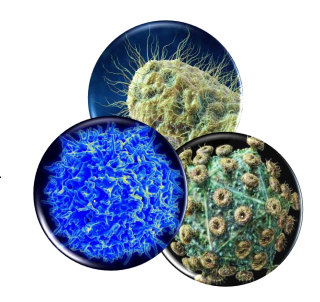
# MICROBIOLOGY, VIROLOGY, IMMUNOLOGY

Laboratory workbook

Student \_\_\_\_ group of dental faculty



MINSK BSMU 2020

#### МИНИСТЕРСТВО ЗДРАВООХРАНЕНИЯ РЕСПУБЛИКИ БЕЛАРУСЬ

БЕЛОРУССКИЙ ГОСУДАРСТВЕННЫЙ МЕДИЦИНСКИЙ УНИВЕРСИТЕТ КАФЕДРА МИКРОБИОЛОГИИ, ВИРУСОЛОГИИ, ИММУНОЛОГИИ

# МИКРОБИОЛОГИЯ, ВИРУСОЛОГИЯ, ИММУНОЛОГИЯ MICROBIOLOGY, VIROLOGY, IMMUNOLOGY

Лабораторный практикум

4-е издание



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# Glossary

aerobic - Using oxygen for growth and metabolism.

agar - A gelling agent used in bacterial growth media that allows liquids to become a gel-like solid.

anaerobic - Not requiring any oxygen for growth.

antigen - Part of an organism that is foreign to our bodies and stimulates an immune response.

**asexual organisms** - Living creatures (usually bacteria) that are neither male nor female, and therefore do not reproduce by exchanging genetic material.

biofilm - A complex community of microorganisms living together and attached to a surface.

**capsule** - A structure that surrounds or encapsulates many bacteria and may serve to protect them from harsh conditions or to assist with adherence to surfaces.

cariology - The study of cavities.

collagenase - An enzyme produced by some bacteria that breaks down the connective tissue collagen.

colonies - Masses of bacteria that arise from a single cell on solid growth media.

colonization - The act of attaching to and inhabiting a surface.

**conjugation** - The process of DNA transfer from one bacterial cell to another.

**culturing -** The act of growing bacteria in a laboratory.

cytokine - Proteins that are made by cells that alter the properties and behavior of other cells.

**cytosol** - The interior of a cell that contains the cell's inner components, or "guts".

dissemination - The process by which a pathogen is transmitted from one host to another.

**DNA fingerprint** - A characteristic sequence of nucleic acid bases (A, G, C, T) that is unique to and defines a given bacterial species.

endodontic infections - Infections that occur within the pulp of the tooth.

endoplasmic reticulum - In a eukaryotic cell, the structure on which ribosomes reside.

extracellular - The environment outside of a cell.

flagella - Flexible rope-like structures that help bacteria swim and move in different environments.

genome - The complete DNA material of an organism.

**genus** - The designation for a group of organisms highly related to each other.

gingivitis - Gum disease.

**glucan** - A general term for sugar or polysaccharide.

**Gram negative** - Bacteria that appear pink after the Gram stain procedure due to their thin peptidoglycan cell wall.

**Gram positive** - Bacteria that appear purple after the Gram stain procedure due to their thick peptidoglycan cell wall.

growth media - The food and nutrients on which bacteria grow in the laboratory.

Hemagglutination - The clumping together of red blood cells.

hemolysin - A bacterial toxin that is able to destroy red blood cells.

hemolysis - The act of lysing, or killing, a red blood cell.

**host** - The organism, usually a human, that a pathogen lives in or on.

**immuno-compromised** - A state where an individual's immune system is weakened, usually by an infection or disease.

incubate - To allow microorganisms to grow in the lab under favorable growth conditions.

**inflammation** - The process whereby immune cells and chemicals accumulate at the site of infection and result in swelling and redness.

inner membrane - The phospholipid-containing structure around a Gramnegative cell.

invasin - A protein that a pathogen uses to enter into a host cell.

**lectin** - A protein that binds to a specific type of sugar.

leukotoxin - A bacterial toxin that is able to destroy white blood cells.

**lipid A** - The innermost portion of lipopolysaccharide (LPS) that anchors it into the outer membrane of Gram-negative bacteria; composed of lipid.

**lipopolysaccharide (LPS)** - The outer part of the outer membrane of Gram-negative bacteria; composed of lipid and sugars.

localized - Found only at a specific location.

macroscopic - Large enough to be seen with the naked eye.

metabolize - To utilize a nutrient source for growth and maintenance.

microbiologist - A professional who studies organisms too small to be seen with the naked eye.

migration - The act of moving throughout the body and occupying a new environment.

mucins - Large proteins in saliva that give it hydrating properties.

normal flora - The community of microorganisms that is found in an environment during good health.

nucleoid - The region of the bacterial cell cytosol that contains the chromosome.

**O-antigen** - The outermost portion of lipopolysaccharide; composed of sugars linked together in chains.

**oligosaccharide core** - The central portion of lipopolysaccharide that links the O-antigen to lipid A; composed of sugars.

organelles - Discrete structures that carry out specific functions within a cell.

**outer membrane** - The outermost layer of a Gram-negative cell that contains both phospholipids and lipopolysaccharide.

pathogen - An organism that can cause disease.

peptide - A short sequence of amino acids linked together in a chain.

**peptidoglycan** - Chemical that makes up a bacterial cell wall; composed of a mixture of amino acids and sugars.

persistent - A state where a pathogen remains in an environment for a prolonged period of time.

**pH** - The measure of how acidic or basic a substance is; acids have low pH values and bases have high pH values.

phagocytes - Cells of the immune system that are able to engulf pathogens and parts of them.

**phospholipid bilayer** - The composition of cell membranes, made up of phosphate groups attached to lipid molecules.

pili - Bacterial hair-like projections that are made of protein and aid in attachment to surfaces and other bacteria.

plaque - The bacterial biofilm that accumulates on teeth.

**polymerase chain reaction (PCR)** - The method by which the amount of genetic material (DNA) can be selectively increased.

**polymicrobial infection** - An infection caused by more than one microorganism.

resolution - The ability to distinguish two objects as separate entities.

**ribosome** - The structure on which amino acids are synthesized into a protein.

saliva - The liquid produced in our mouths by the salivary glands that helps to maintain good oral health.

salivary antibody - Proteins in the mouth that are directed against specific pathogens.

salivary glands - The organs in the mouth that produce saliva.

**secretion systems** - Components that bacteria use to export material from the inside of their cells to the outside.

sialidase - An enzyme produced by some bacteria that breaks apart specific types of sugars.

**species** - The designation for organisms that are biologically identical to each other.

transpeptidation - Linking together sugar chains with peptides.

**vaccine** - A substance that can boost the immune response and protect us from subsequent infection by a specific pathogen.

virulence - The ability to cause disease.

