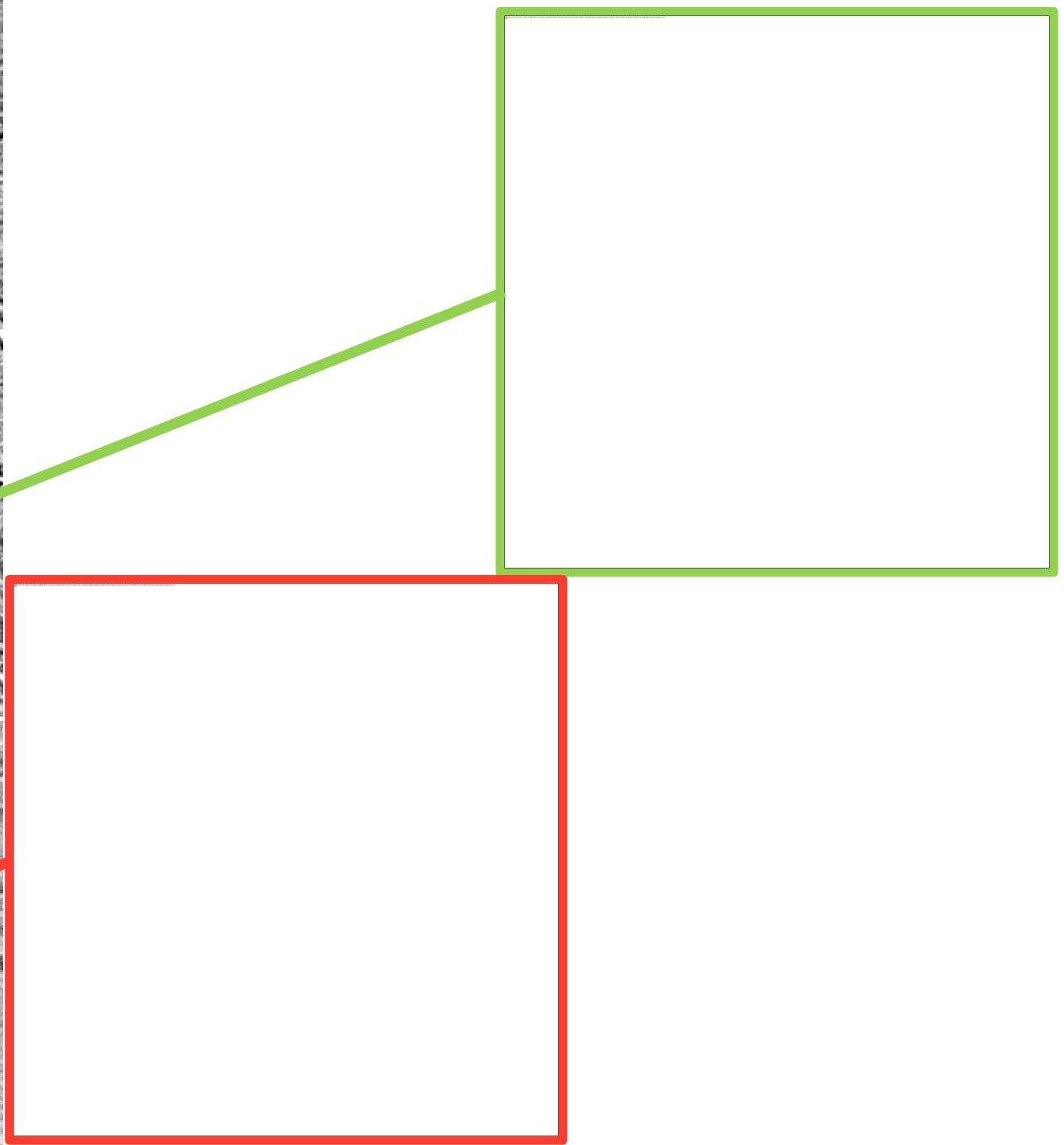
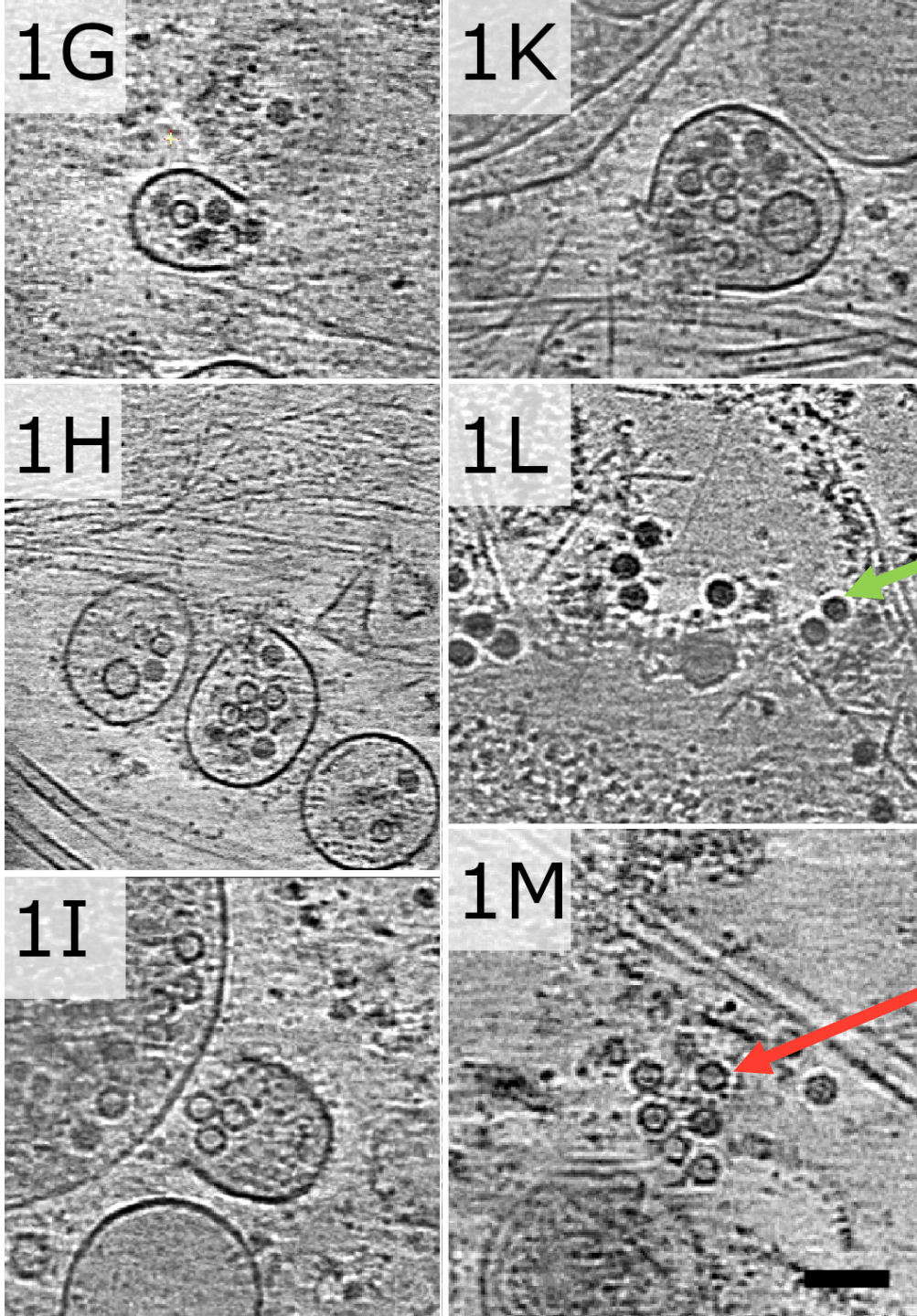




