

آخر محاضرة فاينال

3- Laboratory Diagnosis and Treatment of Viral Infection

Mohammad Altamimi, MD, PhD
The Hashemite University, 2024

تفريغ : أمانة بعارة

Objectives

1. Understand principles of laboratory diagnosis of viral infections
2. Differentiate methods of viral detection and isolation
3. Describe viral reaction to physical and chemical agents
4. Describe and apply common methods of Inactivating viruses
5. Understand principles and classes of anti-viral agents
6. Understand principles, types, and application of viral vaccines

Difficulties

هاي صعوبات زمان

- Can not be seen under light microscope
- Can not be cultivated easily
- Do not grow on culture media
- Treatment was not available

١) ما يقدر نشوف تحت
light micros

٢) ما يقدر نزرع الفيروسات بسهولة

٣) ما يتفوع على ال
culture media

٤) ما كان في علاج

هنا تغير الوضوء

Changed situation

- ١- Rapid techniques فحوصات سريعة
- 2- Screening for Blood transfusion →
- 3- Treatment available
العلاج ما متوفر

مرنا خرف انه يستقل عن
لمريضة الدم

Specimens

(العينات)

(أنا باخذ عينة من المكان يلي فيه ال infection)

According to the disease

● Respiratory – Throat swab ^{مسحة من الحلق}
إذا مرض بالجهاز التنفسي
باخذ عينة عن طريقه
(Throat swab)

● CNS – CSF ^{cerebral spinal fluid}
إذا مرض بال CNS
باخذ عينة من هذا السائل CSF

● Eyes- Conjunctival scrapings

● Viremia – Blood ^{دم مصنف}
عن طريقه ^{دم في الميوس}
blood

● GIT and Liver - Stool ^{GIT}
إذا عندي مرض بالجهاز الهضمي أو بال liver

● Skin - Scrapings ^{عن الجلد / الجرح}
إذا عندي مرض بال skin
باخذ عينة عن طريقه
scraping



Specimen Storage and Transport

هون بقالح احفظ العينات عند درجة حرارة 4°C (في الدم)
يعني زي ال urine مثلاً

- Keep specimens other than blood at 4°C
- If delay $>24\text{hrs}$, freeze at -70°C or below.
← از اسكان التخزين
من 24 ساعة
حفظها عند -70°C
- Avoid any storage at -20°C ; greater loss in infectivity
← لأنه يجعل فقدان الفيروسات الفيروسي
- Nonenveloped viruses more stable than enveloped
← الفيروسات غير المغلفة أكثر استقراراً من المغلفة
- Viral Transport Medium
 - Salt solution – ensures proper ionic concentrations
محلول ملحي ٥ أنه يضمن تركيزات أيونية مناسبة
 - Buffer - maintains pH
 - Protein - for virus stability
كما فعل على استقرار الفيروس البروتين
 - Antibiotics or antifungals – to prevent contamination
لتجنب التلوث

1. Microscopy

يعني العيالك الموجودة جوا
السيوبلارم بعتد الوفا
بس عند لمحة ال light
micro

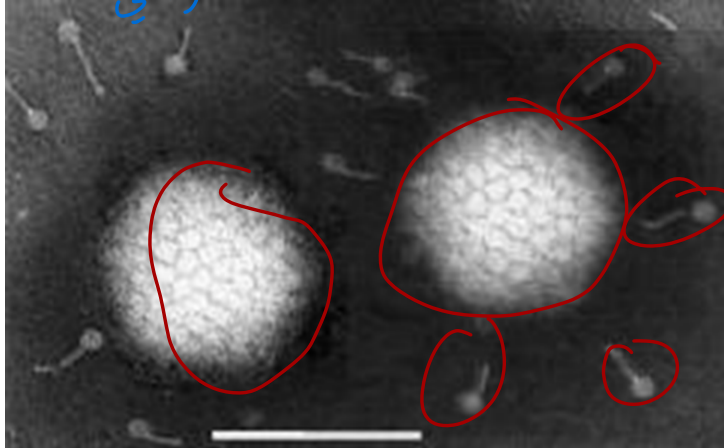
- ① Electron Microscope
- ② Light microscope – Inclusion bodies
- ③ Fluorescent Microscope - Fluorescent antibody technique