# Laboratory safety procedures

- 1. Place all extra clothing, unnecessary books, purses, backpacks, and paraphernalia in an appropriate place. Racks are provided for these materials. The laboratory work area must be kept free of articles not actually in use.
- 2. Eating, drinking, and smoking are forbidden at all times in the laboratory.
- 3. Keep your locker or laboratory door clean. Do not allow your locker drawer to become filled with cultures that have no value in your current work.
- 4. Return all reagents, cultures, and glassware to their appropriate places.
- 5. Wear a laboratory coat, smock, or lab apron when working in the laboratory. This will protect clothing from contamination or accidental discoloration by staining solutions.
- 6. Do not place anything in your mouth while in the laboratory. This includes pencils, food, and fingers. Learn to keep your hands away from your mouth and eyes.
- 7. Avoid contamination of benches, floor, and wastebaskets.
- 8. Clean your work area (laboratory bench) with a phenolic disinfectant such as 5% Lysol or 5% phenol or a quaternary compound such as cetylpyridinium (Ceepyrn) before and after each laboratory period. This standard procedure lessens the chance for accidental infection as well as for contamination of cultures.
- 9. Special receptacles will be provided for infectious materials and used glass slides. Place all discarded cultures and contaminated glassware into these receptacles. Do not let unwanted and unneeded materials accumulate. Tall jars filled with a solution such as 5% Lysol or special receptacles will be provided for pipettes.
- 10. When infectious material is accidentally spilled, cover it immediately with a disinfectant such as 5% Lysol or 5% phenol and notify your instructor at once.
- 11. Flame wire loops and needles before and immediately after transfer of cultures. Do not move through the laboratory with a loop or pipette containing infectious material.
- 12. Wash your hands thoroughly before and after each experiment, using disinfecting soap if possible.
- 13. Label all experimental material with your:

a.	Name				
h	Data	1	1		

- c. Exercise number Ex. 5
- 14. Telephone number to call in case of an emergency 101, 103.

**Practical class 1.** Methods in diagnostic microbiology. Microscopic method of examination (MME). Basic morphological forms of bacteria. Simple methods of staining

#### Suggested reading for self-study:

History of the microbiology, virology, immunology department; main spheres of activity and trends in research. Design and equipment of microbiological laboratory, biosafety levels. Basic rules of work in microbiological laboratory (biosafety in work with class II biohazards). Universal precautions in work with burners and electric supplies.

Taxonomy of microorganisms: classification and nomenclature. Modern approaches to taxonomy of microorganisms. Taxonomic ranks. Vars (types), strains, clones, pure cultures.

Basic morphological forms of bacteria. Morphological characteristics of cocci, rods and spiral-shaped bacteria.

Microscopic method of examination: tasks, procedure, method evaluation. Bright-field light microscope: components and proper use of the microscope. Smear preparation and fixation. Simple methods of staining. The technique of oil immersion microscopy.

Signature of the tutor	

Oral quiz	Laboratory work	Individual work	Tests	Total results

use of the microscope. Smear preparation and fit	xation. Simple methods of staining. The techn	ique of oil immersion microscopy.						
	Laboratory work							
Laboratory exercises	Laboratory report							
<ol> <li>Prepare heat-fixed slide of Escherichia coli, cultured on agar medium, stain with methylene blue, examine under the oil immersion lens and complete the report.</li> <li>Prepare heat-fixed slides of Staphylococcus spp., cultured on liquid medium, stain with basic fuchsin, examine under the oil immersion lens and complete the</li> </ol>	Smear Stain							
report.  3. Complete the drawings of slides seen in demonstration room:  - Streptococcus spp., pure culture, stained with crystal violet;  - Vibrio spp., pure culture, stained with basic fuchsin;  - Bacillus spp., pure culture, stained with crystal violet.	SmearStain	4 Smear Stain	SmearStain					

		INDIVIDUAL WORK						
	Fill the numbers in the table  Biosafety Levels for Infectious Agents BSL							
1 2 3 4 5 6	-	according to the picture above:	Fill in	the empty cells examples of microorganisms in accordance with the level of risk				
※ 八〇 沙沙 時間		bacterium bipolar - staining bacterium clostridium	1	Agents that typically do not cause disease in healthy adults; they generally do not pose a disease risk to humans				
COCCI 8 9 10 11 12 13	14	coccobacterium diplobacterium diplococcus	2	Agents that can cause disease in healthy adults; they pose moderate disease risk to humans				
		fusobacterium micrococcus sarcinae	3	Agents that can cause disease in healthy adults; they are airborne and pose a more serious disease risk to humans				
RODS  15 16 17 18 1	9~	spirollum spirochete (borrelia) spirochete (leptospira)	4	Agents that can cause disease in healthy adults; they pose lethal disease risk to humans; no vaccines or therapy available				
SPIRAL-SHAPED	3,5	spirochete (treponema) staphylococcus streptobacillus streptococcus tetrad		Write the names of the parts of a microscope				
Questions for self-control and discussion:  1. What are the two purposes of heat fixation?  2. What is the purpose of simple staining?		vibrio F THE MICROSCOPIC METHOD OF EXAMINATION (WRITE IN THE CELL)		(5) (4) (6) (6)				
<ul><li>3. Why are basic dyes more successful in staining bacteria than acidic dyes?</li><li>4. Name three basic stains.</li></ul>	1			(10) (9)				
<ul><li>5. Why is time an important factor in simple staining?</li><li>6. How would you define a properly prepared bacterial smear?</li></ul>	3			(17) (8) (11)				
7. Why should you use an inoculating needle when making smears from solid media? An inoculating loop from liquid media?	4			(12)				
<ul> <li>8. Why is oil necessary when using the 90× to 100× objective?</li> <li>9. What are three bacterial shapes that you have observed?</li> <li>10. How can you increase the resolution on your microscope?</li> <li>11. In microbiology, what is the most commonly used objective?</li> </ul>	5			(14)				

# Practical class 2. MME. The morphology and fine structure of bacteria. Differential methods of staining

#### Suggested reading for self-study:

Distinctive features of prokaryotic and eukaryotic cells. Basic bacterial cell structure: components of bacterial cell. The composition, function, detection methods of bacterial cell wall. Gram stain: medical application, principles, procedure for Gram stain.

