

Respiratory infections

Respiratory tract anatomy

THE UPPER RESPIRATORY TRACT

Nose

Sinuses

Mouth

Pharynx

Larynx

THE LOWER RESPIRATORY TRACT

Trachea

Bronchi

Bronchioles

Lungs



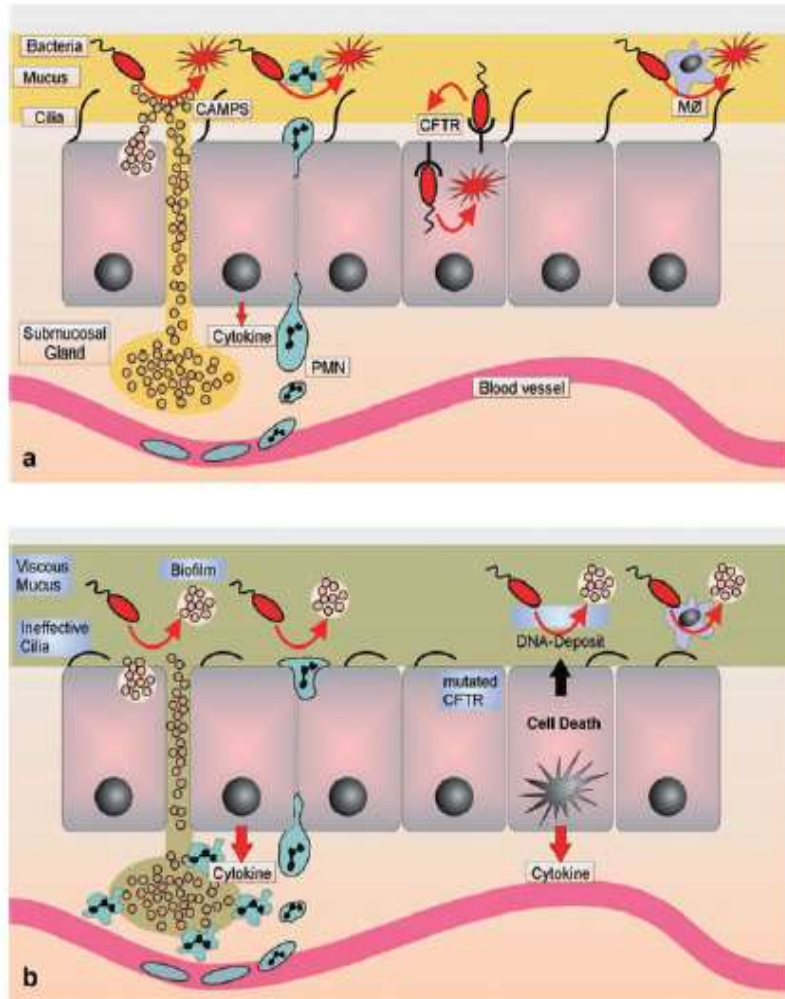
RELEVANCE, ETIOLOGY, EPIDEMIOLOGY

- RELEVANCE – highest morbidity
- seniors, chronic infections – frequent hospitalization, significant mortality
- ETIOLOGY – viruses, bacteria, fungi, parasites
- EPIDEMIOLOGY – etiology, symptomatology and significance are age depended
- transmission – most frequently by droplets or direct contact
- seasonal epidemics (cold periods)
- repeatedly occur – many pathogens and serotypes (short immunity answer)
- transmission is potentiated – high mobility and people concentration, stress factors, emerging and re-emerging pathogens, e.g. coronavirus (SARS 2002-2003) or antigenic shift of flu virus, SARS-CoV-2 emerged in 2019 (China)

PHYSIOLOGY AND PATHOPHYSIOLOGY

- Upper respiratory tract (URT) - large number of commensal bacteria
- Lower respiratory tract (LRT), middle ear, paranasal cavity – nearly sterile, if inflammation – large number of polymorphonuclear cells (PMN) with majority of one bacterial morphology
- defence mechanisms – ciliary epithelia of URT and LRT, lysozyme, mucosal IgA
- bacterial infection – non-specific (innate) mechanisms – e.g. C-reactive protein (CRP), complement, PMN
- viral infection – interferon (blocking of NA metabolism in neighbour cells – inhibition of virus replication)
- NALT – in URT, *nasopharyngeal-associated lymphoid tissue*
- BALT – in LRT, *bronchus -associated lymphoid tissue*
- Lymphatic tissue (part of Waldeyer's ring) – without capsule, directly penetrating epithelia (lymphoepithelium – dendritic cells, Langerhans cells, intraepithelial location – B lymphocytes, plasmatic cells) – antigen monitoring

Innate immunity and microbes elimination



Bacterial killing mechanism of innate immunity in the respiratory tract of healthy individuals and cystic fibrosis (CF) individuals.

A. In healthy individuals, bacteria, entering the mucus layer overlaying the respiratory epithelium, are effectively removed from the airways by a functional mucociliary clearance system and killed by CAMPs derived from submucosal glands or epithelial cells, by functional neutrophils or macrophages or within epithelial cells after uptake via functional CFTR. PMN, polymorphonuclear leukocytes.

B. In individuals with CF defective CFTR leads to a highly viscous mucus layer which impairs mucociliary clearance, the migration of CAMPs, neutrophils and macrophages towards the bacterial targets, the uptake of bacteria by CFTR itself, induces a pH shift in epithelial lysosomes which due to ceramide accumulation results in DNA deposits which in turn serve as adhesion matrices for bacteria. A similar pH shift in the phagolysosomes of macrophages impairs bacterial killing. Ceramide accumulation also triggers cytokine release which induces further neutrophil influx.

Ref. Goring, Gulbins, Cellular Microbiology (2009) 11(2), 208–216

Viral agents & respiratory infections (1)

Orthomyxoviruses (ss RNA)

- **influenza viruses** – ½ acute respiratory illness, major determinant of morbidity and mortality caused by respiratory infections (**Dg** – deep nasopharyngeal swab/PCR, **Th** – **oseltamivir**, baloxavir, zanamivir, **prevention – vaccination**)

Paramyxoviruses – ss RNA, acute resp.inf. & pneumonia 4 mil.children yearly (WHO)

- **parainfluenza virus** - ubiquitous in all ages, nose throat – **common cold, croup-laryngotracheobronchitis, bronchiolitis, pneumonia** (**Dg** – deep nasopharyngeal swab/PCR, **Th** – symptomatic (ribavirin), **no prevention**)
- **respiratory syncytial virus** – **bronchiolitis and pneumonia** in infants under 1 year, seniors (**Dg** – deep nasopharyngeal swab/PCR, **Th** – symptomatic, **prevention – vaccination**)
- **metapneumovirus** – described 2001, wide range of disease – **from mild upper resp.inf. to severe lower resp.tract inf.**

Viral agents & respiratory infections (2)

Adenoviruses

DNA virus, icosahedral symmetry, naked

typical symptoms - cough, nasal congestion, fever, sore throat, pneumonia (**Dg** – deep nasopharyngeal swab/PCR, **Th** – symptomatic, **no vaccination**)