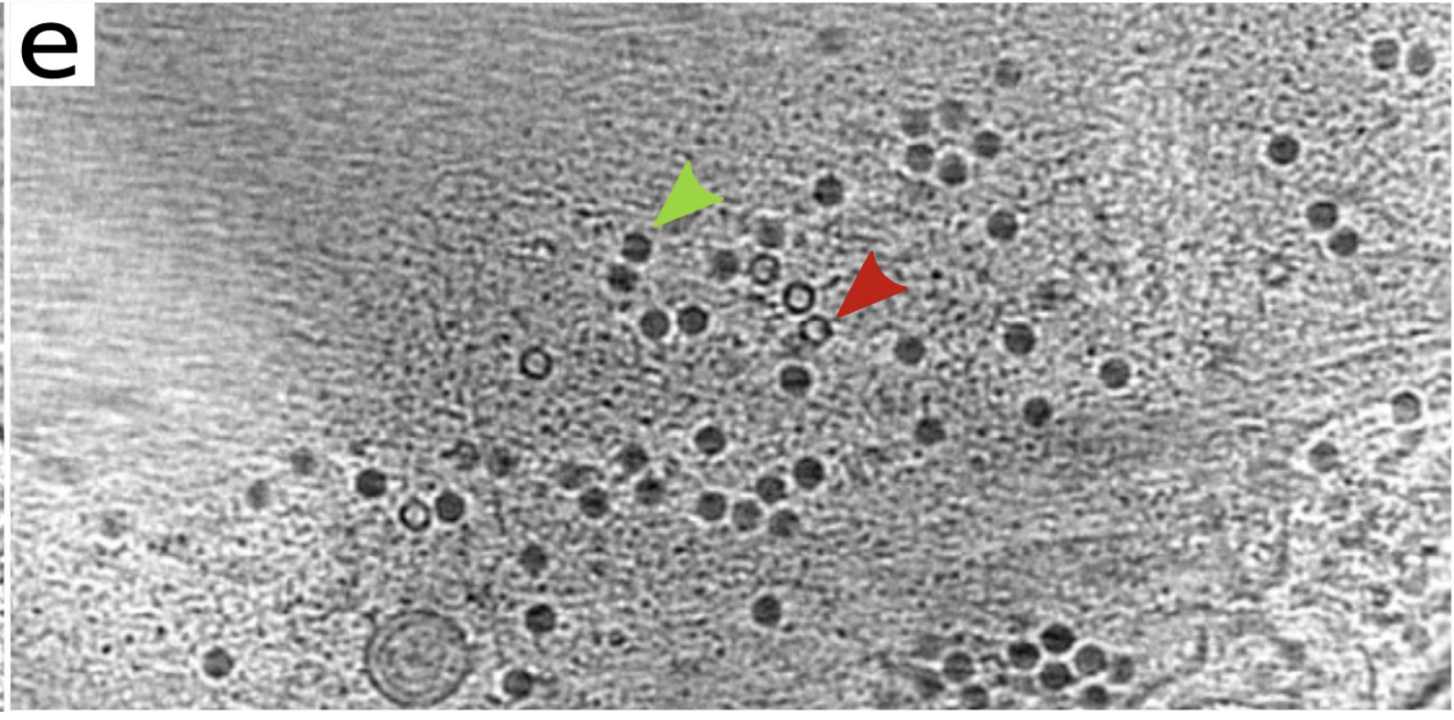
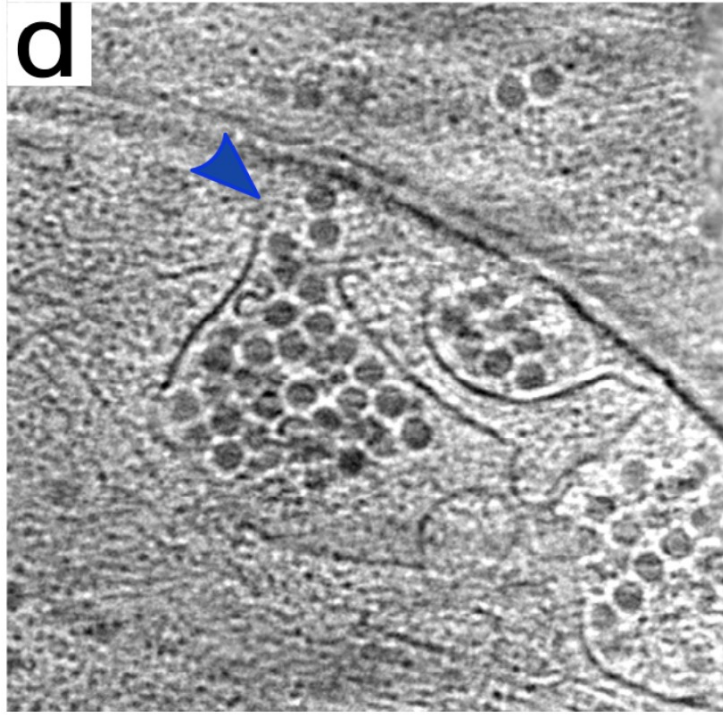
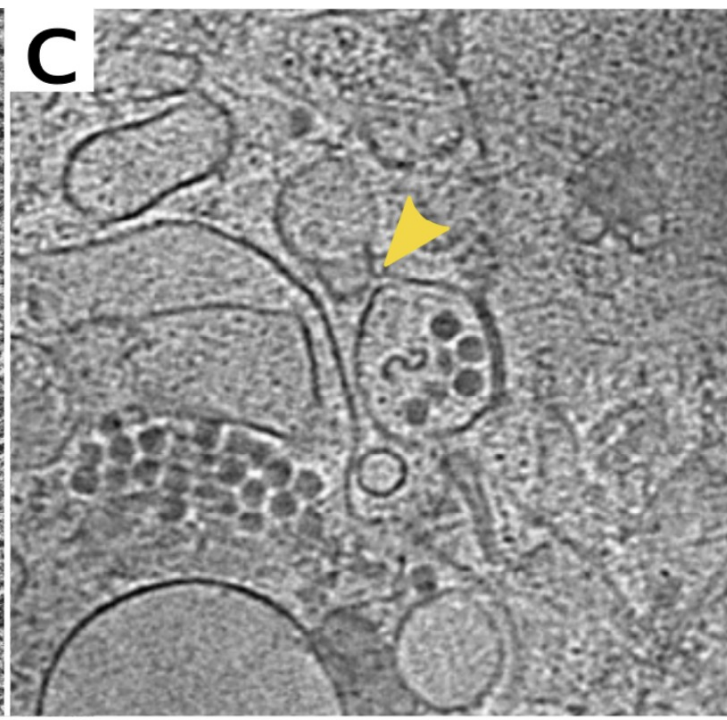
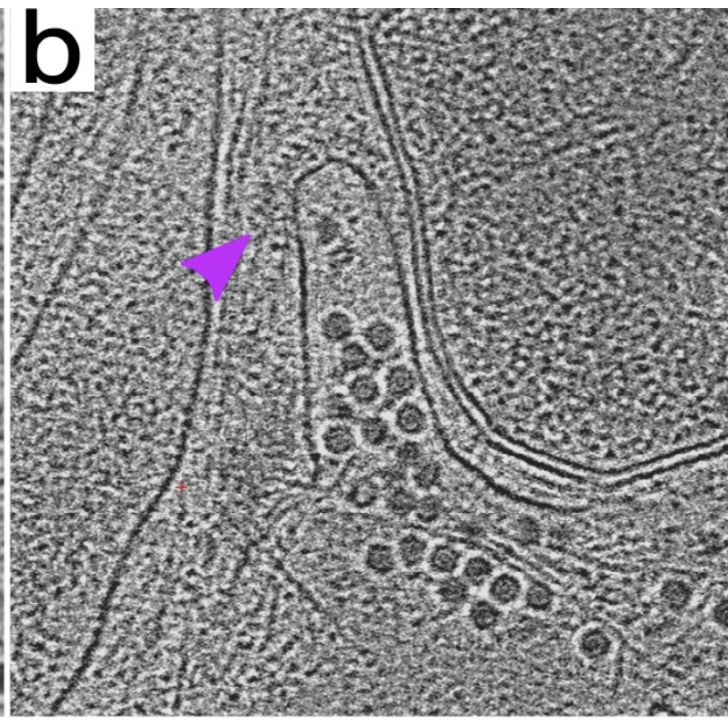
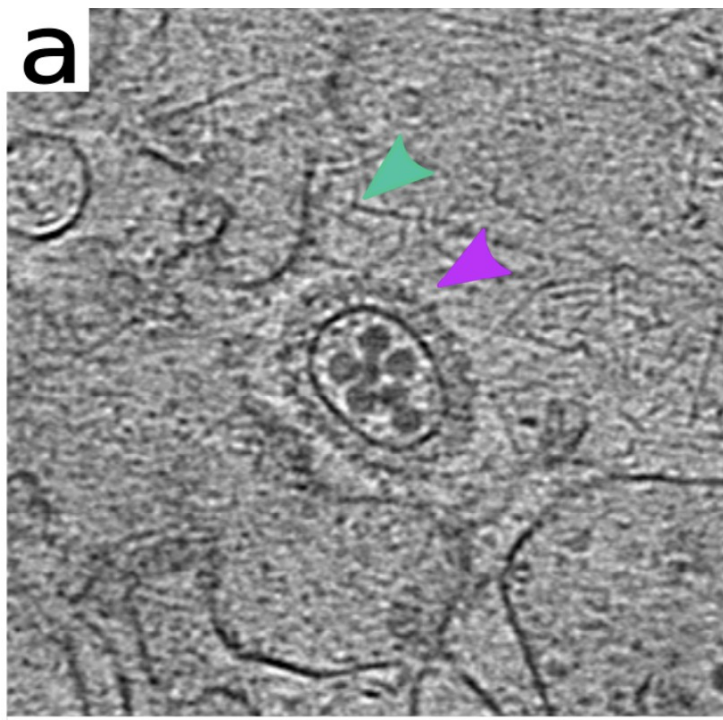
Cell entry of rhinovirus 2