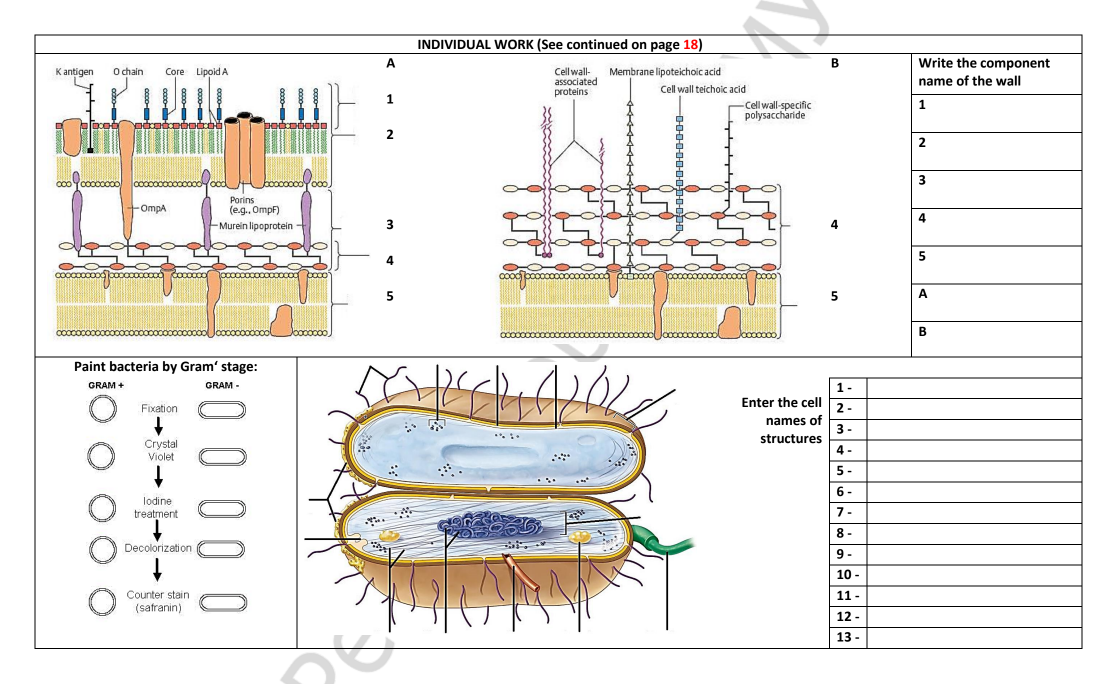
The composition, function of capsule, flagella, pili (fimbriae) and methods for their detection. Detection of capsule using negative staining.

The cytoplasmic membrane: structure, function. The most important bacterial cytoplasmic membrane proteins. Bacterial core: cytoplasm, cytoplasmic structures (nucleoid, plasmids, ribosomes, and mesosomes). Inclusion bodies - storage granules (starch, fat, sulfur, polymetaphosphate (volutin)). Methods for nucleoid and volutin detection. Loeffler and Neisser stain for volutin granules.

Acid-fast bacteria and unique properties of their cell wall. Ziehl-Neelsen acid-fast staining: medical application, principle, procedure.

נו	Signature	of the tuto	<b>r</b>		
:	Oral quiz	Laboratory work	Individual work	Tests	Total results
,					
,					

	Laboratory work					
Laboratory exercises		Laborato	ory report			
<ol> <li>Prepare heat-fixed slide of the mixed culture of <i>Escherichia coli</i> (gram-negative) and <i>Staphylococcus aureus</i> (gram-positive), Gram stain, examine under oil immersion and complete the report.</li> <li>Complete the drawings of slides seen in demonstration room:         <ul> <li>slide with capsule of <i>Klebsiella pneumoniae</i>, negative staining;</li> <li>slide with mixture of <i>Escherichia coli</i> (gramnegative) and <i>Staphylococcus aureus</i> (gram-positive), Gram stain;</li> </ul> </li> </ol>	Stain	2 Smear Stain	3 Smear Stain	4 Smear Stain		
<ul> <li>slide with volutin granules of Corynebacterium diphtheriae, Loeffler staining;</li> <li>slide with volutin granules of Corynebacterium diphtheriae, Neisser staining;</li> <li>slide of the mixed culture of asid-fast and asid-liable microorganisms, staing Ziehl-Neelsen.</li> </ul>	(1111/1111/1111/1111)	6 Smear Stain	7 Smear Stain			



# Practical class 3. MME. The morphology of the spirochetes, actinomyces, rickettsia, chlamydia, mycoplasmas

#### Suggested reading for self-study:

Bacterial forms with defective cell wall (protoplasts, spheroplasts and L forms): factors inducing cell wall removal, medical importance of L-forms.

Resting forms of microorganisms. Bacterial endospores: medical importance, properties of endospore, the periods of endospore formation, detection methods. Spore stain using Ozheshko method: principle, procedure.

Taxonomy, morphology, medical significance of the Spirochetes, Actinomyces, Rickettsiae, Chlamydiae, Mycoplasmas.

Romanowsky-Giemsa stain. Dark-field light microscopy. Phase-contrast light microscopy. Fluorescence microscopy.

# f Oral quiz Laboratory Individual work Tests results

	Laboratory work					
Laboratory exercises		Laborato	ry report			
<ol> <li>Prepare slide of <i>Rickettsia spp.</i>, stain with fuschin, examine under the microscope, complete the report.</li> <li>Complete the drawings of slides seen in demonstration room:         <ul> <li>slide with Treponema denticola in dental plaque, Gram stain;</li> <li>Leptospira spp., dark-field microscopy;</li> <li>Borrelia recurrentis in the blood of patient with relapsing fever, Romanowsky-Giemsa stain;</li> </ul> </li> </ol>	Stain	2 Smear Stain	3 Smear Stain	4 Smear Stain		
<ul> <li>Chlamydia inclusions in cytoplasm of host-cell, Romanowsky-Giemsa stain;</li> <li>slide with Actinomyces spp., pure culture, Gram stain;</li> <li>slide with spores of Bacillus anthracis, Ozheshko staining;</li> <li>slide with E. coli, pure culture, acridine orange stain.</li> </ul>	5 Smear Stain	6 Smear Stain	7 Smear Stain	8 Smear Stain		

		INDIVIDUAL	. WORK	,	
Morphology of Spirochetes (write in cells names of structures)  Endoflagella (axial filaments) beneath outer membrane, Basal body, Outer membrane, Endoflagella, Periplasm, Cell wall (peptidoglycan), Inner (cell/plasma) membrane, DNA in nucleoid, cytoplasm			Confront Gram-positive and Gram-negative bacteria		
1 2	1		Characteristic	Gram-Positive	Gram-Negative
			Number of peptidoglycan layers		
			Overall thickness in nm		
4		Specific compounds			
3 5			Interbridges between tetra peptides of neighbor glycan chains		
4	6				
5 7		Periplasmic space			
7	8	4	Porin proteins		
0.1 μm	9		Permeability		
The technique of Gram stain			Secretion systems		
(write the component and exposure Component: crystal violet, tag water, basic fuchsine or safranin,	-	odine	Flagella fixation in cell envelope		
component		exposure time,	Main mechanisms of genetic exchange		
1			Cell wall deficient forms in vitro		
2	- 4		Ability to produce spores		
3			Ability to produce long filamentous		
4		7	Susceptibility to Lysozyme		
5			Adhesion by pili		
6			Pathogenicity islands		
7 Tag water (wash slide thoroughly)		5	Gram stain (fill)		