الاجسام الادخالية

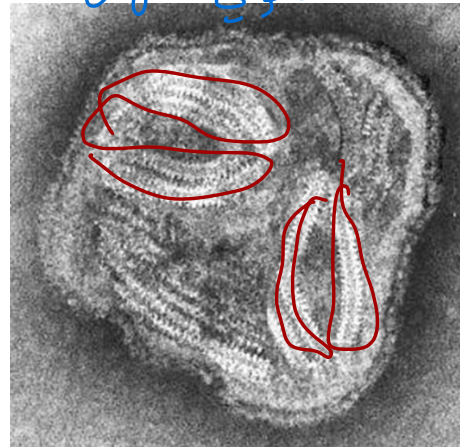
تقنية الاجسام المتعددة الفلورية

المجهر
الفلوري

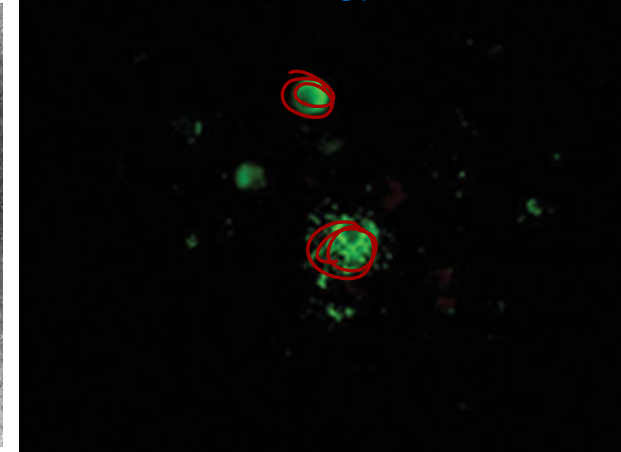
① الميكروني
micro



② الفلوري Light



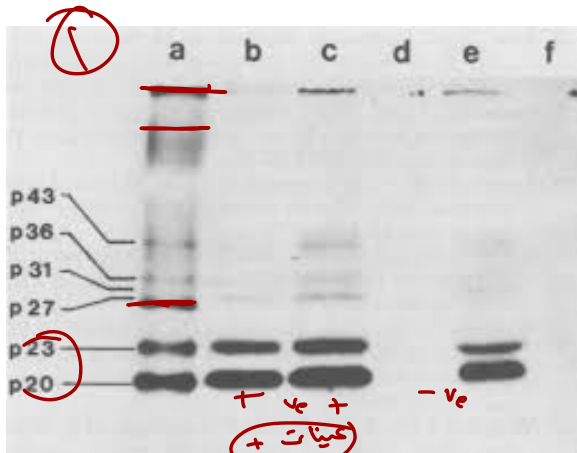
③ الفلوري fluorescent



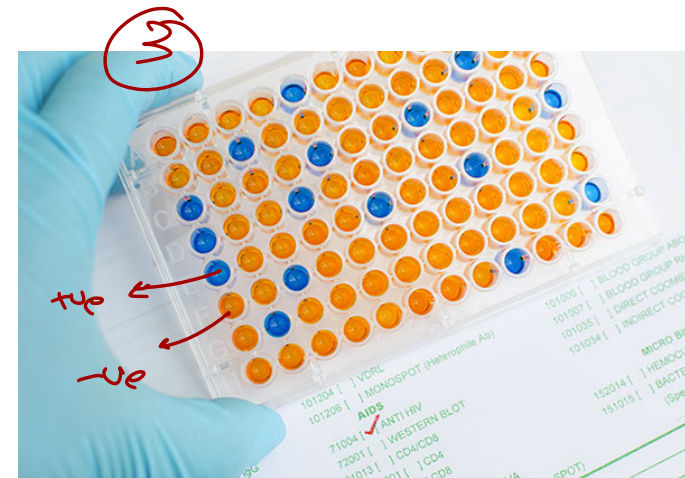
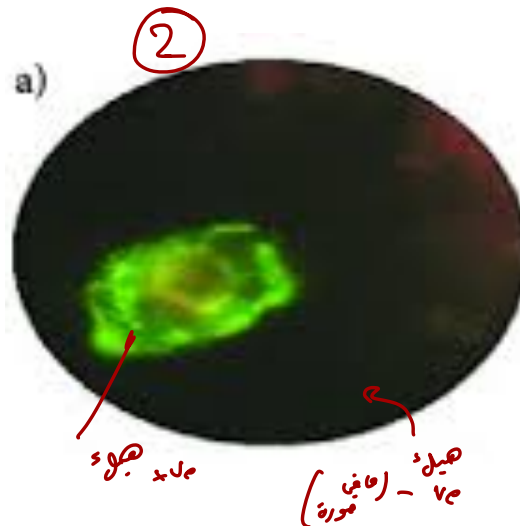
2. Demonstration of Viral Antigens

