General Consideration

Season is from late November to late March

• Acute viral respiratory illnesses are among the most common of human diseases, accounting for one-half or more of all acute illnesses.
• Influenza is an acute respiratory illness caused by infection with influenza viruses.
• One of the most important Emerging and Reemerging infectious diseases.
• The illness affects the upper and/or lower respiratory tract and is often accompanied by systemic signs and symptoms such as fever, headache, myalgia, and weakness.
• Outbreaks of illness of variable extent and severity occur nearly every year. Such outbreaks result in significant morbidity rates in the general population and in increased mortality rates among certain high-risk patients, mainly as a result of pulmonary complications.
Influenza Mainly affects upper respiratory tract, rarely involve lungs (The respiratory tract) exception to this is H5N1 (avian influenza) affects mainly lower respiratory tract and it's a severe disease, not effectively transmitted between humans

**Myxoviruses**

- Orthomyxo viruses
  - Smaller
  - Segmented RNA genome
  - Liable to Agic variation

- Paramyxoviruses
  - Larger
  - Single piece of RNA
  - Not liable to Agic variation

**Influenza viruses**

- Parainfluenza
- Mumps virus
- Measles virus
- Respiratory syncytial virus

**Myxo** = affinity to mucin
Characteristics of Influenza Virus

- Pleomorphic
- Types (A, B, C, D)
- Diameter 80 - 120 nm
- Pleomorphic, spherical, filamentous particles
- Single-stranded RNA
- Segmented genome, 8 segments in A and B
- Hemagglutinin and Neuraminidase on surface of the virion

Affect Humans

Typing based on:
1) Nucleocapsid (ribonucleoprotein)
2) Matrix protein
Influenza Structure

- 8 segments of single-stranded RNA
- Segments combine with nucleoprotein (NP) to form the ribonucleoprotein core
- M1 matrix protein surrounds the core
- Lipid coat surrounds the matrix
- Embedded in the lipid membrane are 2 important viral proteins: hemaglutinin (HA) and neuraminidase (NA)
- RNA segments + nucleocapsid = a nucleocapsid with helical symmetry
Influenza A Virus Structure

- Lipid Bilayer
- NA (Neuraminidase)
- HA (Hemagglutinin)
- M₂ (Ion channel)
- M₁ (Matrix protein)
- Infected cell protein
- NS₁
- PB₁, PB₂, PA (Transcriptase complex)
- NP (Nucleocapsid)
Antigenic structure & Classification

I. Type Specific Ag (core Ag):
- Three serotypes: A, B & C according to internal structure ptns (nucleocapsid & matrix). These ptns don’t cross react.

II. Strain (subtype) specific Ag:
- Two surface glycoproteins, HA & NA are used to subtype the virus.
- Influenza strains are named after their types of HA & NA surface ptns e.g. H1N1.

Circulating Now between Humans H1N1 H2N3
Haemagglutinin (H)
Binds to host cell surface receptor

Neuraminidase (N)
Cleaves neuraminic acid to release virus progeny from infected cells

Works on neuraminic acid a component of the mucus
Cleavage and spreading of the infection

Has the ability to agglutinate RBCs of different species
Initiation of infection, binds to sialic acid found in mucus secreting cells in the respiratory tract
Fusion with Host Membrane

The flu virus binds onto sugars on the surfaces of epithelial cells such as nose, throat, and lungs of mammals and intestines of birds.
Influenza virus Replication cycle

1. Virus (structure shown in a cutaway view) adsorbs to a respiratory epithelial cell by hemagglutinin spikes and fuses with the membrane.

2. The virus is endocytosed into a vacuole and uncoated to release its 8 nucleocapsid segments into the cytoplasm.

3. The nucleocapsids are transported into the nucleus. The (-) sense RNA strand (black) is transcribed into a (+) sense strand (red) that will be translated into viral proteins that make up the capsid and spikes.

4. (+) Sense RNA is used to synthesize glycoprotein spikes inserted into the host membrane.

5. Envelope forming.

6. Release of mature virus occurs when viral parts gather at the cell membrane and are budded off with an envelope containing spikes.