	INDIVID	UAL WORK	
Questions for self-control and	discussion (Practical class 2)	Questions for self-control and discussion (Practical class 3)	
What is the function of the iodine solution in the Gram stain? If it were omitted, how would staining results be affected?	result	For what diseases would you use an acid-fast stain?	
What is the purpose of the alcohol solution in the Gram stain?		What chemical is responsible for the acid-fast property of mycobacteria?	
What counterstain is used? Why is it necessary? Could colors other than red be used? What is the advantage of the Gram stain over the simple stain?  Describe at least two conditions in which an organism might stain gram variable.	result	How should the acid-fast stain of a sputum specimen from a patient with suspected pulmonary Nocardia infection be performed?  Is a Gram stain an adequate substitute for an acid-fast stain? Why?	
Which step is the most crucial or most likely to cause poor results in the Gram stain? Why?	. (	Are acid-fast bacteria gram positive or gram negative? Explain your answer.	
Why must young cultures be used when doing a Gram stain? What is meant by gram variable?		Why is it important to know whether bacterial cells possess flagella, or endospores?	
What part of the bacterial cell is most involved with Gram staining, and why?		What do endospore stains have in common with the Ziehl-Neelsen acid-fast stain? Is bacterial sporulation a reproductive process? Explain.	
What is an advantage of negative staining?	70	What is the purpose of the heat during the acid-fast staining procedure?	
Why is negative staining also called either indirect or background staining?		Why are endospores so difficult to stain?	

# Practical class 4. Ecology of microorganisms. Asepsis. Methods of sterilization, disinfection and antisepsis

#### Suggested reading for self-study:

Ecology of microorganisms. Interspecific and intraspecific relations. Symbiosis, its variants. Antagonistic microbial relationships, its background and medical importance. Bacteriocins.

Definition of terms asepsis, sterilization, disinfection, antisepsis. Methods of sterilization: physical, chemical, mechanical. Differences between sterilization and disinfection. Types and methods of disinfection. Types and methods of antisepsis. Practical antisepsis. Classification of antiseptics, origin and characteristics of groups. Mechanisms of action on microorganisms. Antimicrobial management in dentistry.

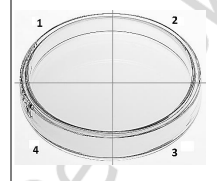
f Oral quiz Laboratory Individual Tests Total		Signature	of the tuto	r		
work work results	s f	Oral quiz	Laboratory work		Tests	Total results

#### Laboratory work

# 1. Test the effectiveness of hygienic and 1. Divide a nutrient agar plate into 4 sections with a marking pen or pencil. Mark each section of the plate with numbers 1, 2, 3, 4. practical class.

**Laboratory exercises** 

- **Laboratory report**
- surgical hand antisepsis. The result is 2. Mark each plate with your group number and your name.
- taken into account in the next 3. On the surface of agar medium at section N 1 make a fingerprint of skin untreated with any antiseptic (control).
  - 4. Wash your hands with soap as you do it usually at home and make a fingerprint on the surface of the agar medium at section N2.
  - 5. Wash your hands with soap twice and then your fingers with antiseptic (1% solution of iodopyron) 2 minutes, neutralize iodopyron with neutralizer (1% solution of sodium thiosulfate) for 2 minutes and make a fingerprint on the surface of agar medium at section N 3.
  - 6. Do not wash your hands and fingers with antiseptic (1% of iodopyron) 2 minutes, neutralize iodopyron with neutralizer (1% of sodium thiosulfate) for 2 minutes and make a fingerprint on the surface of agar medium at section N 4.
  - 7. Incubate Petri dishes at 37°C for 24 hours.
  - 8. After incubation count the amount of colonies grown at each section and fill in the table. Formulate the conclusion regarding effectiveness of hygienic and surgical hand antisepsis.





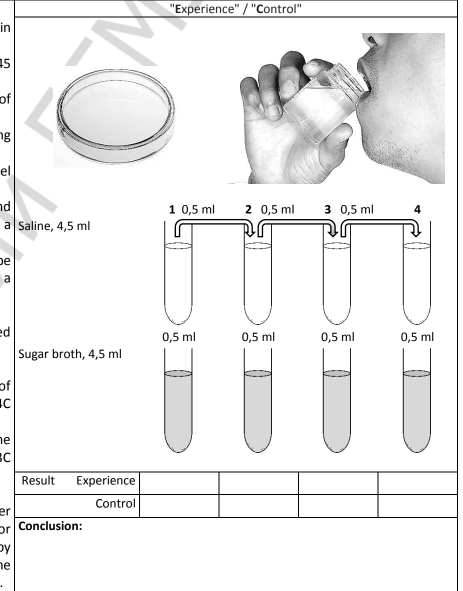
	<del>-</del>	
Section	Experiment description	Quantity of
		CFU
1	Control	
2	Hygienic hand antisepsis (washing with soap)	
3	Surgical hand antisepsis	
4	Antisepsis with iodopyron	

Conclusion:

account in the next practical class.



- 2. Test the effectiveness of hygienic oral 1. Mark the Petri plate "Experience" and "Control".
  - antisepsis. The result is taken into 2. Rinse mouth with sterile saline 45 seconds, and spit in the plate "Control".
    - 3. Rinse the mouth with 1% solution of boric acid 45 seconds and spit into the sink.
    - 4. Rinse mouth with sterile saline, and spit in the plate of "Experience".
    - 5. Using a sterile pipette and spray bulb make breeding materials:
    - a) prepare 4 test tubes with 4,5 ml of sterile saline, label 1C, 2C, 3C, 4C;
    - dial 0,5 ml of material from the plate "Control" and release into the tube 1C. Reset the pipette into a Saline, 4,5 ml porcelain cup;
    - other pipette to mix the contents of the tube 1C, type 0,5 ml tube and release in 2C. Reset the pipette into a porcelain cup. Do this with the other tubes.
    - b) analogous prepare "Experience" material.
    - 6. Use a glass pipette and spray bulb produce seed dilutions on sugar broth:
    - prepare 4 tubes with Sugar broth sign 1C, 2C, 3C, 4C;
    - sterile pipette to stir the contents of the tube 4C gain of diluted material 0,5 ml in a test tube and release 4C broth:
    - without changing the pipette, transfer 0,5 ml of the diluted material from the tube into the tube 3C broth; do this with the other tubes.
    - 7. Analogous prepare "Experience" material.
    - 8. Incubate all tubes at 37°C for 24 hours. After incubation observe each tube for growth (+) or Conclusion: absence of growth (-). Complete the table by recording your own results and formulate the conclusion regarding effectiveness of oral antisepsis.