Coronaviruses

ssRNA, helical, enveloped, high frequency of mutations and recombinations, tropism to respiratory and GIT tract

SARS (Severe Acute Respiratory Syndrome) – acute respiratory distress requiring ventilation support (death 10%)

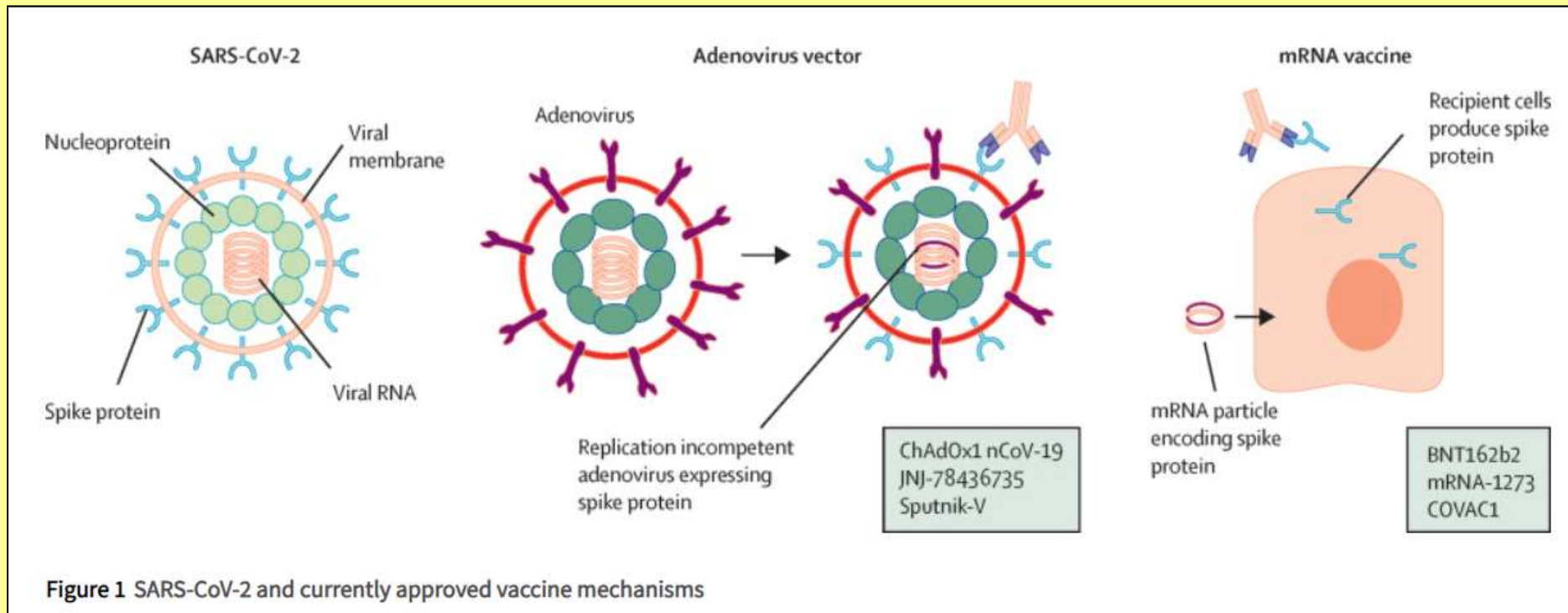
and **SARS-CoV-2 (Covid-19 disease)** (**Dg** – deep nasopharyngeal swab/PCR, **Th** – paxlovid, remdesivir, molnupiravir), **prevention – vaccination, see also next slide)**

Picornaviruses (Th-Symptomatic, prevention - no vaccine)

ssRNA, icosahedral symmetry

rhinovirus group (100 species) – **common cold viruses**

coxsackieviruses – **herpangina - severe febrile pharyngitis**



more information - <https://www.thelancet.com/journals/langas/article/PIIS2468-1253%2821%2900024-8/fulltext>

Bacterial agents & respiratory infections

- * **Staphylococci** (*S. aureus* – sinusitis, pneumonia – PVL production), Th - oxacillin (PVL+ add also clindamycin or linezolid)
- * **Streptococci** (*S. pyogenes* – tonsillitis, pharyngitis – strep throat, scarlet fever, autoimmune complication – rheumatic fever, glomerulonephritis, *S. pneumoniae* – pneumonia but also meningitis and otitis media), Th - penicillin
- * **Hemophilus** (*H. influenza b* – e.g. epiglottitis, pneumonia, non-typable – e.g. sinusitis, chronic bronchitis), Th – cefuroxime, cefotaxime
- * **Bordetella pertussis** – whooping cough, upper respiratory tract (affected also ciliated epithelium trachea, bronchi), Th- macrolides, prevention – toxoid vaccine)
- * **Corynebacterium diphtheriae** – pseudomembranes – tonsils, pharynx, larynx (Th-penicillin or erythromycin and antitoxin!, prevention – toxoid vaccine)
- * **Chlamydia pneumoniae** (Th-macrolides, tetracyclines, macrolides), **Mycoplasma pneumoniae** (Th-macrolides, tetracyclines, macrolides), **Legionella pneumophila** (Th- macrolides, fluoroquinolones, or doxycycline) – atypical pneumonia

Fungal agents & respiratory infections

- e.g. *Cryptococcus neoformans*, *Pneumocystis jirovecii* – pneumonia in AIDS patients (*more in the presentation of fungal infections*)

DIAGNOSTICS OF UPPER RESPIRATORY TRACT INFECTION (1)

RHINITIS – around 100 serotypes of rhinoviruses, **microbiological dg is not significant**, watery nasal secretion, afebrile, secondary bacterial colonisation – purulent character – without antibiotics, dif.dg – allergic rhinitis, symptomatic therapy

TONSILITIS AND PHARYNGITIS – most frequent patients in ordination, most often tonsillopharyngitis, 80% viruses, affecte – often children and young adults, children up to 3 years – mainly adenoviruses. **Viral pharyngitis** – swallowing disorders, fever, conjunctivitis, etiology – herpangina – various serotypes of Coxsackie A (isolation – pharyngeal swab and stool, RT-PCR, neutralisation antibodies, antibodies IF).