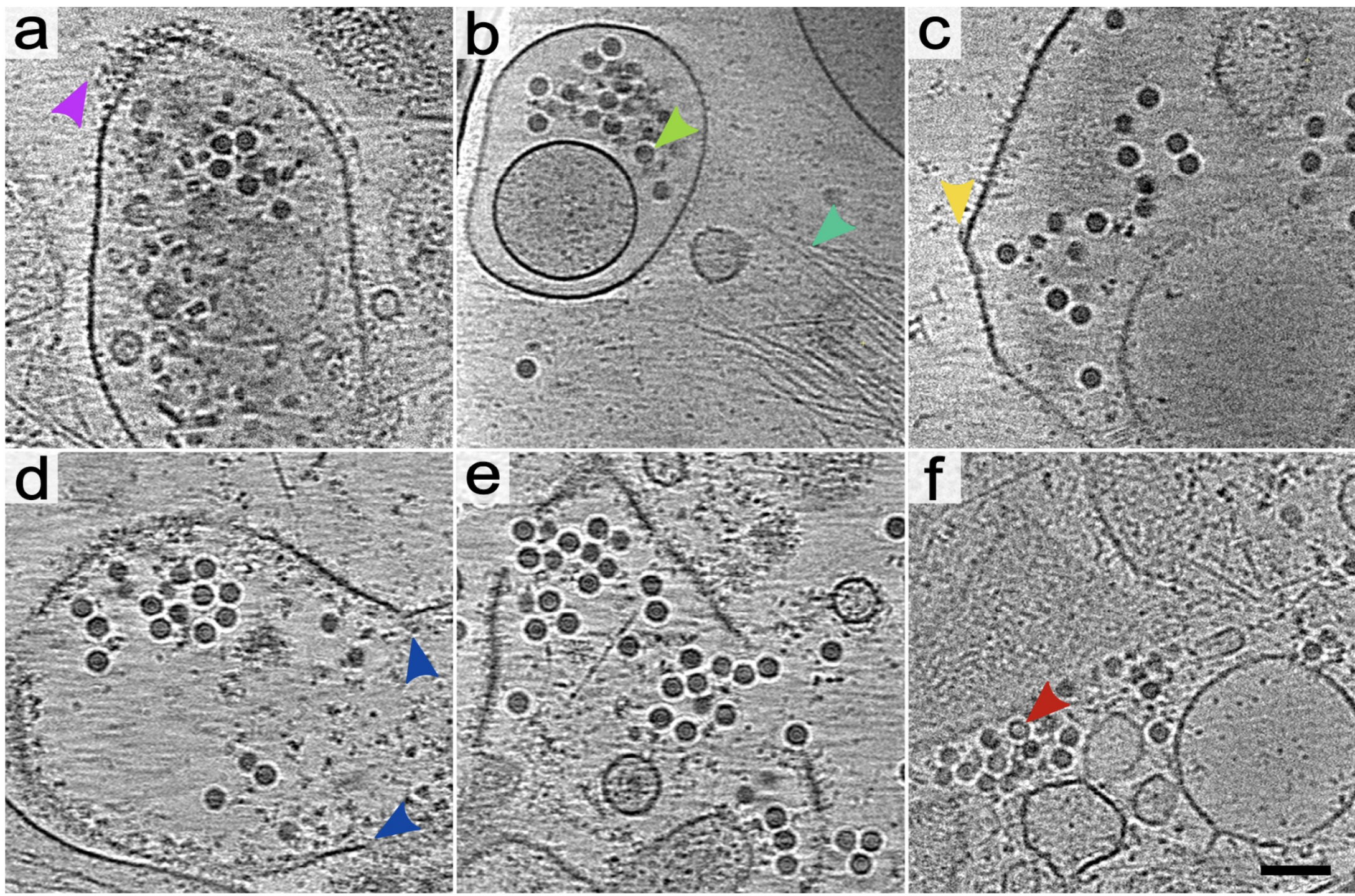
Cell entry of rhinovirus 2



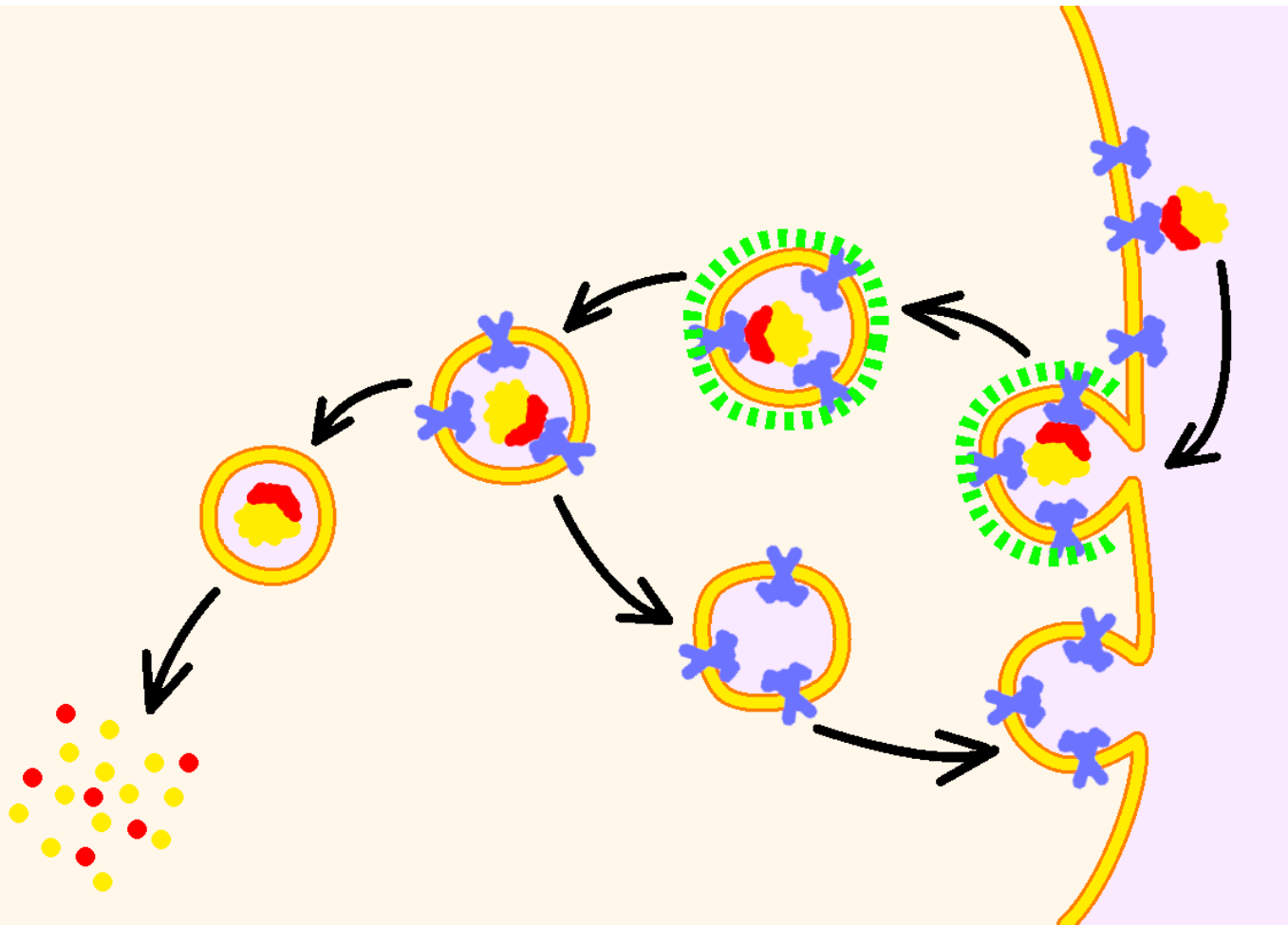
Cell entry of echovirus 30



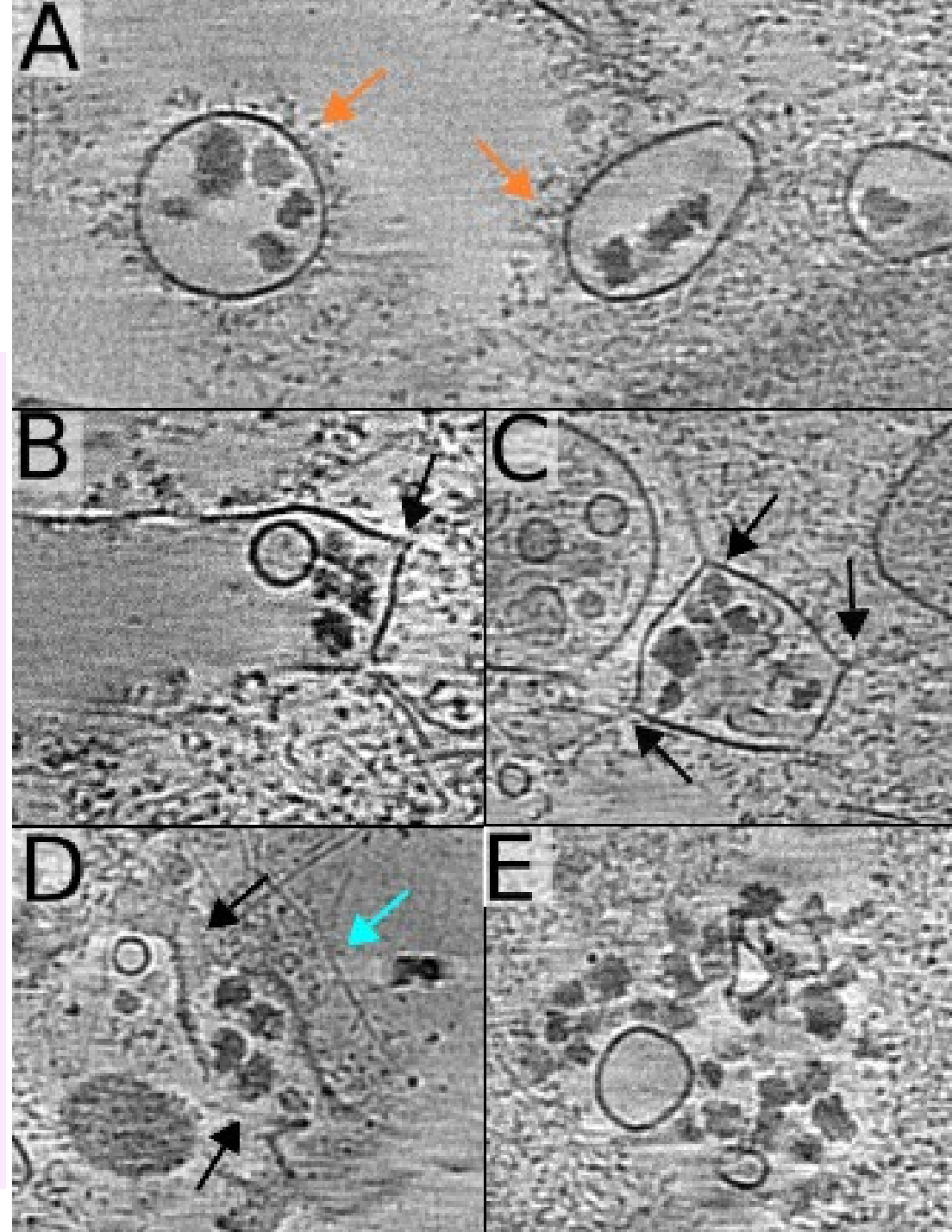
Cell entry of enterovirus 71



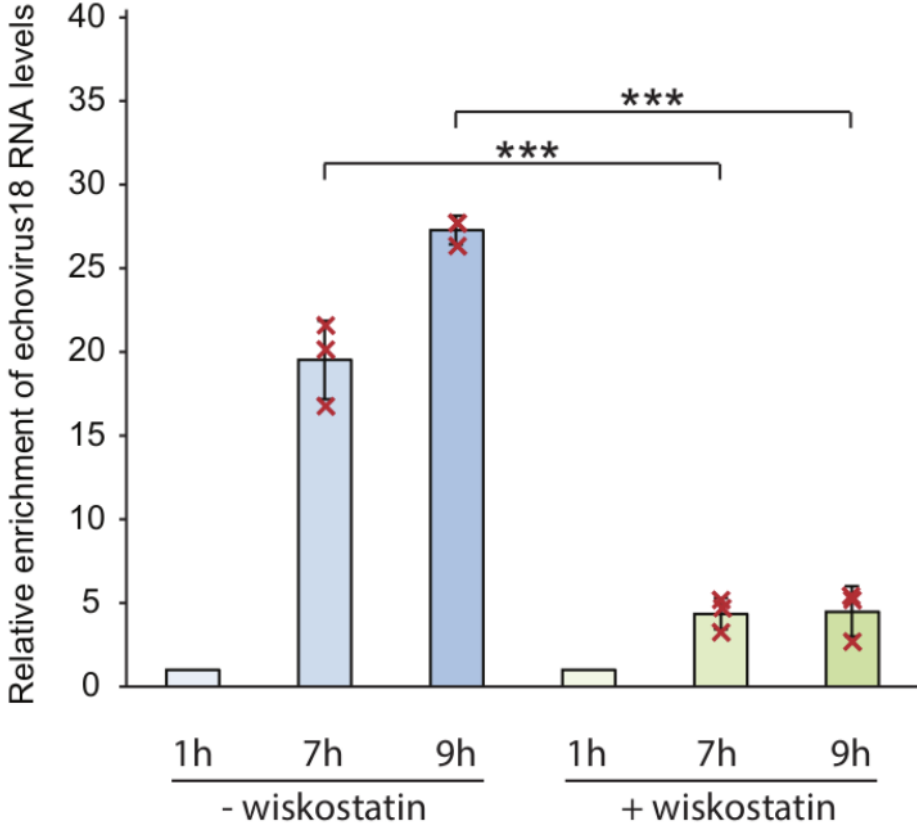
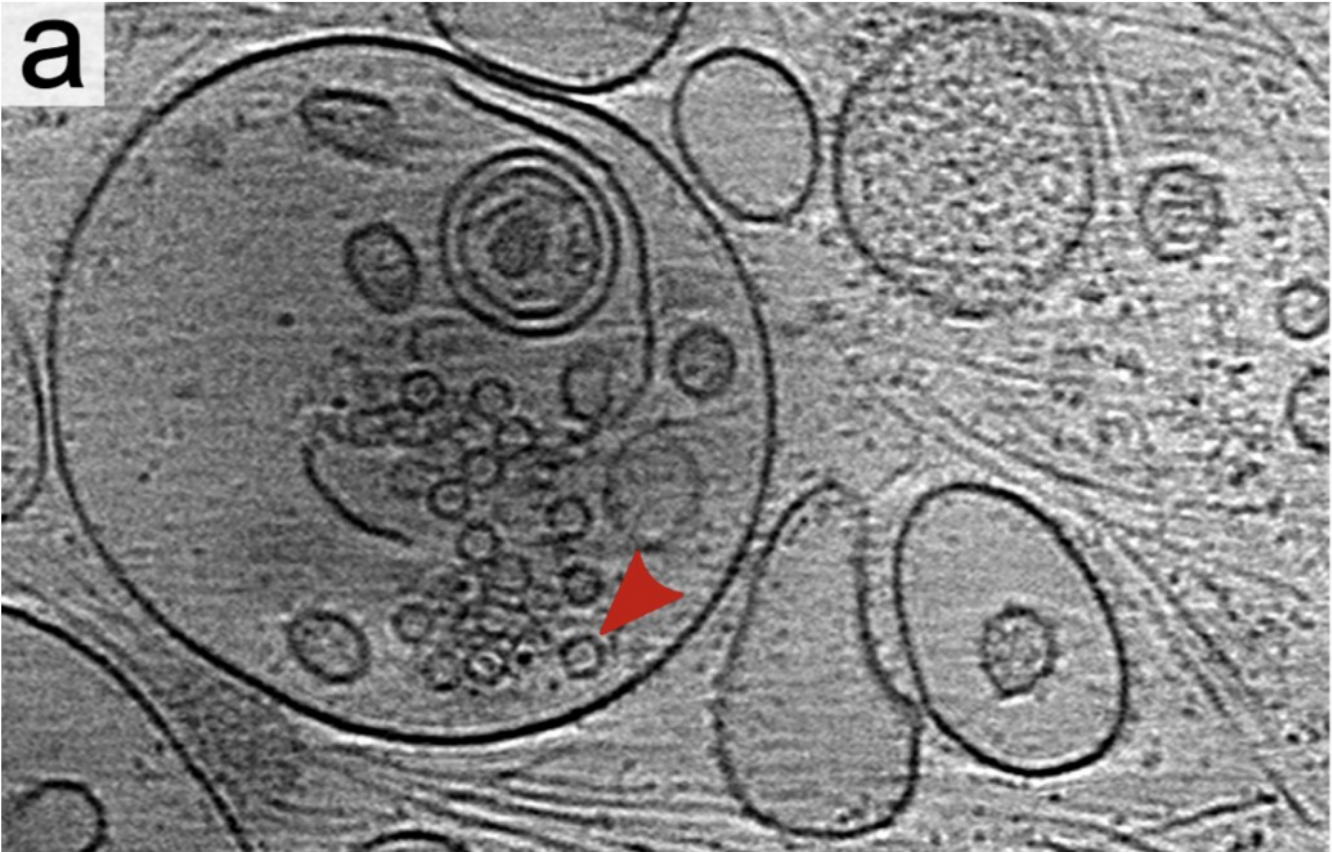
Endocytosis of VLDL - native cargo of VLDLR