	INDIVIDU	AL WORK	
Enter in cells possi	ble methods of sterilization	Give the definition of the	ne following terms:
Bacteriological loops		Antisepsis -	
Gauze, cotton, bandage		Asepsis -	
Rubber, plastic products		Disinfection -	
Glass products		Sterilization -	
Air in operating room		Modes of action of disinfectants	and antiseptics (write in cells)
General-purpose media		Mode	Disinfectants or antiseptics
Enriched media with serum or blood			
Solution which is inactivated at above 60 °C	95		
Borer			
Dental mirror			
Tooth brush			

**Practical class 5.** Bacteriological method of laboratory diagnosis of infectious diseases. Techniques for pure culture isolation and maintenance

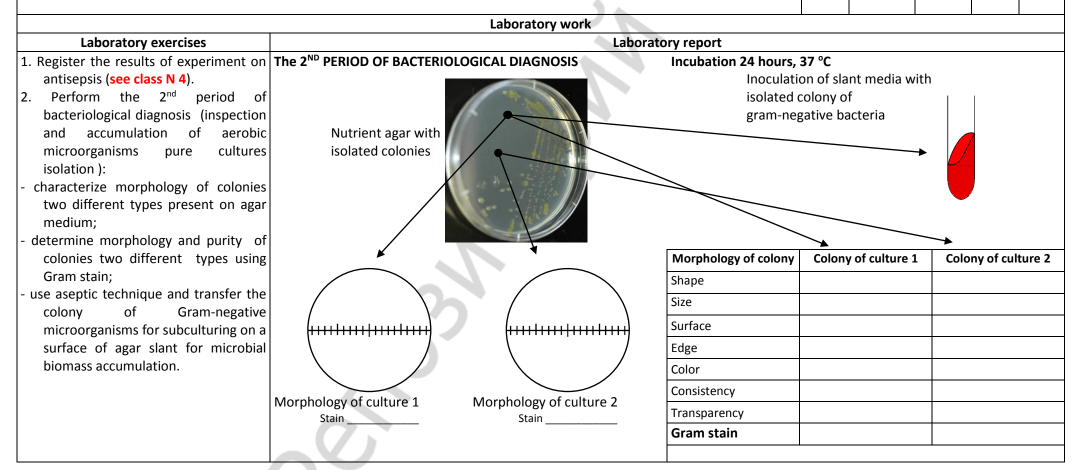
### Suggested reading for self-study:

Metabolism and energy exchange in microbes. Constructive and energy metabolism. Types and methods of feeding, nutrient transport through the membrane. Breathing microbes, breathing apparatus, ways of biological oxidation. Aerobic, anaerobic, facultative anaerobes.

Cultivation of microorganisms. Conditions required for growth. Nutrient media for culturing bacteria: classification and characteristics. Culture media ingredients, procedure of preparation and sterilization. General requirements to bacteriologic nutrient media. Incubator.

Bacteriological method of laboratory diagnosis: tasks, procedure, evaluation of the method. Methods of aerobic and anaerobic microorganisms isolation in pure culture. Bacterial colony characteristics.

t	Signatu	ire of the ti	utor		
	Oral quiz	Laboratory work	Individual work	Tests	Total results



	INDIVIDUAL WORK
Ques	stions for self-control and discussion:
Define a pure culture, a mixed culture.	
Define a bacterial colony. List four characteristics by which bacterial colonies may be distinguished.	
Why should a Petri dish not be left open for any extended period?	
Why does the streaking method of plates inoculation result in isolated colonies?	
Why are culture media sterilized before use?	
Discuss the relative value of broth and agar media in isolating bacteria from mixed cultures.	
At what temperature does agar solidify? At what temperature does agar melt?	
Define a culture medium.	
Discuss some of the physical and chemical factors involved in the composition, and in the preparation, of a culture medium.	
Why is it necessary to isolate individual colonies from a mixed growth?	
Are the large numbers of microorganisms found in the mouth cause for concern? Explain.	
Why are plate cultures incubated in the inverted position?	
How do you decide which colonies should be picked from a plate culture of a mixed flora?	
Why is it necessary to make pure subcultures of organisms grown from clinical specimens?	
How can you determine whether a culture or subculture is pure?	
What kinds of clinical specimens may yield a mixed flora in bacterial cultures?	
When more than one colony type appears in a pure culture, what are the most likely sources of the extraneous organisms?	

# Practical class 6. Bacteriological method of infectious diseases laboratory diagnosis. Techniques for pure culture identification

#### Suggested reading for self-study:

Identification of microorganisms: approaches and methods. Bacterial species: term definition, species criteria and methods for discovering bacterial species.

Biochemical activities of bacteria and methods for the biochemical properties detection of microorganisms. Enzymes of microorganisms: classification, importance for identification: a) proteolytic (proteases, peptidases, decarboxylases, deaminases, cysteine desulfurase, urease, tryptophanase); b) carbohydrate hydrolyses (carbohydralyses, amylase); c) lipolytic (lipases, lecithinase); d) oxidative- reductive (dehydrohenase, oxidase, catalase); e) hemolysins;  $\alpha$ -,  $\beta$ -,  $\gamma$ -, -hemolysis.

Rapid multitest systems for microorganisms identification. Automatic bacteriological analyzers: structure and principle of bacterial identification.

f	Signature	of the tuto	or		
,	Oral quiz	Laboratory work	Individual work	Tests	Total results
f					

		La	aboratory w	ork						
Laboratory exercises				Labor	atory repor	rt				
diagnosis (identification of aerobic microorganisms pure cultures): determine morphology and confirm purity of agar slant culture; using stab technique inoculate Hiss media with sucrose, maltose, mannitol for the determination of bacterial carbohydrate hydrolyses; using stab and streaking technique inoculate			Smear Stain _			\	Key YE	LLOWY 6,8-	RED<8,2 CR	
Kligler Iron agar for the determination of bacterial carbohydrate hydrolyses and H <sub>2</sub> S production; using stab technique inoculate semisolid tube medium to detect motility; inoculate nutrient broth and test the culture for the indole production.  Demonstration: semisolid and liquid Hiss media with different pH indicators; hemolysis on blood agar medium, lecitinase activity, indol detection; differentiate among members of the family <i>Enterobacteriaceae</i> using Kligler Iron agar; rapid multitest systems for identification of microorganisms.	glucose, lactose H <sub>2</sub> S production Carbo hydrases cysteinedesulfu rase	Semiliquid nutrient medium motility detection	Hiss medium sucrose  Carbo hydrase	Hiss medium maltose  Carbo hydrase	Hiss medium mannitol  Carbo hydrase	n Nutrient bullion  indole detection tryptophanase				

DA CEPTION OF COLUMN ASSESSMENT OF COLUMN ASSESSMEN	
BACTERIOLOGICAL METHOD OF LABORATORY DIAGNOSIS – 5 I's	
1 2 3 4	1

**Practical class 7.** Molecular Basis of Bacterial Genetics. Molecular methods of infectious diseases diagnosis and bacterial genetic investigations