Streptococcal tonsillopharyngitis – *S. pyogenes* (beta hemolytic streptococcus of group A, pharyngeal swab – cultivation, direct detection using immunochromatographic test, complication – rheumatic fever and acute glomerulonephritis (ASO – antibodies to streptolysin O), rarely also beta-hemolytic streptococci of group C and G

Infectious mononucleosis – EBV virus, heterophilic antibodies (Paul Bunnell reaction), specific antibodies to EBV, elevated liver enzymes **Arcanobacter tonsillopharyngitis** – *Arcanobacterium haemolyticum*, young adults, culture, gram-positive rods (reverse CAMP test), **Acute retroviral syndrom** - similar to mononucleosis, sex.active, EBV or CMV can not be confirmed, **Angina Plaut-Vincent** – necrotic ulcerative tonsillitis, *Fusobacterium* and *Borrelia vincentii*

DIAGNOSTICS OF UPPER RESPIRATORY TRACT INFECTION (2)

SINUSITIS – when nasopharyngitis - always inflammation in paranasal cavities, viruses - rhinoviruses, adenoviruses, influenza, parainfluenza, bacteria – *H. influenzae*, *S. pneumoniae*, *M. catarrhalis*, *S. pyogenes*, *S. aureus*, gram-negative and anaerobic bacteria, in immunosuppressed – fungi (*Aspergillus*) secondary to infection of URT, dg – microscopy, culture of aspirate, dif.dg – allergic sinusitis, tumors, adenoid vegetation

MESOTITIS – inflammation of middle ear, complication of URT, up to 3 years – 80% of children, complication – spontaneous perforation of eardrum, mastoiditis, meningitis, pathogens - *S. pneumoniae*, non-capsulated *H. influenzae*, *S. pyogenes*, *M. catarrhalis*, *S. aureus*, dg – culture, elevated CRP

EPIGLOTTITIS – peracute, life-threatening, agents – *H. influenzae* typ b (vaccination available today), dg – leukocytoses, elevated CRP, laryngeal swab (after providing respiratory passage) and from hemoculture

LARYNGITIS AND TRACHEITIS (trachea – LRT) – 90% flu viruses, parainfluenza, adenoviruses, RS viruses, *C. diphtheriae* – diphtheria, complication - bacter. superinfection, e.g. *S. aureus*, dg – antigen in sputum, virus isolation from nasopharyngeal wash, laryng. swab, serology – seroconversion of specific titres, therapy – symptomatic (antitussives, antipyretics...)

Upper respiratory tract samples

* **Bacteriological diagnosis**

Pharyngeal and tonsillar swab
Puncture of paranasal sinuses
Nasal swab (exceptionally)

* **Virological diagnosis**

Nasopharyngeal swab (more often swab – pharynx + nose, not from tonsils), PCR

* **Serology**

Coagulated blood (yields serum)

Bacteriology – swabs, spatula, container



**Specimen collection & transport
Respiratory tract infection**

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Charles University, Prague

DIAGNOSTICS OF LOWER RESPIRATORY TRACT INFECTION (1)

TRACHEITIS (LOOK LARYNGITIS A TRACHEITIS)

BRONCHITIS – also very frequent in ordination, **Acute** – 90% viruses, adenoviruses, flu viruses parainfluenza, rhinoviruses, metapneumovirus, *M. pneumoniae*, other bacteria – rarely, occurrence – spring and winter season, pertussis (*B. pertussis*) and parapertussis (*B. parapertussis*) – catharal to necrotic inflammation, dg – clinical – auscultation, bacterial superinfection – larynx swab or sputum, diagnostic culture or serology **Chronic** – persistent cough at least 3 months during the year more than 2 years, acute exacerbation of chron. bronchitis – relapsed cough, increased expectoration of purulent sputum, dyspnoea, mainly viruses (rhinoviruses, influenza a parainfluenza, adenoviruses) and bacteria *S. pneumoniae*, non-capsulated *H. influenzae*, *M. pneumoniae*, CF patients – *P. aeruginosa*, *B. cepacia* complex, *complication* – emphysema, chronic obstructive pulmonary disease, dg – sputum culture, dispensarization by pneumologist

BRONCHIOLITIS – in infants – obstruction of bronchi and bronchioles, pathogens – often RSV, also influenza and parainfluenza viruses, *M. pneumoniae*, dg – direct detection, nasopharyngeal secretion – indirect detection – increased antibodies – ELISA, CFR

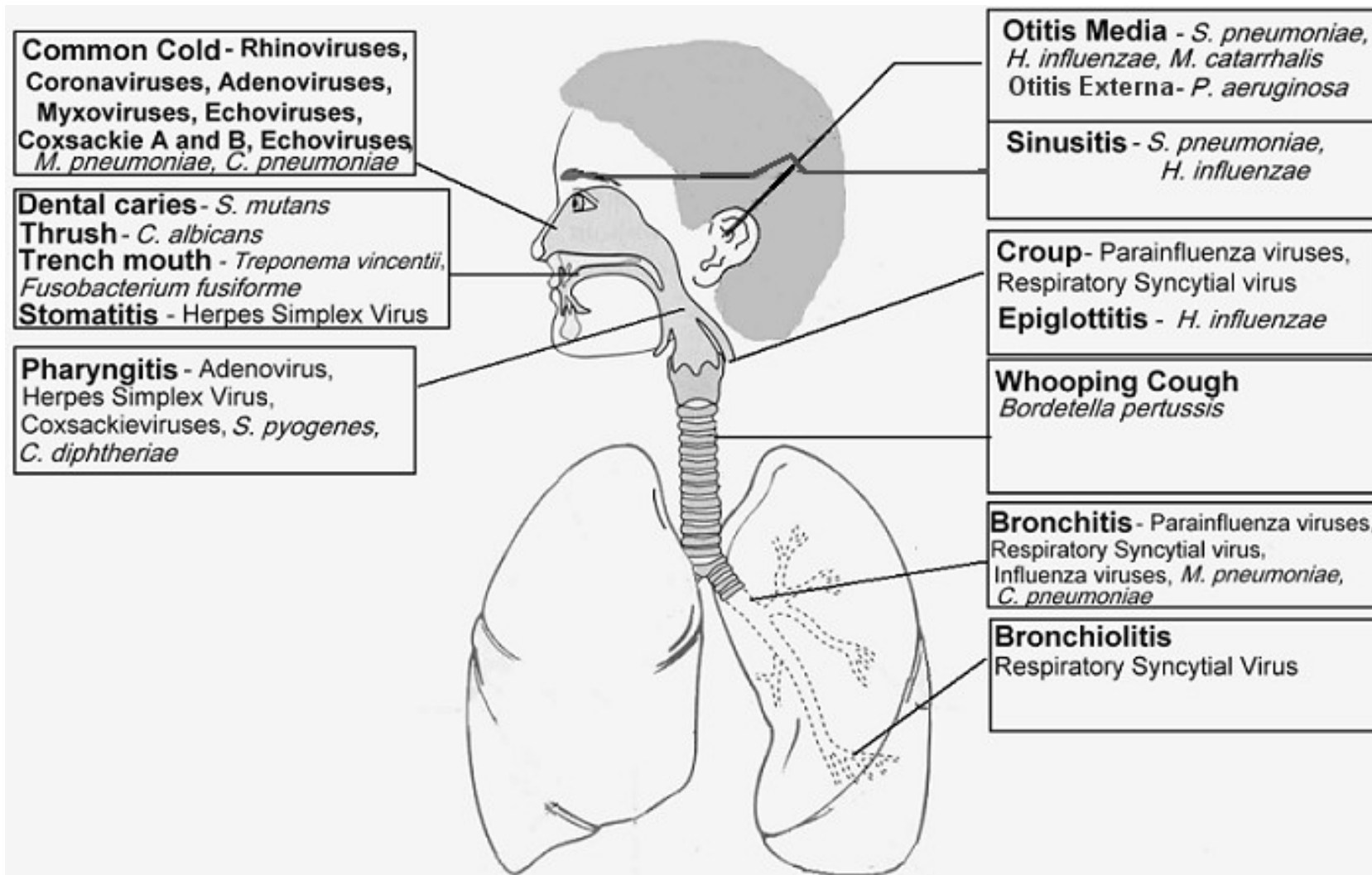
DIAGNOSTICS OF LOWER RESPIRATORY TRACT INFECTION (2)