Jeov et al. 2005



Wiskostatin inhibits enterovirus infection



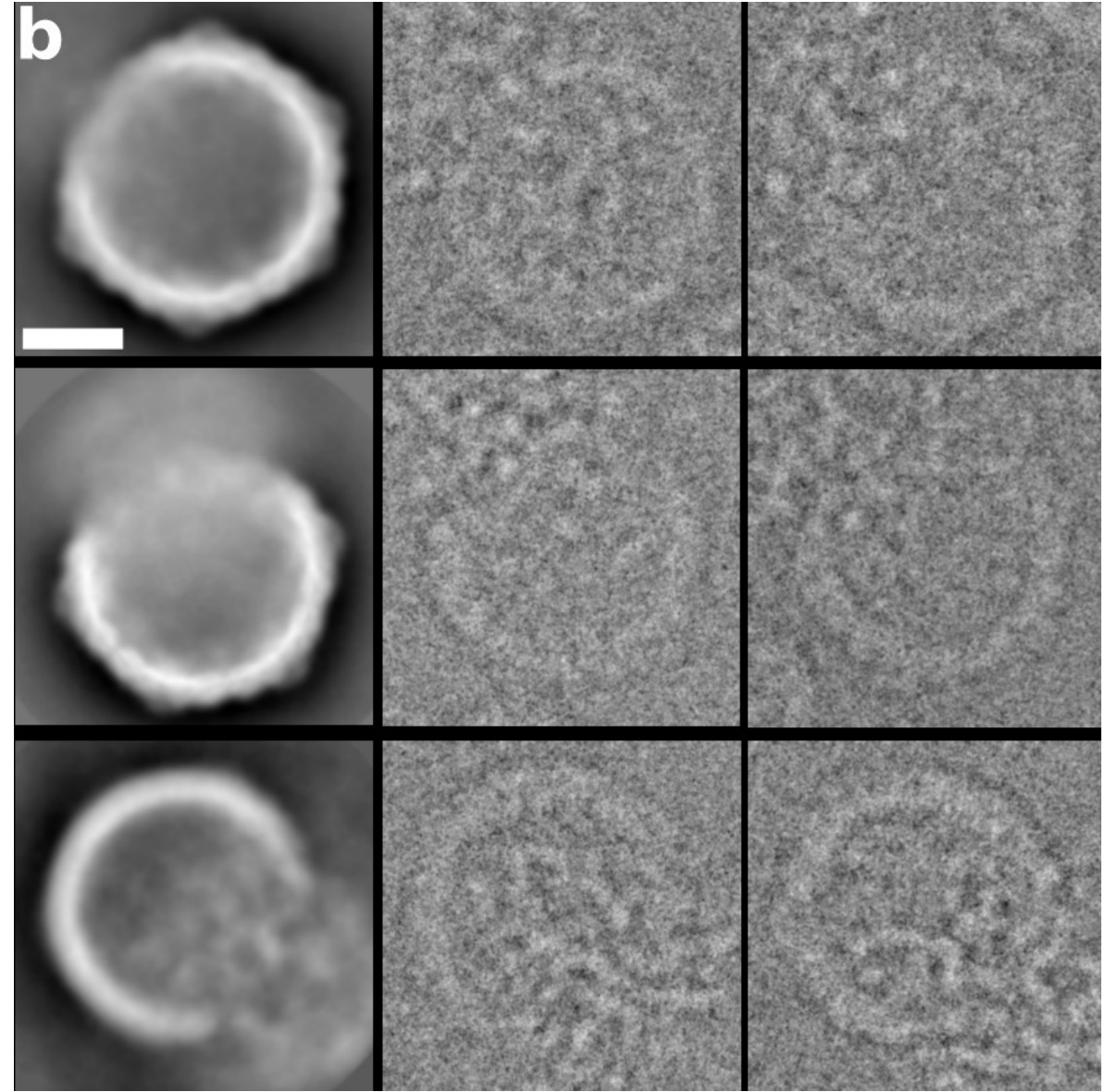
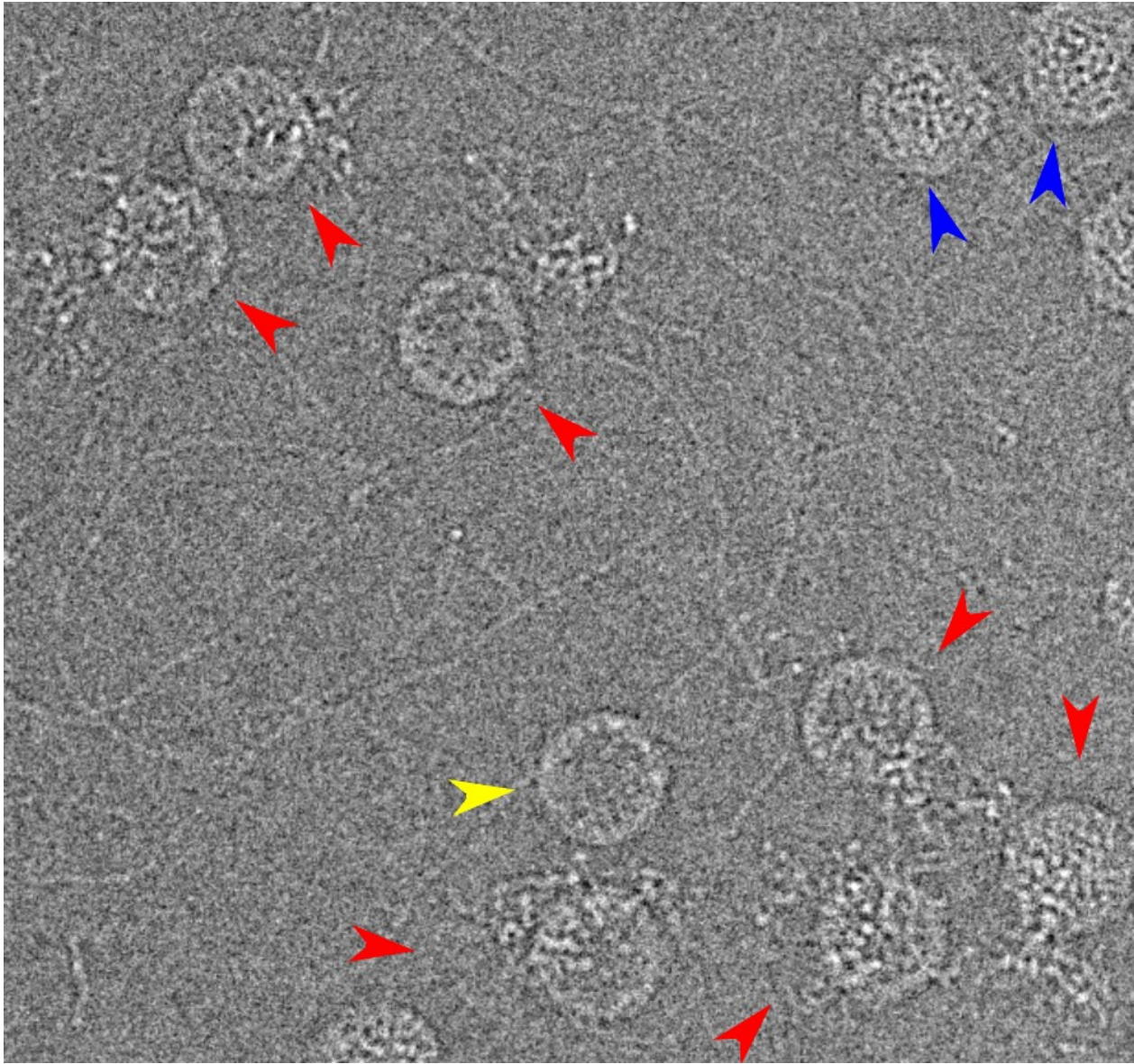
Wiskostatin