6. The (-) sense RNA strands are used to synthesize new (-) sense RNA strands. These are assembled into nucleocapsids and transported out of the nucleus to the cell membrane.

Final assembly cut the host cell membrane budding.
Types of Influenza virus

I- Type A virus: Most common & most severe
- Infects humans as well as animals
- Undergoes continuous Antigenic variations
- Many animal species have their own influenza A virus
- Pigs & birds are the reservoirs playing a role in occurrence of influenza epidemics
Types of Influenza virus

II- Type B virus: Minor outbreaks/epidemics-No pandemics
- Causes milder disease
- Infects human only
- Only undergo antigenic drift
- Not known to undergo antigenic shift

III- Type C virus: 7 segments (lacks Neuraminidase)
- Agntigenically stable
- Known to cause only minor respiratory disease; probably not involved in epidemics
Hemagglutinin

- **Structure**: trimer of “lollipops” with fibrous stem anchored in the membrane and globular protein sphere containing the sialic acid receptor site
- **Function**: Sialic acid receptor sites bind to host cell’s glycoproteins allowing for infection to occur

Of identical dimers (HA1-HA2)

Mediate fusion (initiation of infection)
**Neuraminidase**

- **Structure**: Box-shaped tetramer with stalk that anchors it to the cellular membrane
- **Function**: Cleaves off sialic acid molecules from the surface of cells thereby preventing infected cells from “recapturing” budding virus molecules.
Surface Antigens

Haemagglutinin
- Binds to host cell surface receptor
- The target of neutralizing Abs
- Haemagglutinates RBCs from various animal species

Neuraminidase
- Cleaves neuraminic acid to release virus progeny from infected cells
- Degrades the protective layer of mucin in the respiratory tract
- Plays a minimal role in immunity to influenza
Antigenic Variation

- Ag Variations occurs only in influenza A because it has a wide host range, giving influenza A the opportunity for a major reorganization of its genome & hence its surface Ags.

- Pigs are susceptible to avian, human & swine influenza viruses and they potentially may be infected with influenza viruses from different species. If this happens, it is possible for the genes of these viruses to mix and create a new virus.
Antigenic Variation

1- antigenic shift

- It is the process in which the genetic segment encoding for envelope glycoproteins (HA&NA) is replaced by another one from a different strain through genetic reassortment causing replacement of the original HA or NA by a new one.

- **Major** change, new subtype, May result in pandemic.

- **Genetic reassortment:** the exchange of genetic material between viruses inside a host cell.
Duck Influenza Virus

Human Influenza Virus

Human Influenza Virus with Duck HA

Immune system Has no recall for Duck HA

Antigenic Shift event

This is responsible for appearance of completely new strains to which no one is immune & not covered by annual vaccinations
Example of antigenic shift can happen inside humans.

H2N2 virus circulated in 1957-1967
H3N2 virus appeared in 1968 and completely replaced H2N2 virus
2) Antigenic Drift

• Minor change, same subtype

• Caused by point mutations in gene, minor change of an amino acid sequence of HA or NA. Occurs in influenza A & B produce new strains are referred to as antigenic shifts
  • May result in epidemic

• Example of antigenic drift
  • In 2003-2004, A/Fujian/411/2002-like (H3N2) virus was dominant
  • A/California/7/2004 (H3N2) began to circulate and became the dominant virus in 2005
Antigenic drift

Decreasing serologic relatedness

HA or NA

Antigenic shift

HA or NA

NP

Years

One year
Classification and Nomenclature

Origin is from Avian influenza except HA(17-18)+NA(10+11) from bats

- The standard nomenclature system for influenza virus isolates includes the following information: type, host of origin, geographic origin, strain number, and year of isolation. Antigenic descriptions of the HA and the NA are given in parentheses for type A.

- The host of origin is not indicated for human isolates, such as A/Hong Kong/03/68(H3N2), but it is indicated for others, such as A/swine/Iowa/15/30(H1N1).