Schette investigations														
Suggested reading for self-study:								7						
The structure of bacterial genetic appa	ratus. Regulation of ger	ne expression. General	prope	erties	and va	rietie	s of p	lasmid	ds. Si	gnatu	re of the tutor	•		
Detection of plasmids. Bacterial variability: pl							dec							
genetic variability: Mutation and recombination	<i>7</i> .	_												
Molecular methods: tasks, specimens f	or investigation, advant	ages of the methods.				- •								
Molecular hybridization: test material	s, DNA extraction, com	ponents of DNA hybri	dizatio	on rea	action	mole	cular	probe	es, d	Oral qui	Laboratory	Individual	Tests	Total results
detection of DNA hybrid duplexes, interpretat	ion of results. Equipmen	nt. Practical application	of mo	olecul	ar hyb	ridizat	ion m	ethod		J. u. qu.	work	work		- Otal results
Polymerase chain reaction (PCR): test	naterials, principle, DN	A extraction, componer	nts of	PCR r	eactio	n mix	ture, <sub>l</sub>	orime	rs,					
PCR thermal cycle, detection of amplicons, int	erpretation of results. E	quipment for PCR. Prac	tical a	pplic	ation o	of PCR								
		Labor	ratory	y wor	rk 🦠	$\mathcal{I}$								•
Laboratory exercises				-	la T	abor	atory	repo	rt					
1. Identify isolated pure culture and	Species	Morphology		Bio	ochen	nical o	hara	cteris	tics		Conclusion:			
complete the final report:	- I	,			7						According to	morpholo	gical, cult	ural.
- register the biochemical properties of	I		Glucose	se	Maltose	Mannito	ose		<u>o</u>	Motility	biochemical	•	•	
tested pure culture in the table;	I		)nc	Lactose	alt	an	Sucrose	H <sub>2</sub> S	Indole	lot				)C 13
- analyze the results and determine the	I		<u>5</u>	ت	Į≥	≥ _	. ช	Ĩ	드	2	attributed to	)		
species of tested pure culture.	E. coli	Gram- rods	AG	AG	AG	AG	-	-	+	+				
species of tested pare culture.	S. Typhi	Gram- rods	<b>A*</b>	-	Α	Α	-	+	-	+				
	S. Paratyphi A	Gram- rods	AG	-	AG	AG	-	-	-	+				
	S. Schottmuelleri	Gram- rods	AG	-	AG	AG	-	+	-	+				
	X-microbe		,								* "A" – acid, "G"	- gas		
2. Perform PCR for the detection of	Procedure of PCR	417	1				1			I	,			
M.tuberculosis in the sputum of the	DNA extraction:													
patient with tuberculosis suspected.	Mark the tubes with th	e volume 1,5 ml with l	etters	S (sp	utum)	and N	IC (coi	ntrol).	. Add :	100 µl	of the sputum t	o the tube wi	ith letter S	and 100 µl o
patient with tabelealosis suspected.	negative control to the	tube marked with lette	er NC.	Shake	e the t	ubes t	horou	ighly a	and bo	oil in th	ne water bath fo	r 10 minutes	(in room 5	07).
Identification of M.tuberculosis in	PCR cocktail preparation	on:												
sputum is based on the detection of gen	Mark the tubes with th	e volume 0,5 ml with le	etters	S (spi	utum)	and N	C (cor	ntrol).	These	tubes	contain primer	s, dNTPs, Mg	Cl₂. Add 10	μl of
,		ul of liquid into PCR' tub			-						-	_		•
MPB64 unique for <i>M. tuberculosis</i> and	Detection of PCR prod	-		•		•	•	•			,	•		
M. bovis. PCR amplifies the fragment		products in agarose gel	l. UV d	detect	ion of	specif	fic PCF	R-proc	ducts i	n gel v	vith ethidium br	omide.		
with the size 357 bp. of this gene.		5 1101				•				J				
	Report:													
		ed 357 bp were / not	dete	rted	Snuti	ım iç	nositi	ve /	negat	ive fo	r Mycohacterii	um tuhercul	nsis	
	Specific products size	od 337 bp were / not	acte	cicu.	Spatt	111113	Positi	• • •	LEGUL		i iviyeobacterii	ann taberear	0313.	

Laboratory exercises				Laboratory report			
3. Perform the bacterial conjugation	In bacterial conjugation experiment	E. coli		3	Recombinant <i>E.coli</i>		E. coli
experiment:	donor E.coli is susceptible to streptomycin and synthesize	D (donor)	<b>-</b> +			<b>r</b> .	R (recipient)
- prepare the mating mixture by	threonine and leucine.		F <sup>+</sup>		t de la constant de l	F"	
aseptically transferring 0,5 ml of an overnight meat-peptone both	Recipient E.coli displays complementary properties:		tre⁺ leu⁺		tre leu	tre <sup>-</sup> leu <sup>-</sup>	
culture of donor and recipient <i>E. coli</i>	resistant to streptomycin and		str <sup>s</sup>		str	str <sup>R</sup>	
into the separate tube;	unable to synthesis threonine		J.,		361	30	
- mix and incubate at 37 °C for 1 hours;	and leucine. Recombinants of these two strains will have	1			1- donor		2
- confirm the resistance status and	combination of either the			_	2- Recipient		
leucine and threonine production by	donor or recipient strains'		4	2	3- recombinant		
the culturing donor, recipient and	characteristics and can be readily detected by using		E.				
recombinant E. coli on minimal	selective minimal media.						
medium supplemented with			1		1		
streptomycin.					Registration of THE		
			3		results after 24 hours		
			Mi	nimal medium	incubation at 37 °C		
				thout <b>threonine</b> and			
				ıcine, with			
			str	eptomycin 100 μg/ml			

	INDIVIDUAL WORK									
Bacterial conjugation - Draw a process diagram										
0 min	2 min	10 min	15 min	20 min						
	Pilus formation	DNA replication with continued pilus formation	DNA transfer	Conjugates separate						

	INDIVIDUAL WORK
The	e polymerase chain reaction (PCR), complete cells
Stages	Amplification
Evaluation of method	Practical application

Practical class 8. Infections. Application of laboratory animals in microbiology. Antibiotic susceptibility testing of microorganisms

#### Suggested reading for self-study:

Defenition of infection. Classification of infections. Bacterial pathogenicity and virulence. Measurements of virulence: ID50, LD50, DLM. The genetics of bacterial pathogenicity. Pathogenicity islands. Pathogenicity factors: adhesins, invasins, impedins, agressins, modulins. The role of bacterial biofilms. Methods of adhesins, capsule, invasins, toxigenicity detection.

Biological method (application of laboratory animals in microbiology): tasks, phases, evaluation of the method. Animal models for infectious diseases. Routs for animal infection. Ethical, humane and legal considerations involved in the use of laboratory animals.