COMMUNITY ACQUIRED PNEUMONIA – pyogenic bacteria, mycoplasma and chlamydia, viruses, legionella, *M. tuberculosis*, *Ch. pneumoniae*, *Pneumocystis jirovecii*, microb. dg – sputum culture, tracheal or aspiration culture (intubated patient), BAL, hemoculture (20% sensitivity), exudate

NOSOCOMIAL PNEUMONIA – 48h after admission, often *S. aureus*, *P. aeruginosa*, VENTILATORY – 48h after intubation, pseudomonads, acinetobacters, MRSA, microb. dg – as indicated above

ASPIRATION PNEUMONIA – polymicrobial etiology, **LUNG ABSCESS** – often anaerobes, *S. aureus*, *K. pneumoniae*, **PLEURITIS AND EMPYEMA, ATELECTASIS**

Respiratory infections



Lower respiratory tract samples

* **Sputum** – simple collected expectorated sputum. Validity of the sputum should be evaluated (number of leukocytes, epithelia, contaminating nasopharyngeal flora)

Nonvalid sputum

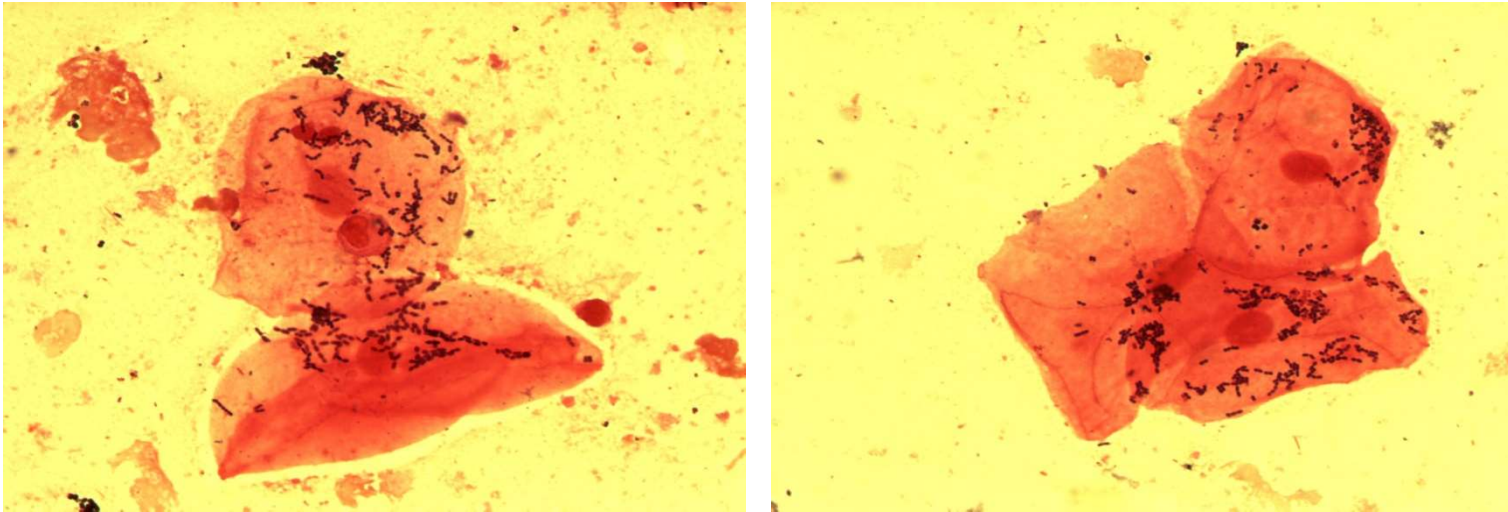


Fig. Large epithelia of upper respiratory tract (saliva in the sample) covered with adhered grampositive cocci in chains (usually comensal Viridans streptococci)(photo Melter)