- So far, 18 subtypes of HA (H1–H15) and eleven subtypes of NA (N1–N9), in many different combinations, have been recovered from birds, animals, or humans. Four HA (H1–H3, H5) and two NA (N1, N2) subtypes have been recovered from humans.
Pathogenesis

Influenza

↓

Sudden+ common complaints
(fever/athralgia/chills...)
+ minor respiratory symptoms
(cough/ runny nose/...)

Common cold

↓

Gradual, patient complains of respiratory problems and no systemic signs (fever/chills/...)

Epithelial cells of respiratory tract

Viral NA degrades the protective mucin layer
Allowing the virus to enter the cells

Replication inside the cells
Cilia damage
Epithelial desquamation

The infection is limited to the respiratory tract → Might cause Metaplasia

There are proteases there essential for HA to be active

Despite systemic symptoms, no viremia

Those symptoms are due to cytokines production

Recovery takes a few months
Mode of transmission

• Highly contagious disease with person to person transmission
• Three modes of transmission

Droplet

Air-Borne

Contact

Direct
Indirect

Short Incubation Period 1-3 days
Duration of shedding

- In otherwise healthy adults with influenza infection, viral shedding can be detected 24 to 48 hours before illness onset, but is generally at much lower titers than during the symptomatic period.
- In a review of 56 studies of 1280 healthy adults who were experimentally challenged with influenza virus, shedding of influenza virus increased sharply one-half to one day following exposure, peaked on the second day, and then rapidly declined.
- The average duration of shedding was 4.8 days. Shedding ceased after six or seven days in most studies but occurred for up to 10 days in some. Studies of natural infection in healthy adults have shown similar results.
Clinical Findings

- High fever
- Non-productive as well as productive cough
- Shortness of breath
- Dyspnoea
- Hypoxia
- Evidence of lower respiratory tract disease with opacities, consolidation, and infiltrates noted on chest imaging
- More severe infections (i.e. pneumonia) are sometimes associated with Influenza because of the increased susceptibility to other infections as a result of a damaged airway
Pulmonary complications

- Primary influenza pneumonia
  - Primary influenza pneumonia occurs when influenza virus infection directly involves the lung, typically producing a severe pneumonia.
  - Clinical suspicion for primary influenza pneumonia should be raised when symptoms persist and increase instead of resolving in a patient with acute influenza.
  - High fever, dyspnea, and even progression to cyanosis can be seen.

- Secondary bacterial pneumonia (*Streptococcus pneumoniae, Staphylococcus aureus, and Haemophilus influenzae*).
  Most common cause of superimposed infection pneumonia post influenza is staph aureus

- Mixed viral and bacterial pneumonia
Complications of pneumonia

- Septic shock,
- Respiratory failure,
- Acute respiratory distress syndrome,
- Refractory hypoxemia,
- Acute renal dysfunction,
- Multiple organ dysfunction,
- Rhabdomyolysis,
- Encephalopathy (Reye syndrome)
- Bacterial and fungal infections such as ventilator-associated pneumonia and blood-stream infection sometimes by multi-drug resistant bacteria

Dont give children who have Flu like syndrome (salicylic drugs like aspirin)
<table>
<thead>
<tr>
<th>Groups at high risk for influenza complication</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ <strong>Children &lt;2 years</strong>*</td>
</tr>
<tr>
<td>▪ <strong>Adults ≥65 years of age</strong></td>
</tr>
<tr>
<td>▪ Persons with chronic pulmonary (including asthma), cardiovascular (except hypertension), renal, hepatic, hematologic (including sickle cell disease), metabolic (including diabetes mellitus), neurologic, neuromuscular, and neurodevelopmental disorders (including disorders of the brain, spinal cord, peripheral nerve and muscle such as cerebral palsy, epilepsy, stroke, intellectual disability [mental retardation], moderate to severe developmental delay, muscular dystrophy, or spinal cord injury)</td>
</tr>
<tr>
<td>▪ Immunosuppression (including immunosuppression caused by medications or by human immunodeficiency virus)</td>
</tr>
<tr>
<td>▪ <strong>Women who are pregnant or postpartum (within 2 weeks after delivery)</strong></td>
</tr>
<tr>
<td>▪ <strong>Children &lt;19 years of age and receiving long-term aspirin therapy</strong></td>
</tr>
<tr>
<td>▪ <strong>Native Americans and Alaskan Natives</strong></td>
</tr>
<tr>
<td>▪ <strong>Morbidly obese (body mass index [BMI] ≥40 for adults or BMI &gt;2.33 standard deviations above the mean for children)</strong></td>
</tr>
<tr>
<td>▪ <strong>Residents of nursing homes and other chronic care facilities</strong></td>
</tr>
</tbody>
</table>
Laboratory Diagnosis