Endosomes
Cytoplasm

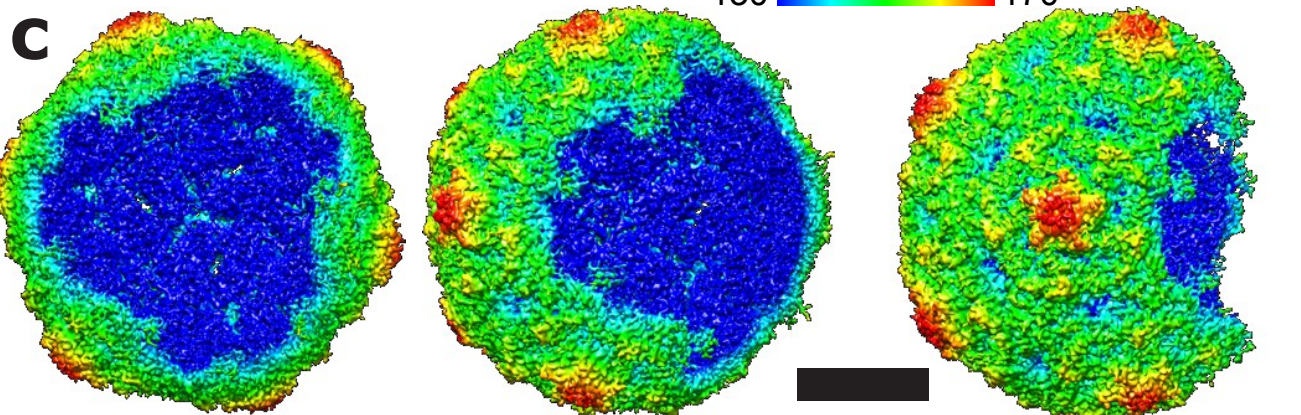
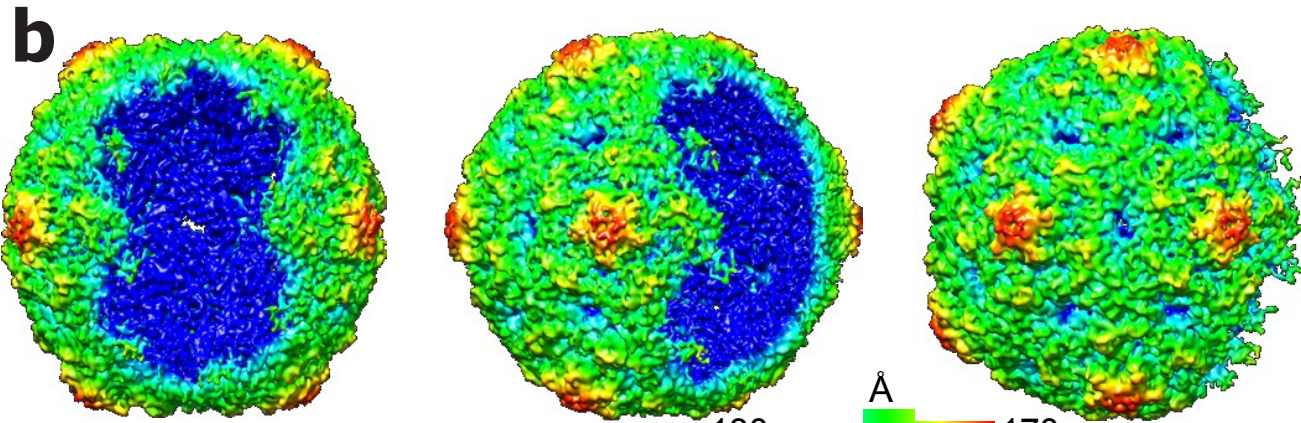
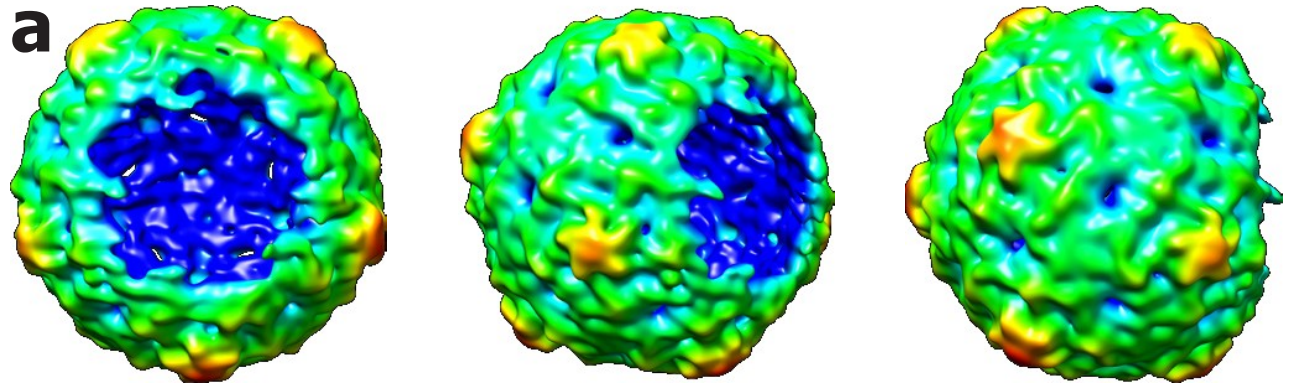


Fold change in the odds of genome release relative to infection of untreated cells

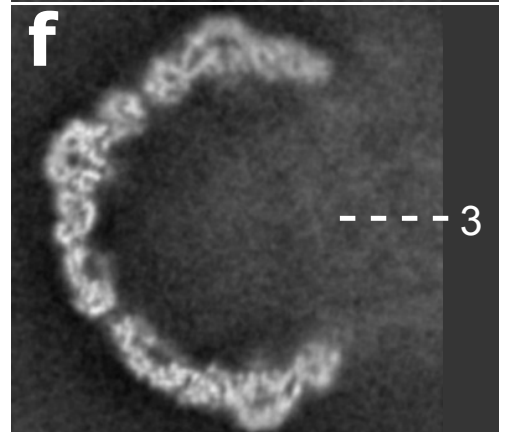
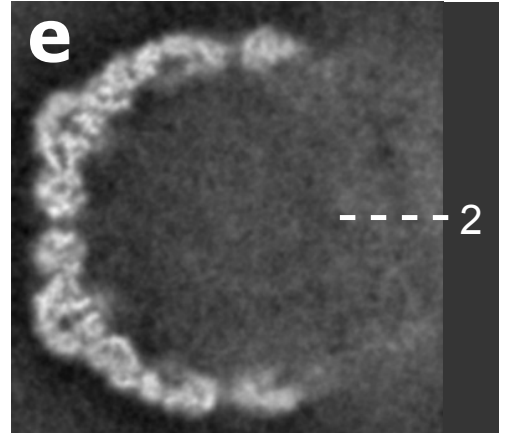
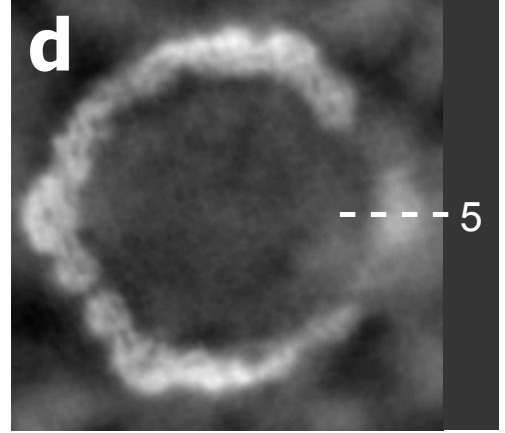
Genome release intermediates of echovirus 18



Open particles of echovirus 18



A
130 170

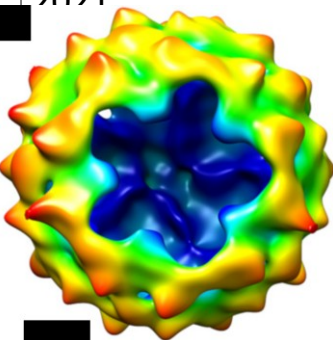
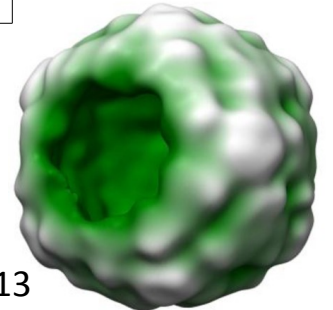


Echovirus 30
Picornaviridae
Buchta et al. 2019

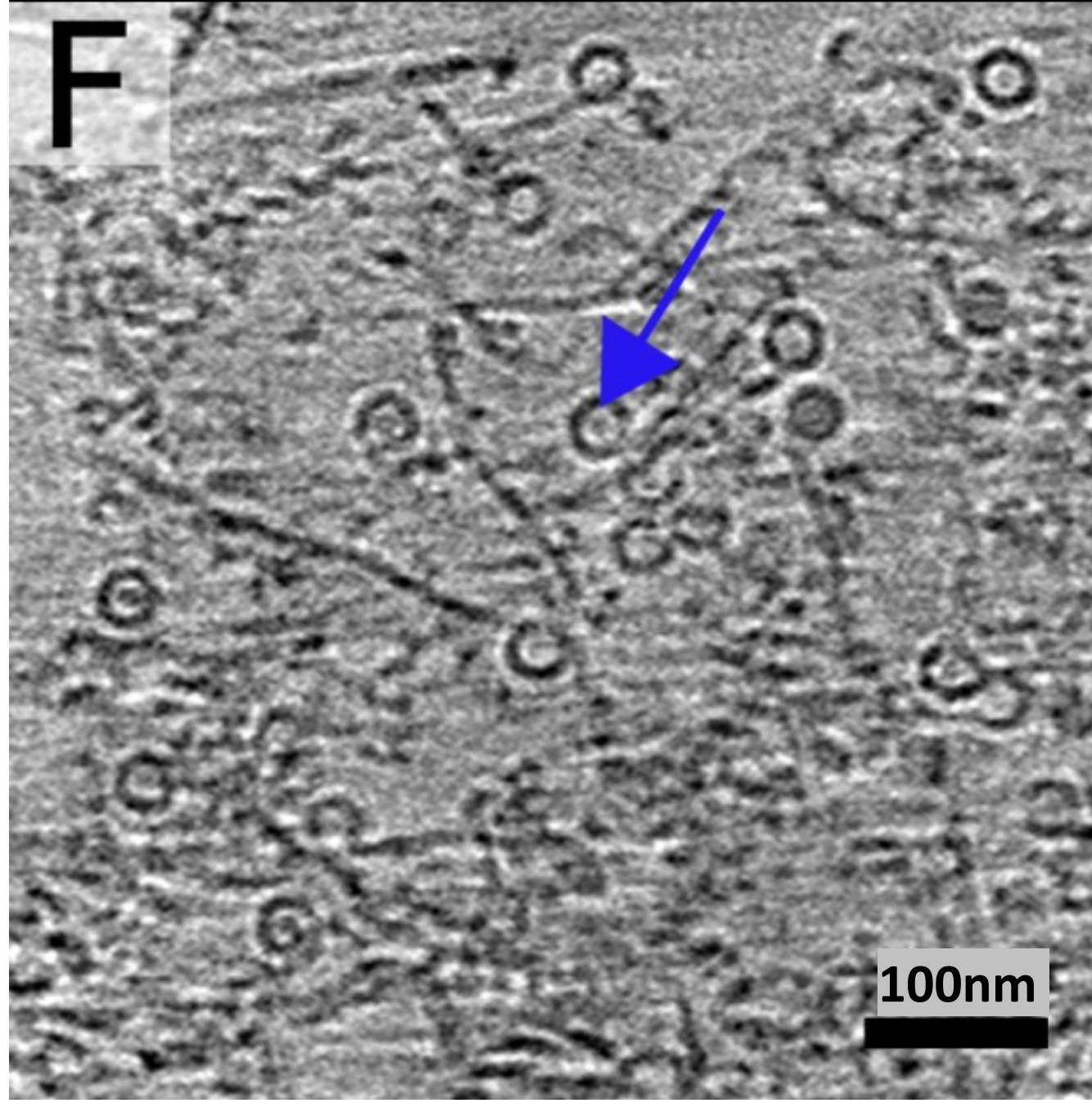
Human
rhinovirus 2
Picornaviridae
Harutyunyan et al. 2013