Valid sputum

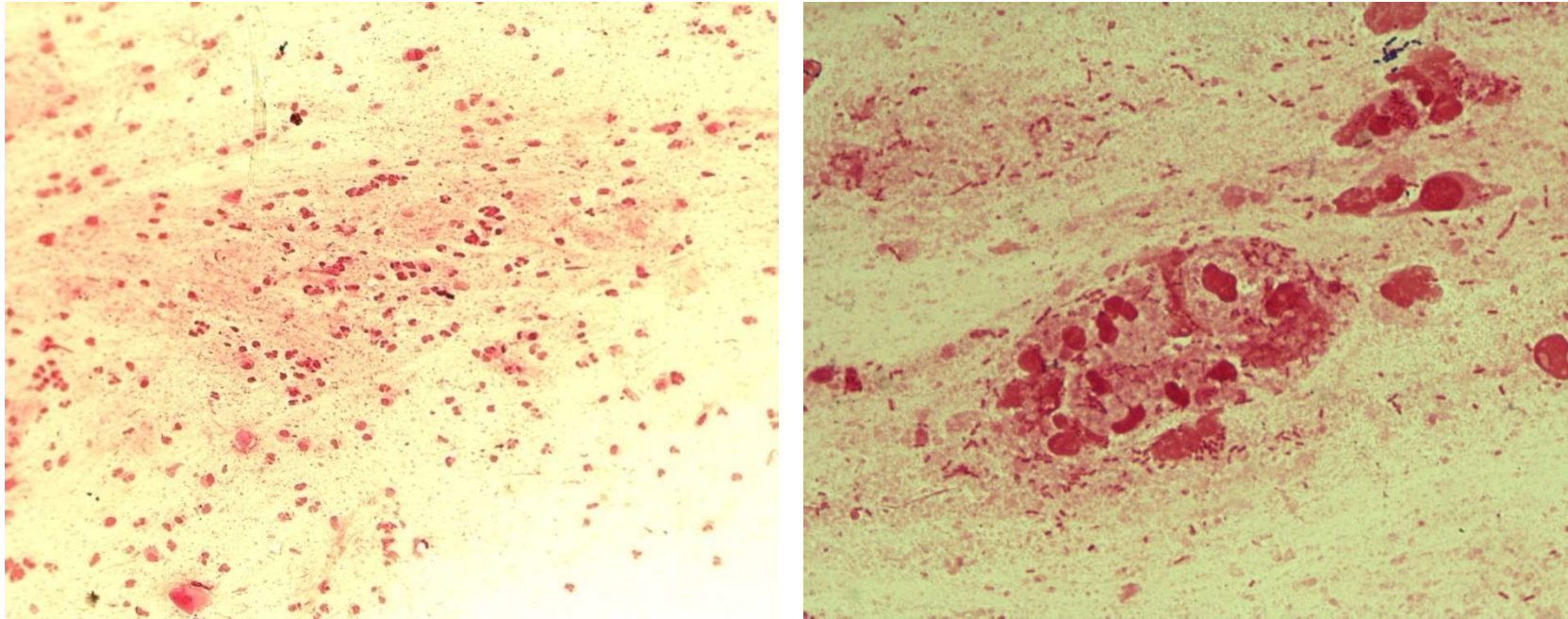
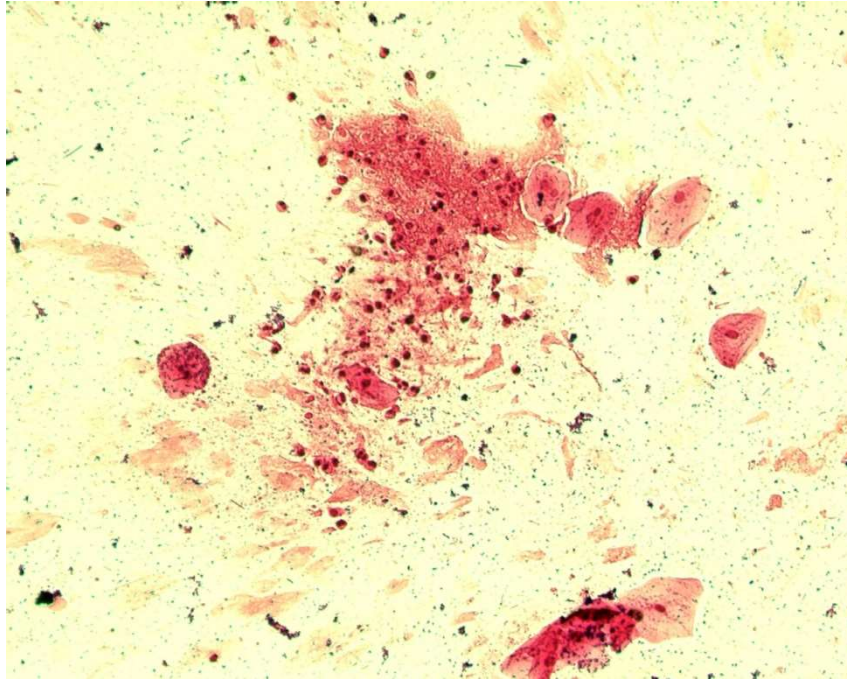
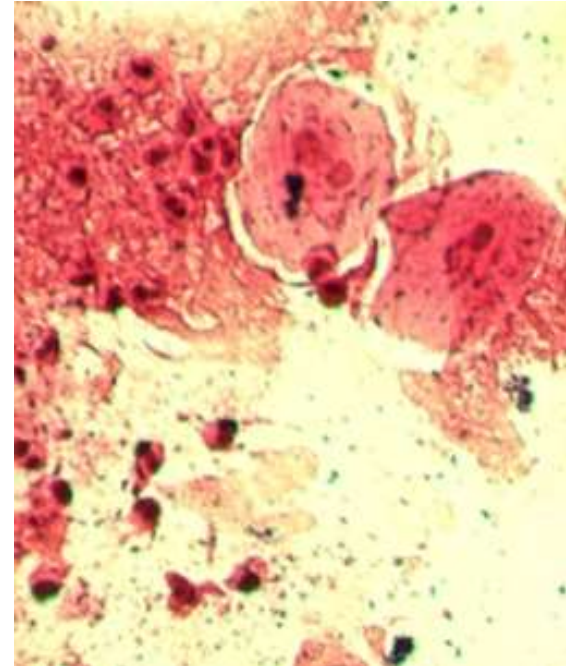


Fig. Large number of leucocytes with lower respiratory tract origin (small magnification, on the left) and uniform morphology of the bacterial cells (large magnification, gram-negative rods, on the left)(photo Melter)

Poorer valid sputum (still available for analysis)



A – small magnification



B – large magnification

Fig. In case large epithelia and leucocytes are included in the sputum is still valid for analysis (photo Melter)