A. Polymerase Chain Reaction

- Rapid tests based on detection of influenza RNA in clinical specimens using reverse transcription polymerase chain reaction (RT-PCR) are preferred for diagnosis of influenza. RT-PCR is rapid (<1 day), sensitive, and specific.

B. Isolation and Identification of Virus

- Viral culture procedures take 3–10 days. Classically, embryonated eggs and primary monkey kidney cells have been the isolation methods of choice for influenza viruses, although some continuous cell lines may be used. In the presence of trypsin, which cleaves and activates the HA so that replicating virus will spread throughout the culture. Cell cultures can be tested for the presence of virus by hemadsorption 3–5 days after inoculation, or the culture fluid can be examined for virus after 5–7 days by hemagglutination.
C. Serology

• Antibodies to several viral proteins (hemagglutinin, neuraminidase, nucleoprotein, and matrix) are produced during infection with influenza virus. The immune response against the HA glycoprotein is associated with resistance to infection.

• Routine serodiagnostic tests in use are based on haemagglutination inhibition (HI) and enzyme-linked immunosorbent assay. Paired acute and convalescent sera are necessary because normal individuals usually have influenza antibodies. A fourfold or greater increase in titer must occur to indicate influenza infection. Human sera often contain nonspecific mucoprotein inhibitors that must be destroyed before testing by HI.
# Hemagglutinin Subtypes of Influenza A Virus

Seasonal flu now called (H1N1 - H3N2)

<table>
<thead>
<tr>
<th>Subtype</th>
<th>Human</th>
<th>Swine</th>
<th>Horse</th>
<th>Bird</th>
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</tr>
</tbody>
</table>

15 subtypes in total.

H5N1 + H7N9 most dangerous.

LD from Avian.

Not efficient in Humans.

Origin of all influenza viruses.
## History: Known Flu Pandemics

<table>
<thead>
<tr>
<th>Name of pandemic</th>
<th>Date</th>
<th>Deaths</th>
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</thead>
<tbody>
<tr>
<td>Spanish Flu</td>
<td>1918-1920</td>
<td>40 - 100 million</td>
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<tr>
<td>Asian Flu</td>
<td>1957-1958</td>
<td>1 - 1.5 million</td>
</tr>
<tr>
<td>Hong Kong Flu</td>
<td>1968-1969</td>
<td>0.75 - 1 million</td>
</tr>
<tr>
<td>Swine Flu</td>
<td>2009-2010</td>
<td>0.15 - 0.6 million</td>
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</table>
Treatment and Prevention
Influenza Vaccines

- Whole virus vaccines: inactivated forms of virus with the predicted HA, are grown in embryonated eggs
- Subunit vaccine: uses both HA and NA subunits extracted from recombinant virus forms
- Split-virus vaccines: purified HA (lessens the side-effects)
- Recommended for health care workers, elderly/people in nursing homes, asthmatics, chronic lung disease patients, some pregnant women, and anyone who is susceptible to infection
Influenza Vaccines

- **Inactivated subunit (TIV)**
  - Intramuscular
  - Trivalent
  - Annual

- **Live attenuated vaccine (LAIV)**
  - Intranasal
  - Trivalent
  - Annual

Flu shot = killed
Trivalent (in Jordan) = 2 subtypes of A and 1 Subtype of B
Quadrivalent vaccine (in USA) = 2 subtypes of A and 2 Subtypes of B
Contraindicated in pregnancy

FLUMIST also has a quadriivalent version.
WHO recommends annual vaccination for (in order of priority)

- Nursing-home residents (the elderly or disabled)
- Elderly individuals
- People with chronic medical conditions
- Other groups such as pregnant women, health care workers, those with essential functions in society, as well as children from ages six months to five years