Slow bee
paralysis virus
Iflaviridae
Škubník et al.
2021

Kashmir bee virus
Dicistroviridae
Mukhamedova et al.
2021
Buchta et al. 2019

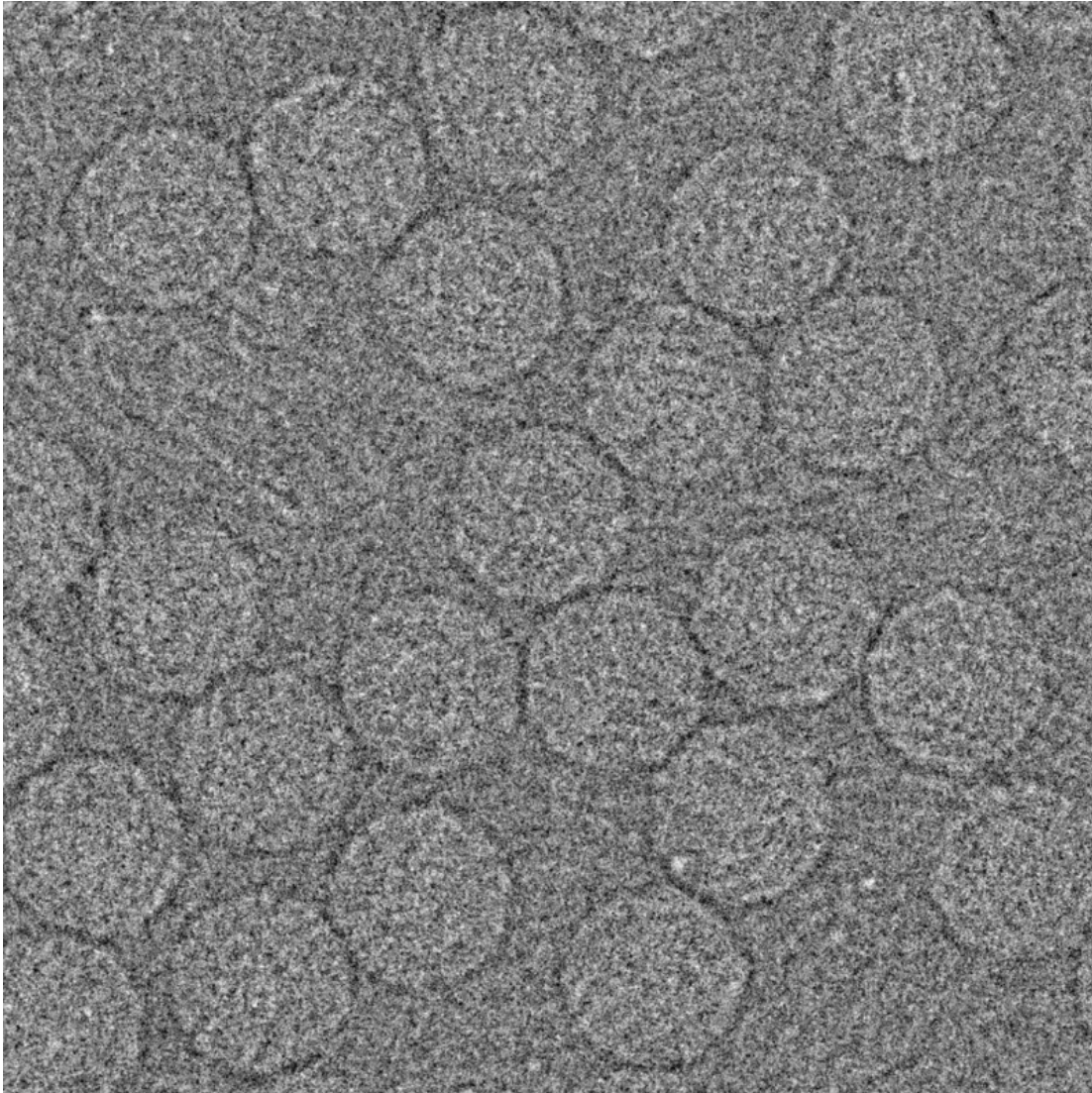


Cell entry of echovirus 18

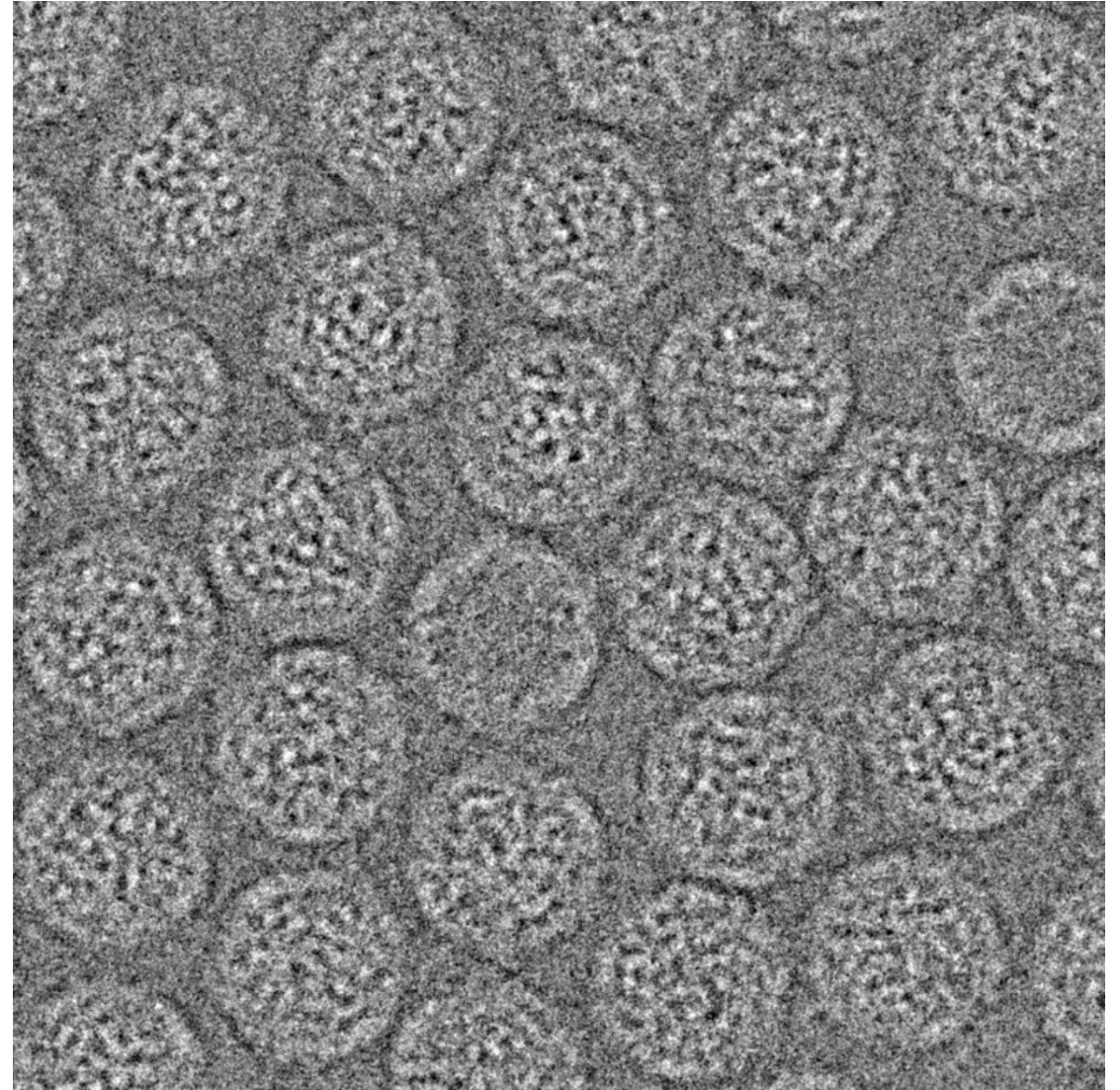


Acidic pH induces genome reorganization

Virions at neutral pH



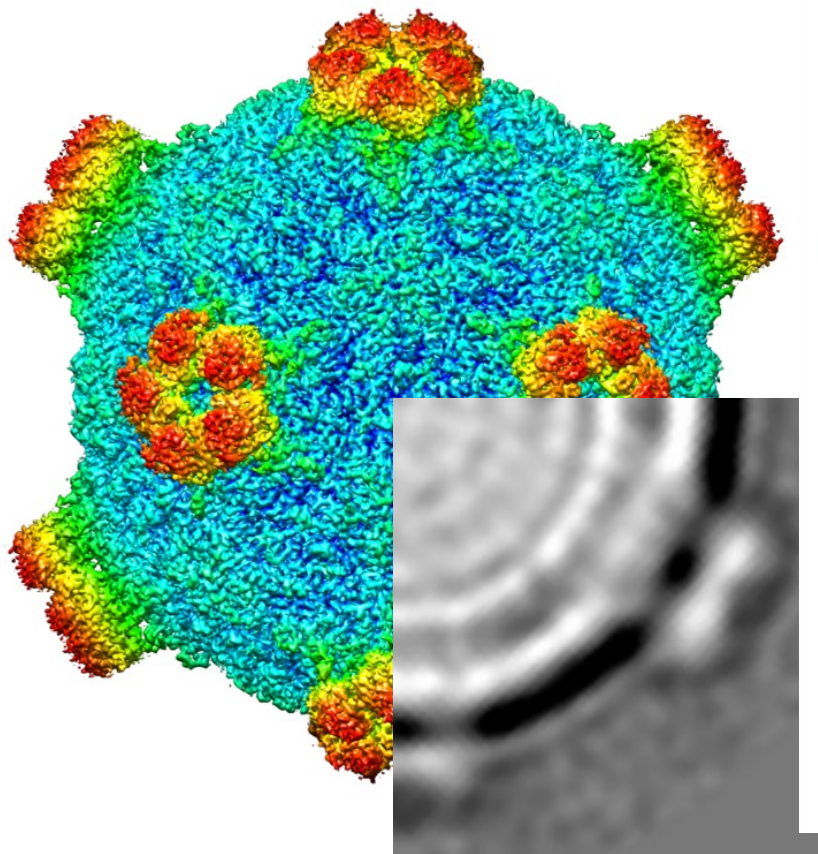
Activated particles at acidic pH



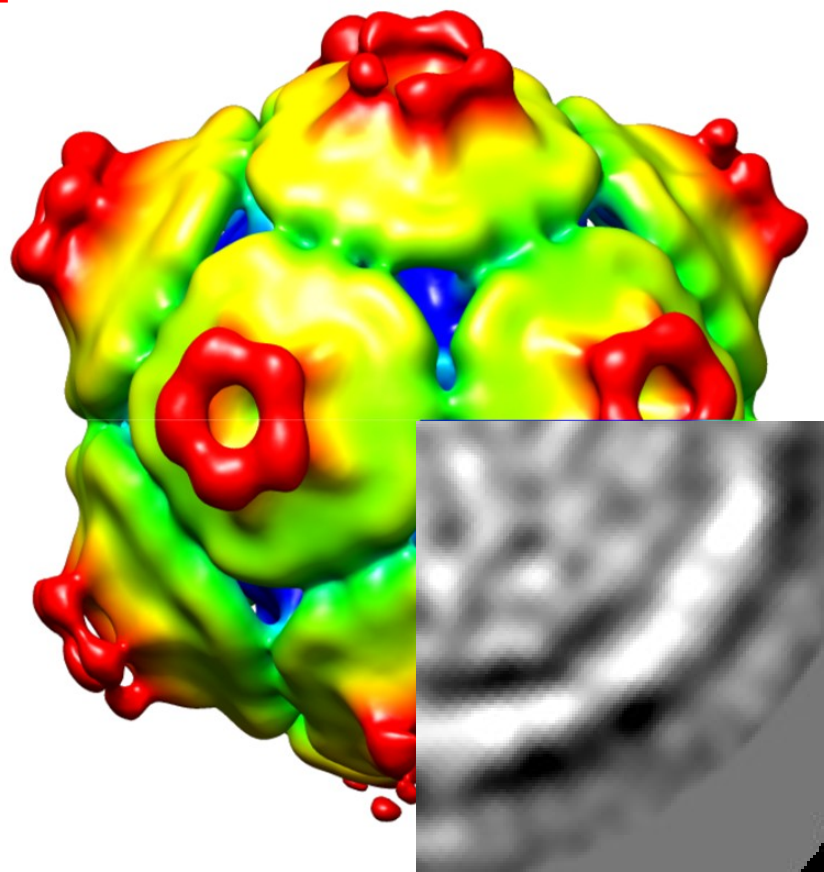
Deformed wing virus (*Iflaviridae*)