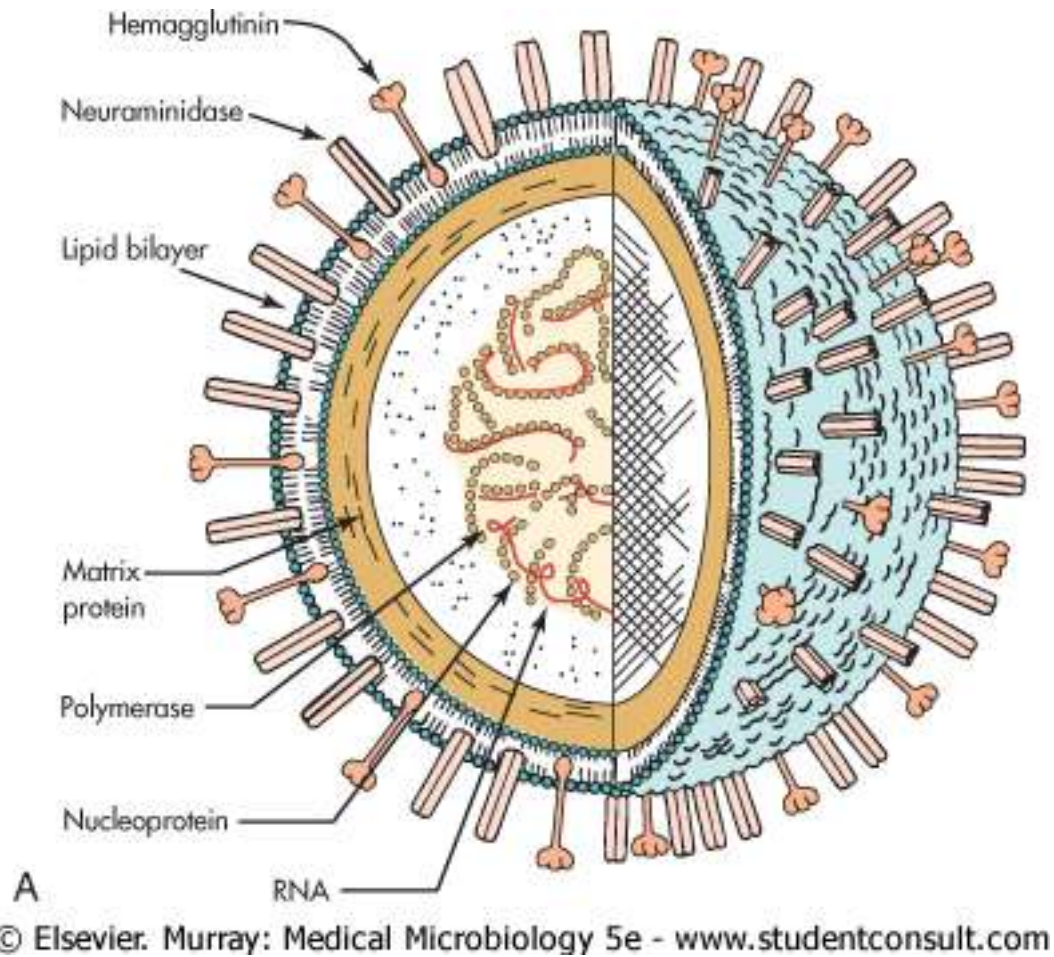
Lower respiratory tract samples

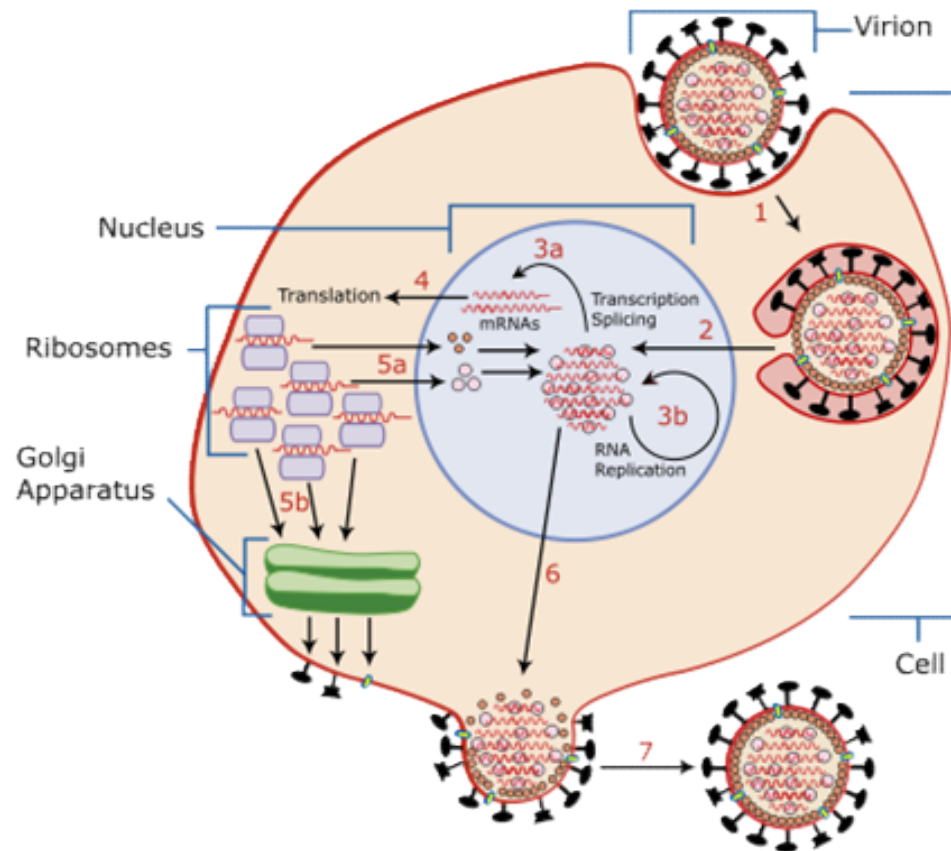
- * **Bronchoalveolar lavage** – from bronchoscopy or intubated patient
- * **Other invasive samples**
- * **Blood for hemocultivation** – in case of bacterial pneumonia
- * **Serum** – serology (detection of specific IgM and IgG antibodies)
- * **Nasopharyngeal swab for viral detection**
- * **Urine – antigen detection** (pneumonia - *S. pneumoniae*, *L. pneumophila*)

Orthomyxoviruses – structure & function

- ssRNA, spherical, 100 nm
- the envelope contains two **glycoproteins, hemagglutinin (HA) and neuraminidase (NA)**, and is internally lined by the matrix (M1) and membrane (M2) proteins.
- **genome occurs as 8 separate segments of RNA !!!**
- most of the segments code for a single protein
- eight different helical nucleocapsid segments, each of which contains a negative-sense RNA associated with the nucleoprotein (NP) and the transcriptase (RNA polymerase components: PB1, PB2, PA)

Structure of influenza A





Scheme of Influenza A virus replication (NCBI): "A virion attaches to the host cell membrane via HA and **enters the cytoplasm** by receptor-mediated endocytosis (**STEP 1**), thereby forming an endosome. A cellular **trypsin-like enzyme cleaves HA** into products HA1 and HA2 (not shown). HA2 **promotes fusion of the virus envelope and the endosome membranes**. A minor virus envelope protein M2 acts as a ion channel thereby making the inside of the virion more acidic. As a result, the major envelope protein M1 dissociates from the nucleocapsid and **vRNPs are translocated into the nucleus (STEP 2)** via interaction between NP and cellular transport machinery. In the nucleus, the viral polymerase complexes transcribe (**STEP 3a**) and **replicate (STEP 3b)** the vRNAs. Newly **synthesized mRNAs migrate to cytoplasm (STEP 4)** where they are **translated**. Posttranslational processing of HA, NA, and M2 includes **transportation via Golgi apparatus to the cell membrane (STEP 5b)**. NP, M1, NS1 (nonstructural regulatory protein - not shown) and NEP (nuclear export protein, a minor virion component - not shown) move to the nucleus (STEP 5a) where bind freshly synthesized copies of vRNAs. The newly formed **nucleocapsids migrate into the cytoplasm** in a NEP-dependent process and eventually interact via M1 with a region of the cell membrane where HA, NA and M2 have been inserted (**STEP 6**). Then the **newly synthesized virions bud from infected cell (STEP 7)**. **NA destroys the sialic acid moiety** of cellular receptors, thereby **releasing the progeny virions.**"