Medical personnel
Antiviral Treatment Recommendations

• Treatment with oseltamivir (Tamiflu) or zanamivir is recommended for:
  • All patients requiring hospitalization
  • Patients at increased risk of complications
    • Children 0-4 years
    • Pregnant women
    • Persons with immune suppression, chronic pulmonary (including asthma), cardiovascular (except hypertension), renal, hepatic, hematological (including sickle cell disease), neurologic, neuromuscular, or metabolic disorders (including diabetes mellitus) or > 65 years

• Early treatment is the key
  • Clinicians should not wait for confirmatory tests to treat
  • Postexposure prophylaxis should generally not be used
    • Consider for high-risk person with close unprotected exposure
    • Do not use if more than 48 hours after exposure

Specially for old people because you can't vaccinate them
Healthy Habits

• When Healthy:
  • Avoid close contact with those who are sick
  • Wash your hands often
  • Avoid touching your eyes, nose and mouth to decrease the spread of germs

• When Ill:
  • Cover your mouth and nose with a tissue (or upper sleeve) when you sneeze or cough
  • Stay home from work or school when you are sick
Key facts

- Influenza is an acute viral infection that spreads easily from person to person.
- Influenza circulates worldwide and can affect anybody in any age group.
- Influenza causes annual epidemics that peak during winter in temperate regions.
- Influenza is a serious public health problem that causes severe illnesses and deaths for higher risk populations.
- An epidemic can take an economic toll through lost workforce productivity, and strain health services.
- Vaccination is the most effective way to prevent infection.
Avian Influenza

• A contagious viral infection and/or disease of many avian species including poultry, wild and exotic birds, ratites, shore birds and migratory waterfowl.

• The highly pathogenic form of the disease is characterized by severe depression, decrease in egg production, high mortality, edema, hemorrhage, and frank necrosis.

• All H5 and H7 infections are reportable to the World Organization for Animal Health (OIE).
Where does AI virus come from?

- All known subtypes of influenza A viruses circulate among wild birds, especially migratory waterfowl (e.g. ducks and geese) which are considered natural reservoirs for influenza A viruses.

- Domestic poultry like chickens and turkeys are not natural reservoirs for AI virus and usually develop clinical disease when infected with AI virus.
Mode of transmission contact, droplet (not sure/no studies)
Main route is their feces (the virus stays alive for 3 months)
How does AI virus spread?

• Exposure of poultry to migratory waterfowl
• Exposure of commercial poultry to AI-infected backyard, game bird, or hobby flocks
• Contact with AI-infected live bird markets
• Bird to bird contact (through feces)
• Aerosol droplets
• Manure, equipment, vehicles, egg flats, crates, contaminated shoes and clothing
• Wildlife vectors/scavengers
What are the types of Avian Influenza in domestic poultry?

Both of them can affect humans

- **Low pathogenic avian influenza (LPAI)**
  - Mild or no clinical signs
  - Low to moderate mortality
  - However, the low pathogenic H5 and H7 strains are capable of mutating under field conditions into highly pathogenic strains

- **Highly pathogenic avian influenza (HPAI)**
  - Sudden onset
  - Severe clinical signs
  - High mortality

No effective transmission cycle between humans

(concorded with birds)

\[ H_{6N14} \text{ and } H_{7N9} \] from birds
H1N1/H5N1

H1N1

Upper respiratory tract

Easily spread
Rarely fatal

H5N1

Lower respiratory tract (lung) so its more severe

Spreads slowly
Often fatal

from birds (most common in Avian)
Criteria for pandemic
1) New subtype (Antigenic shift)
2) Efficient human to human transmission
3) It produces a severe disease

No one can predict when the pandemic is going to happen or what it will be
WHAT IS SWINE FLU?

Swine Influenza (swine flu) is a respiratory disease of pigs caused by type A influenza viruses (H1N1 subtype) that causes regular outbreaks in pigs.

People do not normally get swine flu, but human infections can and do happen.

Swine flu viruses have been reported to spread from person-to-person, but in the past, this transmission was limited and not sustained beyond three people.
The End