Neutral pH

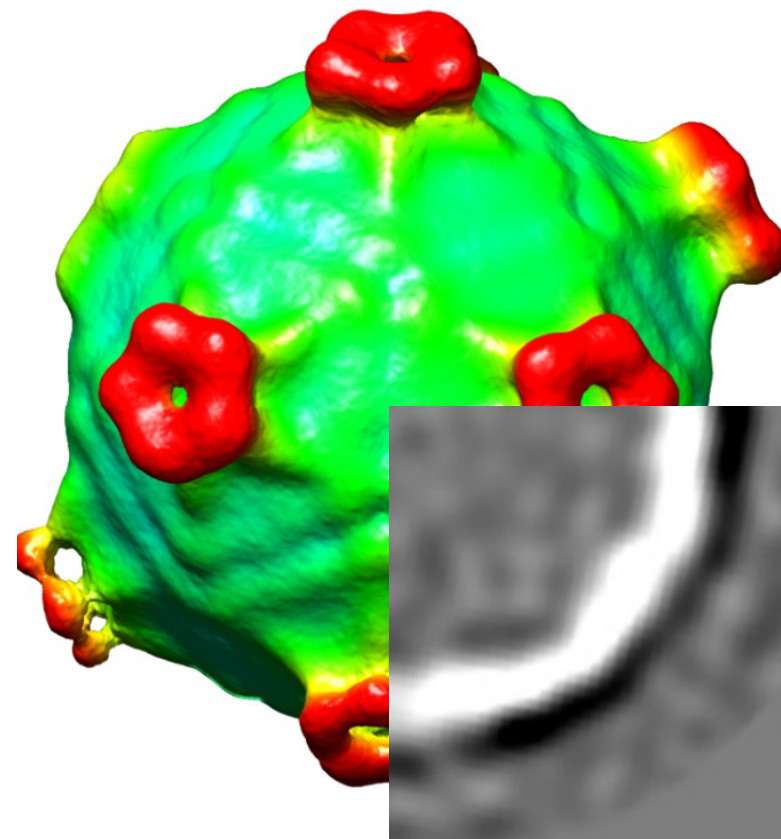
Acidic pH



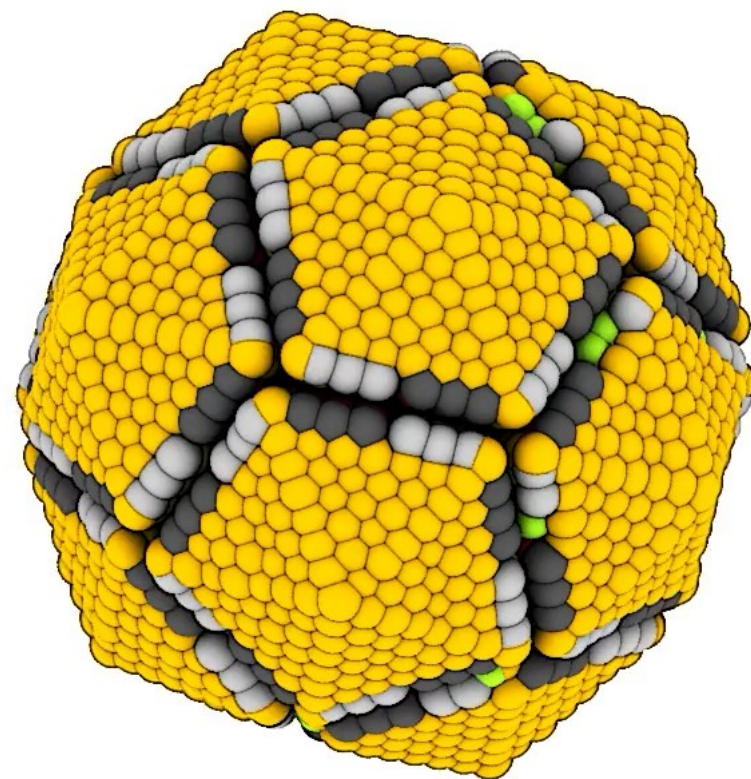
Virion



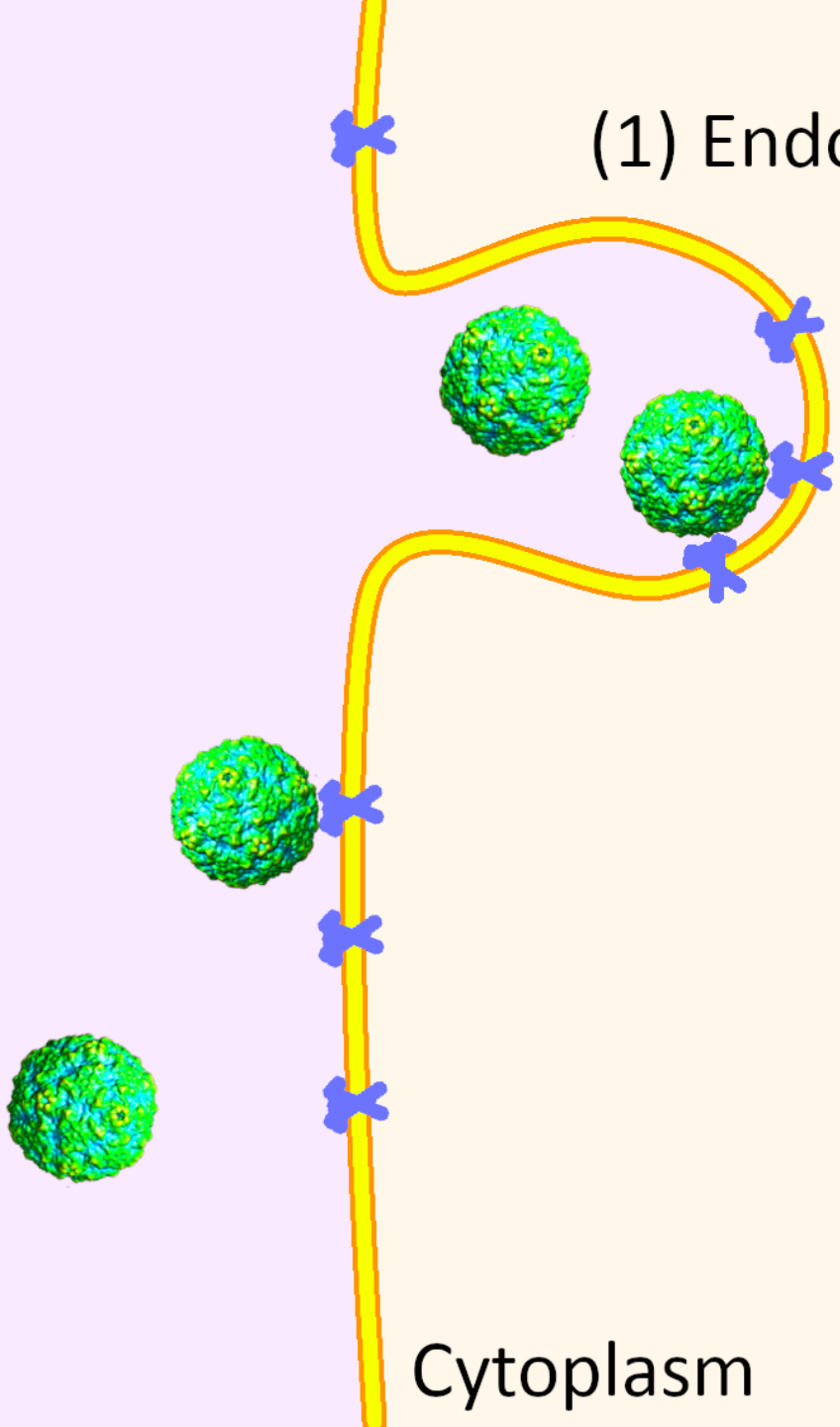
Full particle



Empty particle



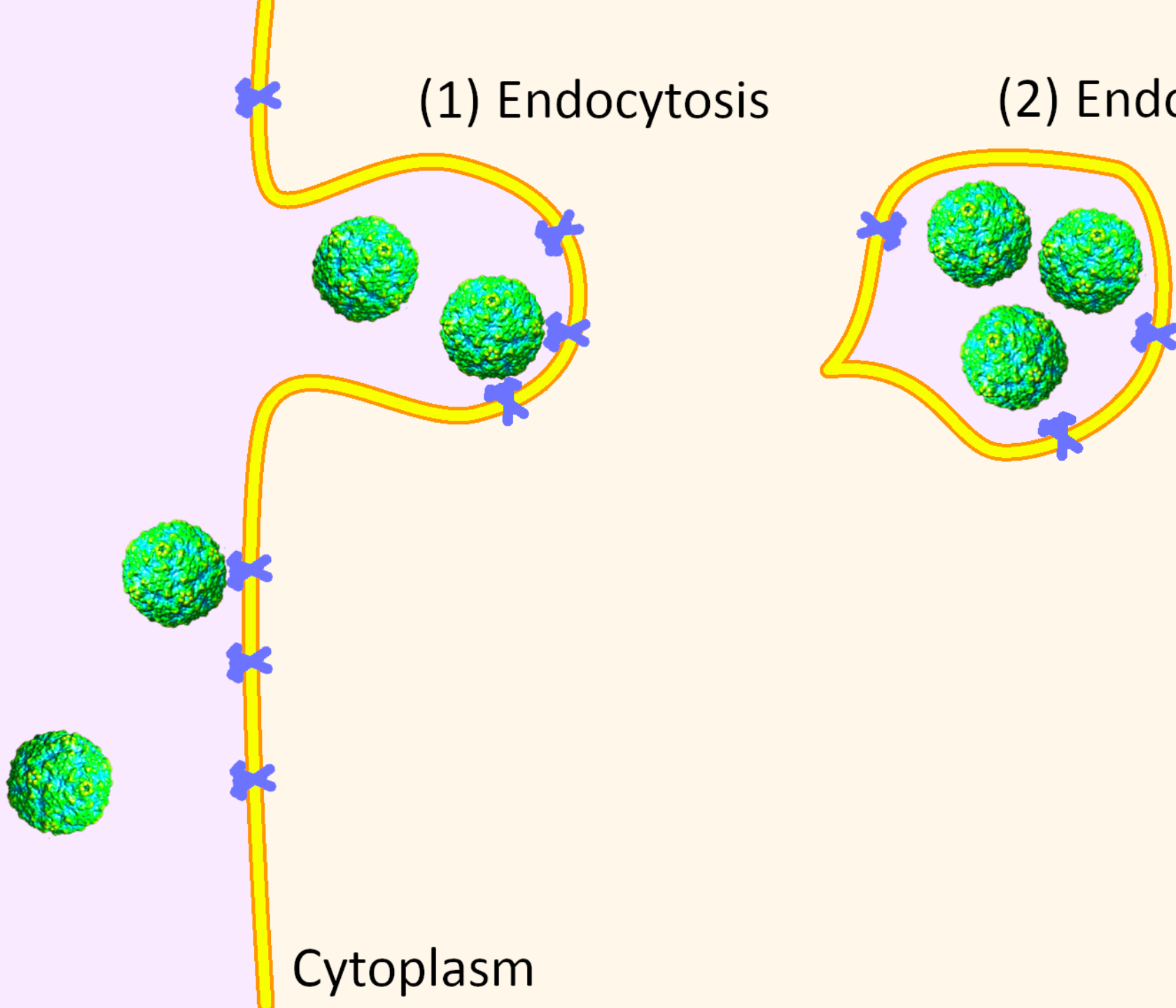
(1) Endocytosis



Cytoplasm

(1) Endocytosis

(2) Endosome



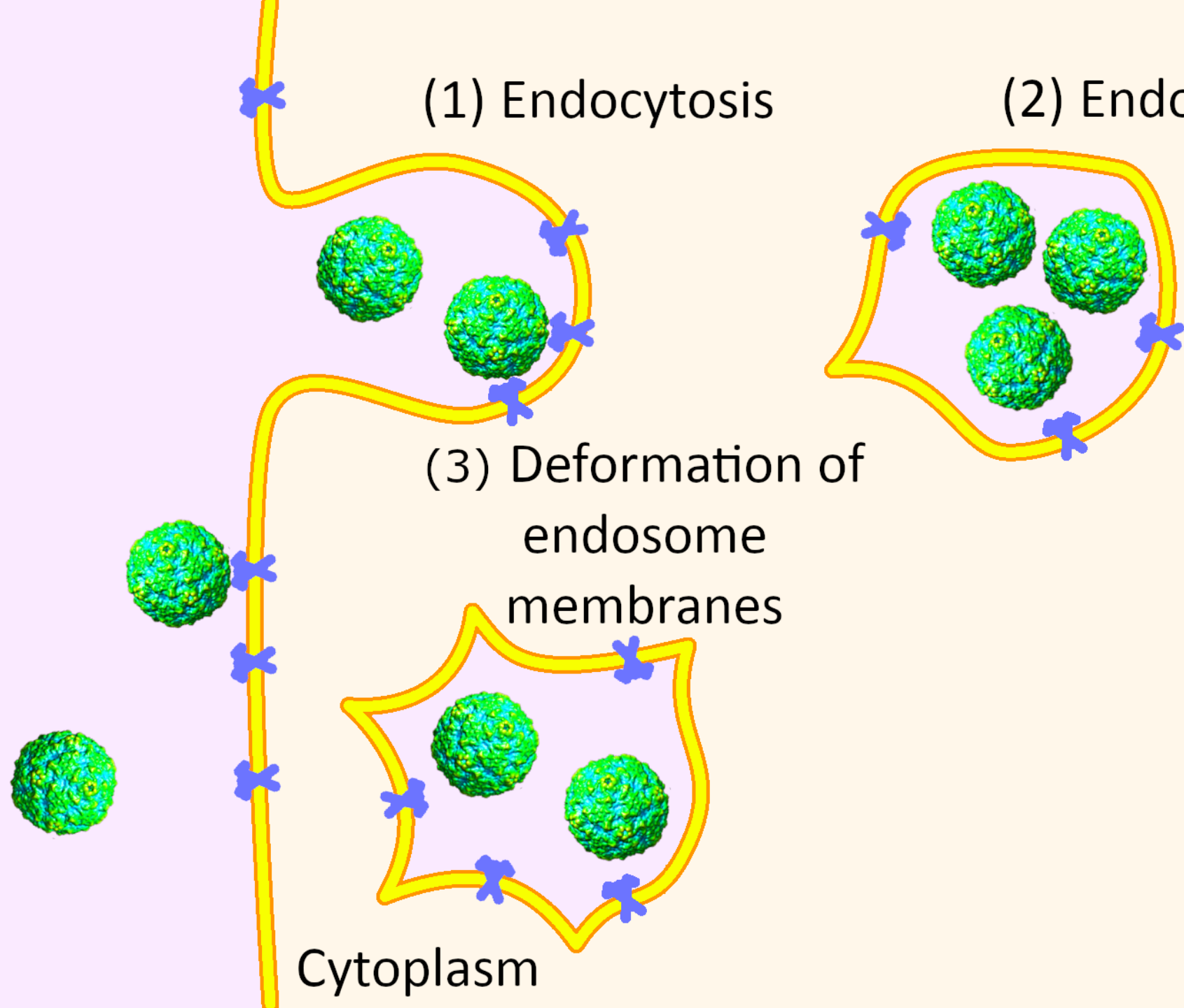
Cytoplasm

(1) Endocytosis

(2) Endosome

(3) Deformation of endosome membranes

Cytoplasm



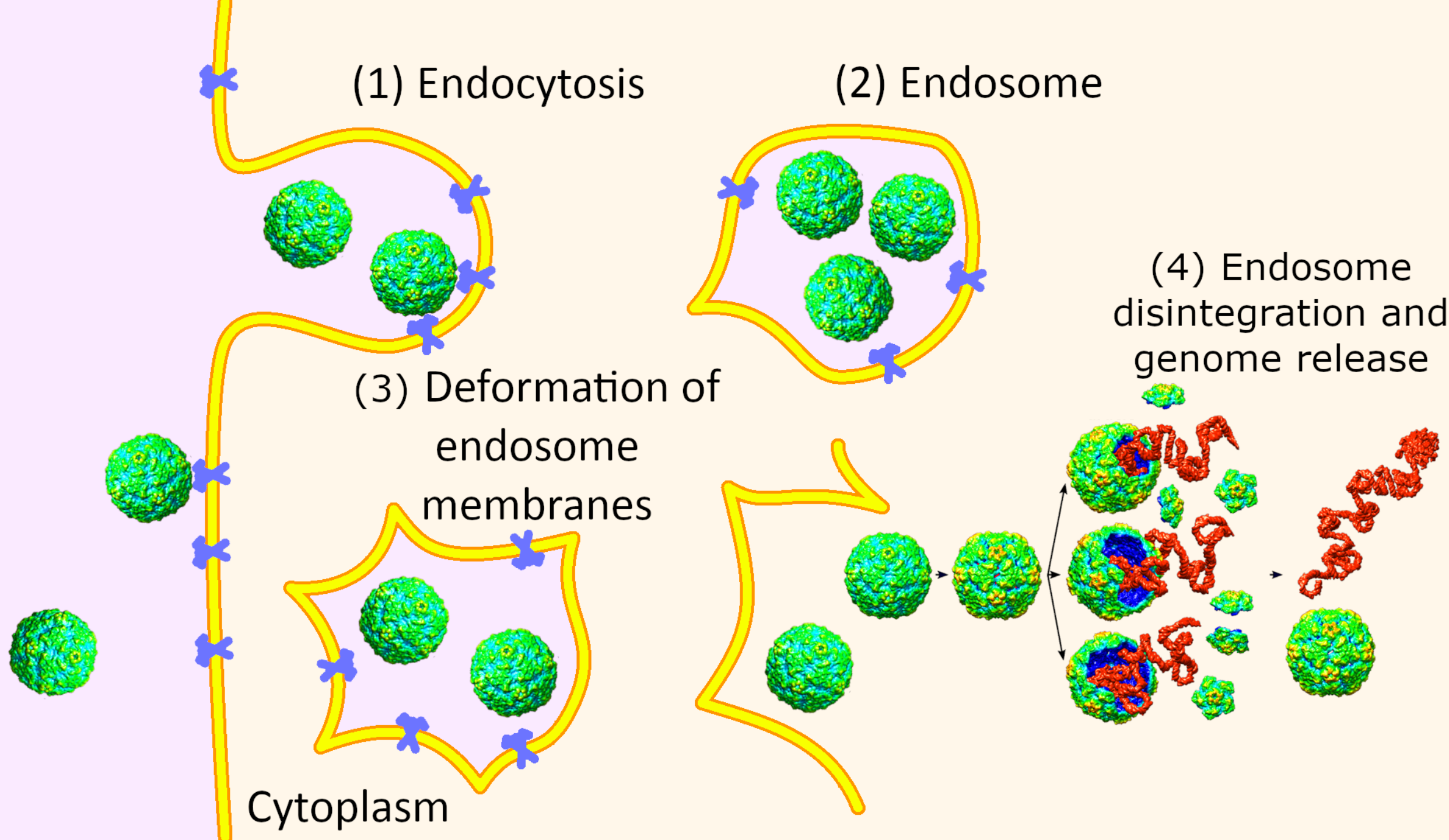
(1) Endocytosis

(2) Endosome

(3) Deformation of endosome membranes

(4) Endosome disintegration and genome release

Cytoplasm



1. Ktere z nize uvedenych zarení a castic interaguje nejsilneji s biologickym materialem?

a) elektrony

b) viditelne svetlo

c) paprsky X

2. Jake je v soucasnosti nejvyssi rozliseni dosazene pri studiu makromolekul pomoci kryo-elektronove mikroskopie?

a) 1.0 Å

b) 0.23 μm

c) 0.5 nm

3. Jake je nejvyssi mozne rozliseni obrazku, který ma velikost pixelu 1.1Å?

a) 1.1 Å

b) 2.2 Å

c) 3.3 Å