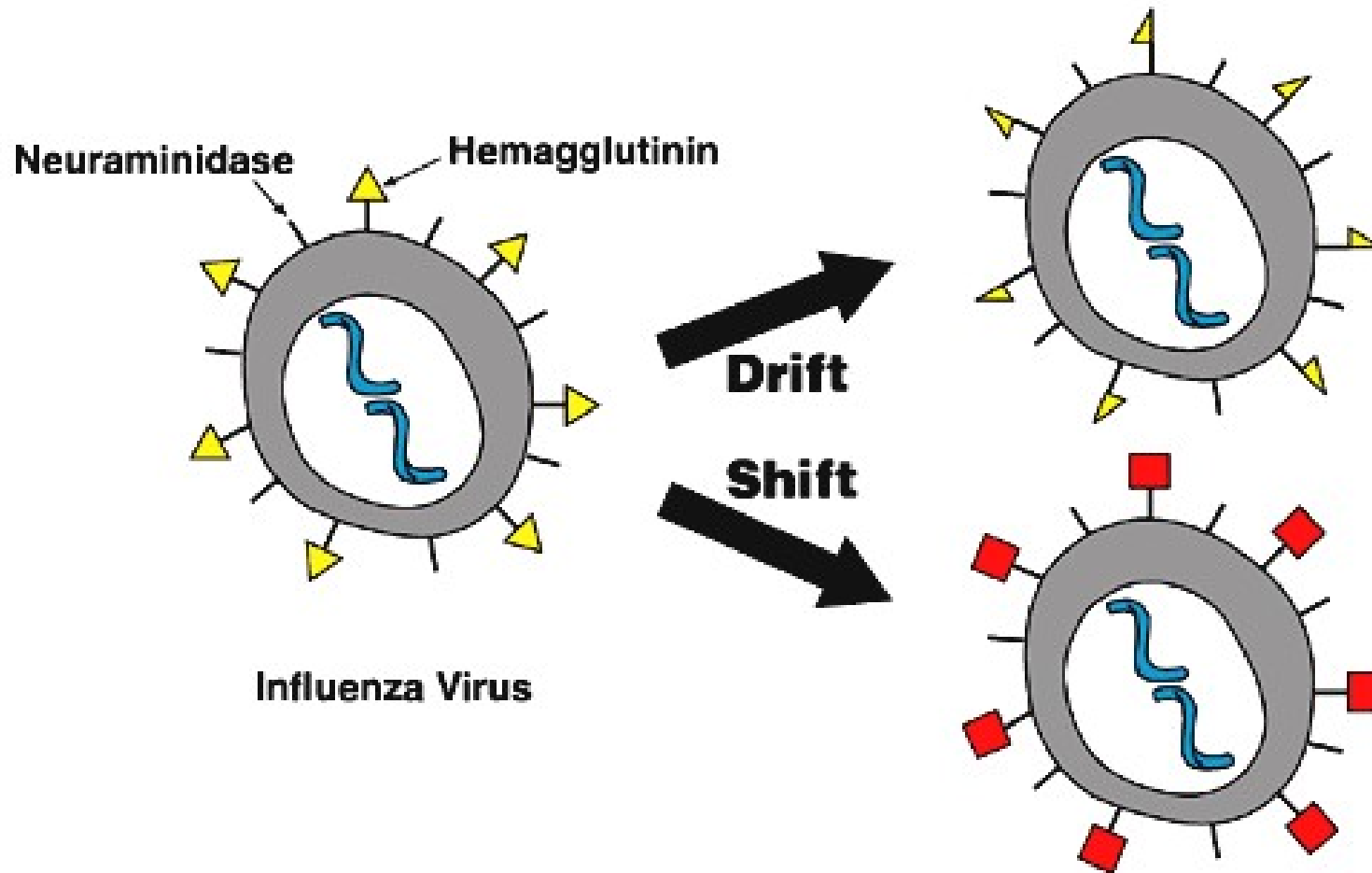
Structure & function of hemagglutinin (HA)

- glycoprotein - name from its ability to agglutinate erythrocytes
- the protein **binds virus particles to susceptible cells**
- **major Ag for neutralizing Ab - responsible for continual evolution of the HA**
- the HA spike on the virus is a trimer – cleavage by protease is necessary to be virus particle infectious (because the proteases is common in only respiratory tract – the site of infection)

Structure & function of neuraminidase (NA)

- glycoprotein, the spike is a tetramer
- its a sialidase enzyme – **releasing infected viral particules by budding from host cells**

Unusual antigenic variation of influenza A virus



Consequences of the unusual properties

- **Antigenic drift** - accumulation of point mutations resulting in amino acid changes in protein, a variant must sustain 2 or more mutations before a new, epidemiologically significant strains emerges
- **Antigenic shift** – reflect drastic changes in the sequence of viral surface protein, **genome segments are reassorted readily in doubly infected cells – result in epidemics !!!**
- **every 10-40 years when a new type of influenza A appears a PANDEMIC RESULTS**
- **1918 (H1N1, Spanish flu), 1957 (H2N2), 1968 (H3N2), 1977 reemerged H1N1 (Russian flu)**
- **Since 1977 virus A (H1N1) and H3N2 and influenza B have been in global circulation**
- **new PANDEMIC with the 2009 H1N1 type (swine/mexican flu)**

Pathogenesis and pathology

- spread: air-borne droplets, direct (hands) or undirect (surfaces) contact
- if deposited viral particles avoid removal by cough reflex and neutralization by preexisting IgA – virions spread to cells
- viral NA lowers viscosity of mucous film – bare cell surface – promoting the spread
- incubation period 1-4 d
- viral shedding 1 d before symptoms appear, duration 5 days seasonal but 10 days mexican flu
- cellular destruction – reparation 1 month (could also secondary bacterial infection – staphylococci, streptococci, *Hemophilus influenzae*)