للمرئقة الثانية بالتشخيص

- ① ● Precipitation on gel eg HBsAg ترسيب الجل
- ② ● Immunofluorescence
- ③ ● Enzyme Linkes Immuno Sorbant Assay (ELISA)

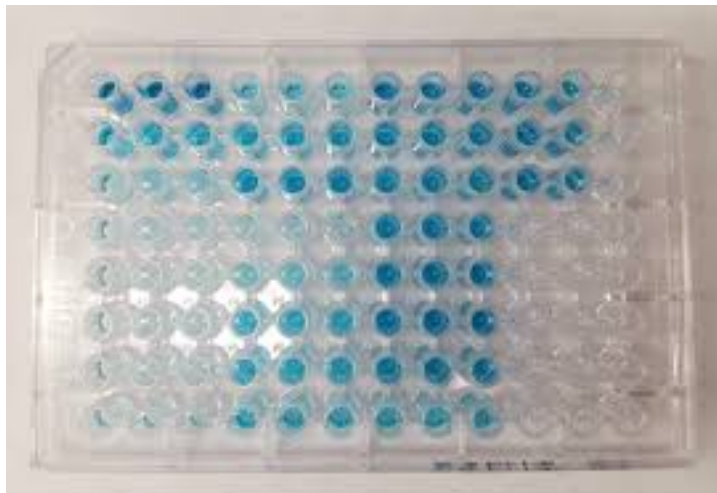


(از موجوده نتیجه +)

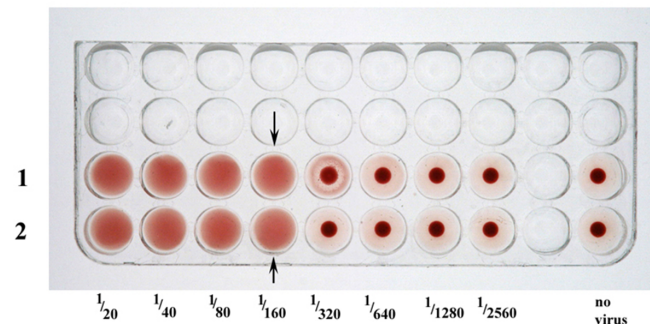


3. Serological Reactions (anti-viral antibodies)

- زيادة العيار للأضداد الخفيفة • Rising titre of antibody in paired sample of sera for IgG antibody
 - First sample – At the earliest
 - Second sample – After 2 weeks
 - Single sample IgM type of antibody detection
 - Techniques – ELISA, Haemagglutination Inhibition (HAI) Test
- اختبار تثبيط التجلط الدموي



Haemagglutination assay (HA assay)



1 and 2 are duplicates

The arrows show the last well in which haemagglutination was seen.

4. Molecular Techniques

لَقْنِيَات تَقْضِيْم الكَمْف النَوَوِي

- Nucleic acid amplification techniques such as polymerase chain reaction (PCR) can be used to detect viral genomes in clinic material.

- To detect RNA, an initial reverse transcription step is performed (converts RNA into cDNA). After this, PCR can be performed.

Molecular assays are very sensitive (able to detect only a few viruses in a clinical sample.)

- They can also be used to measure the amount of virus (viral load) in a patient's sample.