Sources of antibiotics. Spectrum of action. Chemical classification of antibiotics. Mechanisms of action. Side effects. Principles for rational antimicrobial therapy. The problem of resistance to antimicrobials: definitions (intrinsic, acquired resistance), incidence, significance. Resistance mechanisms: non-genetic and genetic origin of drug resistance. Antibiotic susceptibility testing of microorganisms: methods and principles.

1. of	Signat	ure of the	tutor 		
al	Oral quiz	Laboratory work	Individual work	Tests	Total results
e					

#### Laboratory work **Laboratory exercises** Laboratory report 1. Perform the disk diffusion method Pure culture Inoculation on Incubation at 35°C-24 h (Kirby-Bauer) for determination of Müeller-Hinton agar antibiotic susceptibility of four different microorganisms which often infect humans - Staphylococcus aureus. Escherichia coli. Müeller-Hinton agar Pseudomonas aeruginosa, and (composition): Klebsiella pneumoniae. 1,0 ml of inoculum of Diameret of meat extract - 2,0 g; microorganisms casein hydrolysate – 17,5 q; Registration of results corn starch - 1,5 g;

agar – 17,0 g;

pH 7,4±0,2

aqua distillate – 1 l;

Müeller-Hinton agar

application of antimicrobial discs to the

surface of the inoculated agar plate

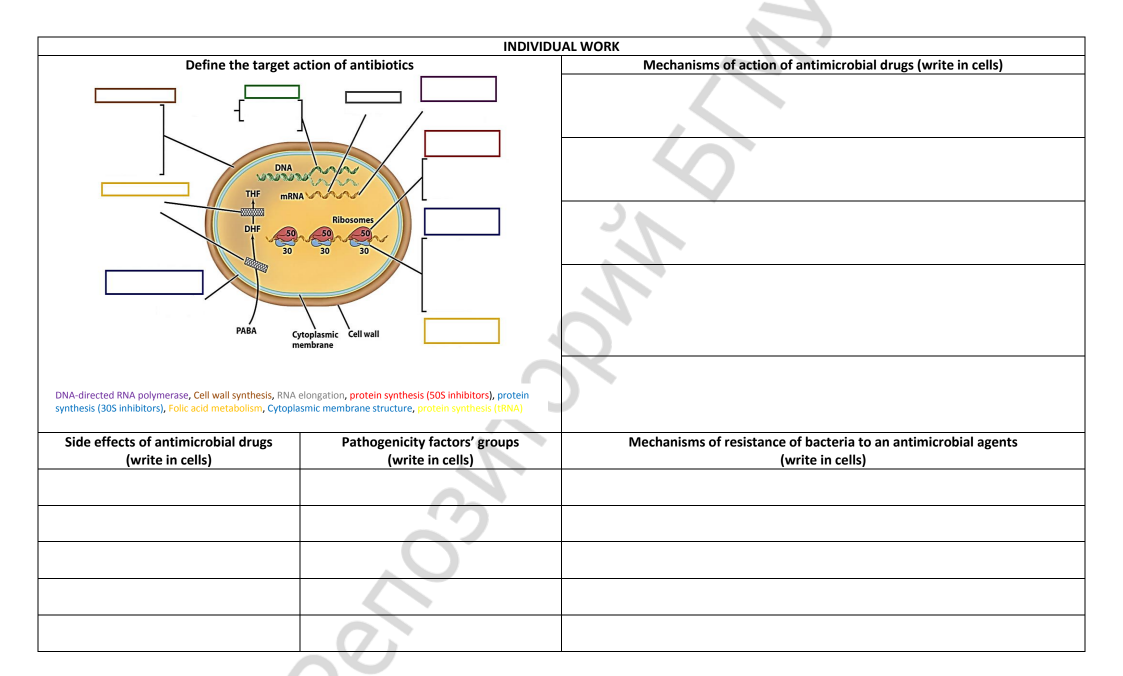
2. Determine antibiotic susceptibility of microorganisms by agar dilution test. Complete the report.

8 mcg/l 16 mcg/l 32 mcg/l control

Conclusion:

Petri dishes with serial doubled dilutions of Ampicillin in agar media Interpretation of results, MIC, mcg/l antibiotic resistant susceptible ≥32 Ampicillin Microbial culture MIC, mcg/ml Interpretation of results Culture 1 Culture 2 Culture 3 Culture 4

3. Determine antibiotic susceptibility of	Results of pure cultur	e		testi	ng by dis	c diffusio	method				
microorganisms by disk diffusion								Antibiotic	Diame	ter of inhibition zones (mm)	
•	Antibiotic		Diameter o	f inhibitio	n Inter	nrotation	of results	Antiblotic	resistant	suscep	otible
method, complete the report	Antibiotic		zone	, mm	inter	pretation	or results		Staphyloc	occus spp.	
(perform it at classes N 9).								Penicillin	≤28	≥2	9
								Oxacillin			
								S.aureus	≤10	≥1:	.3
								CNS	≤17	≥1:	.8
								Canamycine	≤13	≥1	.8
								Gentamicin	≤12	≥1	.5
								Ciprofloxacin	≤15	≥2	.1
						4		Tetracycline	≤14	≥1	9
								Erythromycine	≥23	≥2:	.3
4. Demonstration:						4		Lincomycine	≤13	≥2	.1
- agar disk diffusion test for antibiotic		2,0	4,0	8,0	16,0	32,0	Control	Chloramphenicol	<17	≥1:	.8
susceptibility testing of microorganisms;	μg/ml μg/ml μ	g/ml	μg/ml	μg/ml	μg/ml	μg/ml			Enterobac	teriaceae	
- rapid test for antibiotic susceptibility testing						_		Ampicillin	≤13	≥1	.7
of microorganisms; - slide of <i>Bacillus anthracis</i> in tissues of white mouse, Gram stain;	5s						M	Cefazolin	≤14	≥13	.8
		$\Rightarrow$						Cefotaxime	≤14	≥2:	.3
								Canamycine	≤13	≥18	.8
- slide of <i>Y.pestis</i> in tissues of white mouse,						ы		Gentamicin	≤12	≥1.	.5
-				1 1				Ciprofloxacin	≤15	≥2:	.1
Gram stain;				Т И	ЦЛ	III I	$\cup$	Lomefloxacin	≤18	≥2:	
- slide of Klebsiella pneumoniae		اك						Tetracycline	≤14	≥19	.9
rhinoscleromatis in tissues of white								Doxicycline	≤12	≥1	.6
mouse, Gram stain.								Chloramphenicol	≤12	≥1	.8
	DDM report (form therapy):  BDT report: minir µg/i	nal i						Stain		4-2 SmearStain	_