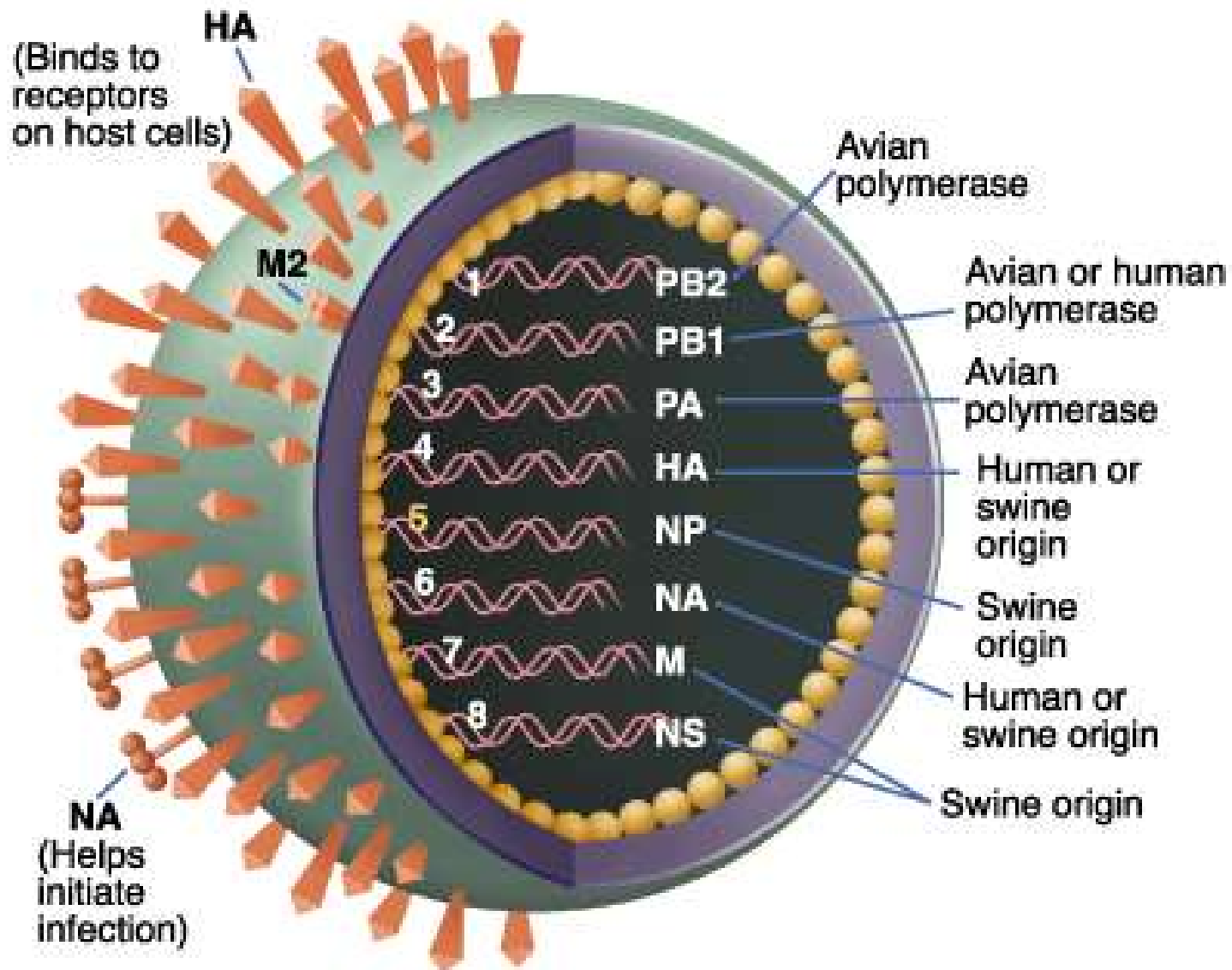
Clinical findings

- attacks mainly – upper respiratory tract
 - seasonal flu – very young, elderly people, mexican flu – mediate age
- a) uncomplicated influenza** – appear abruptly, headache, dry cough, high fever, muscular aches. Many of the classic "flu" symptoms (e.g., fever, malaise, headache, and myalgia) are associated with interferon induction.
- b) pneumonia** – elderly and debilitated and underlying chronic diseases, the pneumonia could be **viral**, secondary **bacterial** or **combination**
- c) Reye's syndrome** – **acute encephalopathy** of **children** and **adolescents**, usually between 2-16 years of age (high mortality 10-40%), possible relation between salicylate use and development of the syndrome

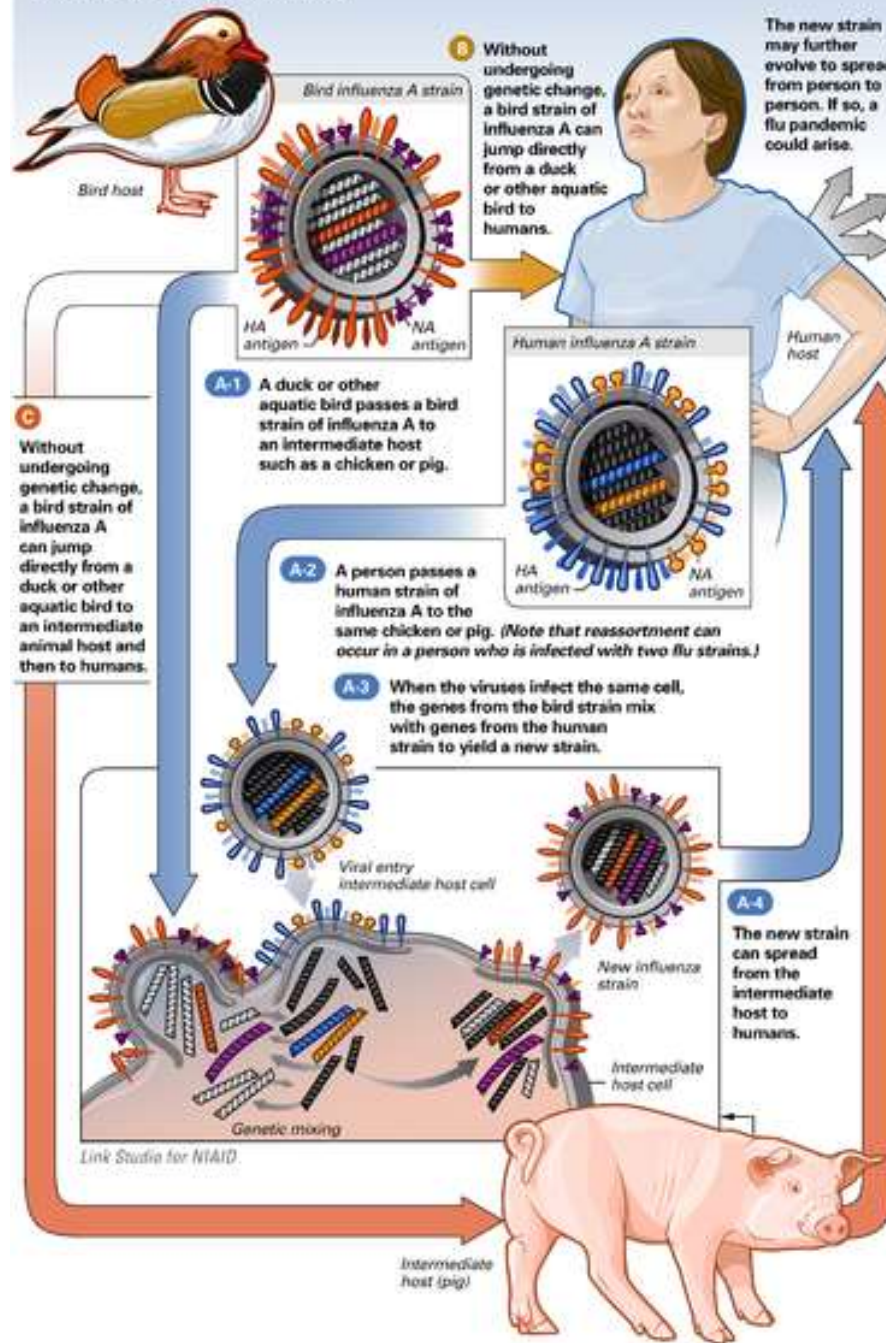
Ecology of the swine flu

- most of the genome segment appear fro swine but some also from birds and humans
- it originated/evolved in pig but probably because of more appropriate receptors the virus can cause clinical disease in human
- it is not agent of pig infection!!!

The 2009 H1N1 influenza virus A



The genetic change that enables a flu strain to jump from one animal species to another, including humans, is called "ANTIGENIC SHIFT."
Antigenic shift can happen in three ways:



Treatment

Hundreds of millions of dollars are spent on acetaminophen, antihistamines, and similar drugs to relieve the symptoms of influenza. The antiviral drug **amantadine** and its analogue **rimantadine** inhibit an uncoating step of the influenza A virus but do not affect the influenza B and C viruses. The target for their action is the M2 protein. **Zanamivir** and **oseltamivir** inhibit both influenza A and B as enzyme inhibitors of the neuraminidase. Without the neuraminidase, the hemagglutinin of the virus binds to sialic acid on other viral particles to form clumps, thereby preventing virus release. Zanamivir is inhaled, whereas oseltamivir is taken orally as a pill. These drugs are effective for prophylaxis and for treatment during the first 24 to 48 hours after the onset of influenza A illness. Treatment cannot prevent the later host-induced immunopathogenic stages of the disease.

Prevention

Natural immunization, which results from prior exposure, is **protective for long periods**. A **killed-virus vaccine** representing the "strains of the year" influenza vaccine is **available every year**. Killed whole-virus vaccines are prepared from **virus grown in embryonated** eggs and then **chemically inactivated**. Ideally the vaccine incorporates antigens of the A and B influenza strains that will be prevalent in the community during the upcoming winter. Persons with allergies to eggs should not get the vaccine. A **live vaccine** is also available for administration as a nasal spray. The trivalent vaccine consists of reassortants for the HA and NA gene segments of different influenza strains with a master donor virus that is **cold adapted** to optimum growth at 25°C. **This vaccine will elicit a more natural protection**, including cell-mediated, antibody and mucosal-secretory immunoglobulin (Ig)A antibody.