لَقْيَاس كَمِيَّة الْفِيرُس



هون يغير
الحكمي
يتحول
RNA
DNA

5. Viral Isolation and Culture

اهداف الزراعة

عزل الفيروسات
من العينات السريرية

Primary purposes of viral cultivation

- ① To isolate and identify viruses in clinical specimens
- ② To prepare viruses for vaccines
- ③ To do detailed research on viral structure, multiplication cycles, genetics, and effects on host cells

عشان نعمل دراسات مفصلة
Structure, multiplication, genetics

Using Live Animal Inoculation

استخدام حقن الحيوانات الحية

Specially bred strains of white mice, rats, hamsters, guinea pigs, and rabbits

ضائير غينيا

Animal is exposed to the virus by injection



Using Bird Embryos

اجنة الطيور

Enclosed in an egg, nearly perfect conditions for viral propagation

ملائمة لتكاثر فيروسات

Chicken, duck, and turkey are most common

Egg is injected through the shell using sterile techniques

Cell culture for viral identification

زراعة الخلايا لتحديد الفيروس

يتم حقن البيض
بتقنيات معقمة



Cell Culture

Routinely used for growing viruses

Classified into 3 types:

① Primary cell culture – normal cells freshly taken from body & cultured, limited growth

Rhesus monkey kidney

→ Chick embryo fibroblast

② Human amnion cell culture

Diploid cell strains – cells of single type (fibroblast cells) that can be subcultivated for limited number of times, mostly 50

WI-38: human embryonic lung cell

HL-8: Rhesus embryo cell

③ Continuous cell lines – malignant cells, indefinite subcultivation

HeLa: Human Ca of cervix cell line

HEP-2: Human epithelioma of larynx

بتأخذ من الجسم
و بتزرع
عطون

نحوها محدود

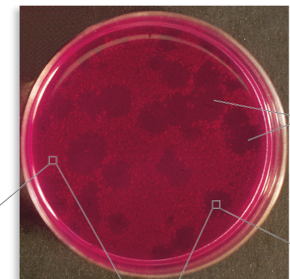
خلايا كلية
قرد الريسوس

→ الليفات جنين الدجاج

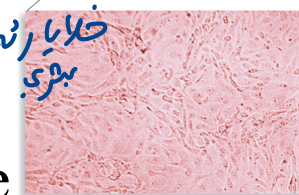
→ خلايا
الأمني البشري

خلايا جنين داء

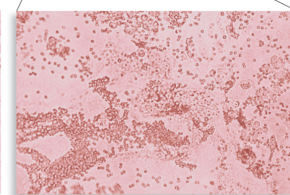
خلايا رئية جنين
بشري



Plaques



(b) Normal



(c) Infected

حنق الرحم

خلايا
الكلى

حنجرة

تحليلكم ترجمة هون قريوها وشوفوا Detection of Virus Growth Cultures



- **Cytopathic effects (CPE)** – morphological changes in cultured cells, seen under microscope, characteristic CPE for different groups of viruses
- **Metabolic Inhibition** – no acid production in presence of virus
- **Hemadsorption** – influenza & parainfluenza viruses, by adding guinea pig erythrocytes to the culture
- **Interference** – growth of a non cytopathogenic virus can be tested by inoculating a known cytopathogenic virus: growth of first virus will inhibit the infection by second
- **Transformation** – oncogenic viruses induce malignant transformation
- **Immunofluorescence** – test for viral Ag in cells from viral infected cultures.

Reaction to physical and chemical agents

1. Heat and cold:

- Icosahedral viruses tend to be stable, while Enveloped viruses are much more heat labile
- Viral infectivity is generally destroyed by heating at 50–60°C for 30 minutes بمنعطل نشاط الفيروسات عن طريق التسخين
- Viruses can be preserved by storage at subfreezing temperatures تحتفظ الفيروسات بحالة درجتها حرارة منخفضة

2. Salts:

- Many viruses can be stabilized by salts in order to resist heat inactivation و الملح بخلية تقاوم امتثال الحرارة

3. pH:

- Viruses are usually stable between pH values of 5.0 and 9.0. Some viruses (eg, enteroviruses) are resistant to acidic conditions. All viruses are destroyed by alkaline conditions. غير وسان محوية

لهم كالم بتعطلوا

4. Radiation:

فوق بنفسيه
الاشعاع
جسيمات عالية الطاقة تعمل تدمير الفيروسات
Ultraviolet, x-ray, and high-energy particles inactivate viruses

5. Detergents:

المنظفات
Solubilize lipid constituents of viral membranes and disrupt capsids into separated polypeptides

6. Formaldehyde:

يتشكل الكسولان البروتيني
Formaldehyde destroys viral infectivity by reacting with nucleic acid

7. Quaternary ammonium, organic iodine, low-concentration chlorine, and Alcohols are relatively not effective against viruses

Common Methods of Inactivating Viruses

-
- ^{التحقيق} Sterilization may be accomplished by ^{بخار تحت الضغط} steam under pressure, dry heat, ethylene oxide, and γ-irradiation
 - ^{مطهرات سطح} → Surface disinfectants include sodium hypochlorite, glutaraldehyde, and formaldehyde
 - ^{مطهرات الجلد} → Skin disinfectants include chlorhexidine, 70% ethanol, and iodophors
 - ^{تعطيل الفيروسات بانتفاخ لقاحات} → Vaccine production may involve the use of formaldehyde, ultraviolet irradiation, or detergents to inactivate the vaccine

Treatment and Prevention of Viral Infections

As bacteria and protozoa do not rely on host cellular machinery for replication, processes specific to these organisms provide ready targets for developing antibacterial and antiprotozoal drugs. However, because viruses are obligate intracellular parasites, antiviral drugs must be capable of selectively inhibiting viral functions without damaging the host, making the development of such drugs very difficult. Furthermore an ideal drug would reduce disease symptoms without modifying the viral infection so much as to prevent an immune response in the host. There is a need for antiviral drugs active against viruses, for which vaccines are not available or are not highly effective.

Anti-viral Development

ادرسوها بیست ما فیها
انفی مع

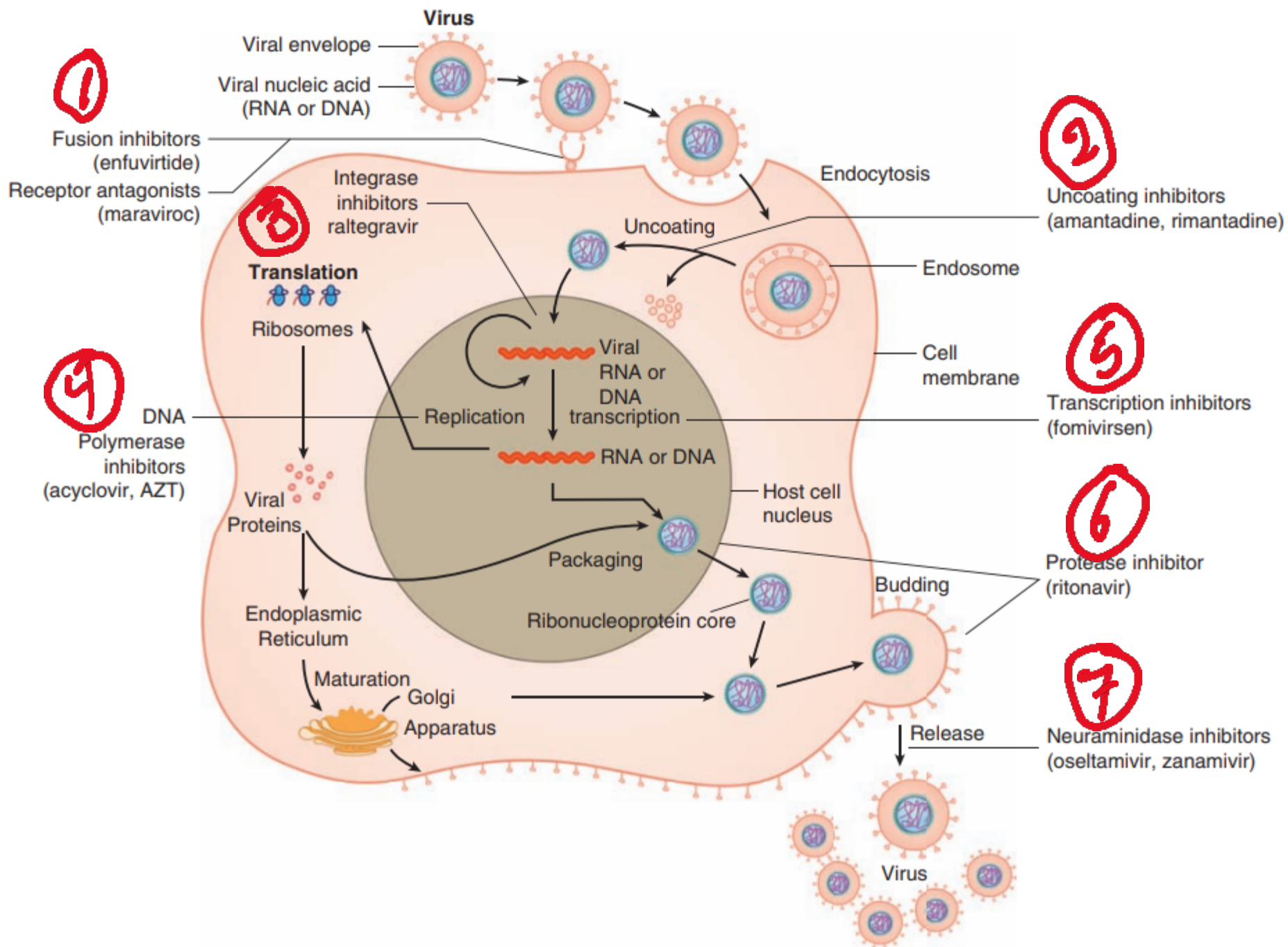
- Viruses are now becoming better understood and several viral genomes have been properly mapped. Scientists are now looking for the best drug targets
- The main point of interest is any viral protein that the host organism does not normally produce
- Once these viral proteins are identified they are tested using a large-scale screening process to test for effectiveness

← بیست های مهمه

Anti-viral Targets

سهمین

- There are several known methods that the makers of Antiviral drugs are looking at, including:
 1. Inhibitors of Attachment
 2. Inhibitors of Cell Penetration and Uncoating
 3. Neuraminidase Inhibitors
 4. Protease Inhibitors
 5. Inhibitors of Nucleic Acid Synthesis
 6. Nucleotide Analogs
 7. Stopping the release of the mature viruses from the host cell



1. Oseltamivir (Tamiflu)

- → Prevents the ^{يمنع} mature viruses from leaving the cell
- → It is a neuraminidase inhibitor, it works on both influenza A and B
← قتيلا لانزيم neuraminidase
- → Neuraminidase is an enzyme found on the virus which cleaves sialic acid from cell membrane, leading to a more effective release of viruses
- → Used to battle avian flu and influenza



فعال لمكافحة انفلونزا الطيور
والانفلونزا الموسمية

2. Acyclovir (Zovirax)

استخدم بشكل اساسي
A widely used antiviral with main implications in the treatment of herpes
Seen as a "new age" in antiviral therapy, Gertrude Elion, its creator, was given the Nobel prize for medicine in 1988

It is a nucleoside analogue and prevents viral replication in infected cells
Inhibits viral DNA polymerase and terminates viral DNA chain growth



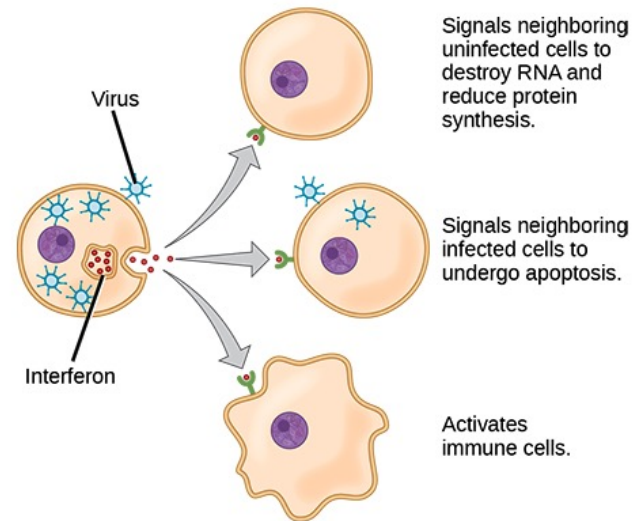
3. Interferons

α and β interferons are produced by all the cells in response to viral infections

γ interferons are produced only by T lymphocyte and NK cells in response to cytokines

The action of interferons leads to an inhibition of translation

Pegylated interferon- α (Peg-IF α) is given for 6 to 12 months to treat chronic hepatitis C disease



Viral Vaccines

General Principles

Types:

1. Killed-Virus Vaccines
 2. ضعيفة / الموهنة Attenuated Live-Virus Vaccines
 3. لقاحات جينية Genetic vaccines
- Proper Use of Vaccines
 - Vaccine development and future direction