	INDIVIDUAL WORK	
Interacting factors of antimicrobial therapy (write in circle)	Characteristics of ideal antimicrobial drug:	Analyze the circuit in the picture (in the middle) and answer next. Which of the resistance mechanisms are shown in the figure?
Give the definition of the following terms:	= antibiotic	Methods of the antibiotic susceptibility testing (write
Antibiotic -	= altibulic	methods and indicate possibility to determine MIC)
Specific - antibacterial therapy Minimal -		
inhibitory concentration Multiple -		
resistance Pathogenicity -		

# Practical class 9. Credit "Morphology and physiology of microorganisms"

List of questions	Oral quiz	Script	Tests	Total results
List of questions				

- 1. History of microbiology as a science. Periods. The founders of microbiology main routs.
- 2. Microscopic method of examination: tasks, procedure, evaluation of the method.
- 3. Bright-field light microscope: components and proper use of the microscope. Dark-field light microscopy: the 26. Bacterial variability: phenotypic and genetic. Practical significance of bacterial variability. Population variability. microscopy. Fluorescence microscopy: principles behind the fluorescence microscopy. The technique of oil immersion microscopy.
- Type of microscopic preparations. Smear preparation and fixation. Simple methods of staining.
- Differential stains of microorganisms. Gram stain: medical application, principles, procedure for Gram stain.
- 6. Morphology of bacteria. Distinctive features of prokaryotic and eukaryotic cells. Basic morphological forms of bacteria. Morphological characteristics of cocci, rods and spiral-shaped bacteria. Motility of bacteria, methods of | 30. Role of microorganisms, social and physical factors in infection emergence. detection.
- Structure and function of cell envelope and appendages. Capsule. Detection methods of the capsule.
- The composition, function, detection methods of bacterial cell wall. The cell wall of gram-positive bacteria. The cell wall of gram-negative bacteria. Bacterial forms with defective cell wall. Factors inducing cell wall removal, medical importance of L-forms.
- Bacterial core: cytoplasm, cytoplasmic structures; their functions and detection methods. Acid-fast bacteria and unique properties of their cell wall. Methods of acid-fast staining: medical application, principle, procedure.
- 10. Resting forms of microorganisms. Bacterial endospores: medical importance, properties of endospore, the periods 36. Ecology of microorganisms. Basic terminology of ecology. of endospore formation, detection methods (principles, procedures).
- 11. Taxonomy of microorganisms: classification and nomenclature. Modern approaches to taxonomy of microorganisms. Taxonomic ranks. Vars (types), strains, clones, pure cultures.
- 12. Taxonomy, morphology, medical significance of the spirochetes. Methods for spirochetes detection.
- 13. Taxonomy, morphology, medical significance of Actinomyces.
- 14. Taxonomy, morphology, medical significance of Mycoplasmas. Methods for Mycoplasmas investigations.
- 15. Taxonomy, morphology, medical significance of Chlamydiae and Rickettsiacea.
- 16. Nutrition of microorganisms. Source of macro- and micronutrients, growth factors. Nutritional types. Transport mechanisms for nutrient absorption.
- 17. Energy strategies in microorganisms. Aerobic and anaerobic respiration. Structures involved in respiration in 4. microorganisms.
- 18. Reproduction of microorganisms. Mechanisms and phases of bacterial division.
- 19. Bacteriological method of laboratory diagnosis: tasks, procedure, evaluation of the method.
- 20. Cultivation of microorganisms. Conditions required for growth. Nutrient media for culturing bacteria: classification | 8. and characteristics. Culture media ingredients, procedure of preparation and sterilization. General requirements to bacteriologic nutrient media.
- 21. Methods of aerobic microorganisms isolation in pure culture.
- 22. Methods of anaerobic microorganisms isolation in pure culture. Cultivation of anaerobic bacteria: culture media. techniques, equipment.
- 23. Identification of microorganisms: morphological, cultural, serologic, biological, genetic.
- 24. Biochemical identification of microorganisms. Detection of: a) proteolytic enzymes; b) carbohydrate hydrolyses enzymes; c) lipolytic enzymes; d) oxidative- reductive enzymes; e) hemolysins. Automatic stations for identification of bacteria.

- 25. The structure of bacterial genetic apparatus. Phenotype, genotype, genome, genes. Regulation of gene expression. General properties and varieties of plasmids. Detection of plasmids.
- principle behind dark-field microscopy. Phase-contrast light microscope: basic principles behind phase-contrast 27. Molecular methods in diagnosis of infection diseases: aims, methods, advantages. Molecular hybridization and polymerase chain reaction: principles of the methods.
  - 28. Doctrine regarding infections. Terms for emergence of infectious disease. Basic terminology of infectology. Classification of infections.
  - 29. Role of microorganisms in infection emergence. Bacterial pathogenicity and virulence. The genetics of bacterial pathogenicity. Pathogenicity islands. Pathogenicity factors: adhesins, invasins, impedins, agressins, modulins.

  - 31. Biological method (application of laboratory animals in microbiology): tasks, phases, evaluation of the method.
  - 32. Chemoprophylaxis and chemotherapy; antimicrobial chemotherapeutic agents and antibiotics. Sources of antibiotics. Especially the use of antibiotics in dentistry.
  - 33. Mechanisms of antibiotics action. Side effects of antibiotics. Principles for rational antimicrobial therapy.
  - 34. The problem of resistance to antimicrobials: definitions (intrinsic, acquired resistance), incidence, significance. Resistance mechanisms.
  - 35. Antibiotic susceptibility testing of microorganisms: methods and principles.

  - 37. Asepsis: definition, surgical, medical asepsis, asepsis in microbiological laboratory.
  - 38. Sterilization: definition, methods of sterilization (physical, chemical, mechanical), quality control.
  - 39. Disinfection: definition, methods of disinfection.
  - 40. Antisepsis: definition, methods of antisepsis. Disinfectant and antiseptics: classification and modes of action.

#### List of practice.

- Prepare heat-fixed slide of bacteria, cultured on agar medium, stain with methylene blue.
- Prepare heat-fixed slides of bacteria, cultured on liquid medium, stain with basic fuchsin.
- Prepare heat-fixed slides of bacteria, cultured on liquid medium, stain by Gram.
- Technology immersion microscopy.
- Determine the morphology of Staphylococcus, pure culture, Gram stain.
- Determine the morphology of E. coli, pure culture, Gram stain.
- Determine the morphology of Gram+ and Gram- bacteria into the mix, Gram stain.
- Determine the morphology of the culture in smear colored by Ginsu-Burri.
- Define streptobacill pure culture morphology, Gram stain coloring.
- 10. Determine antibiotic susceptibility of microorganisms by disk diffusion method.
- 11. Characterize morphology of two different types of colonies present on agar medium.

# Practical class 10. Immune system. Innate immunity. Methods for innate immunity factors evaluation

#### Suggested reading for self-study: Signature of the tutor Human immune system: organs, cells, molecules (CD; receptors; MHC I, II, III; cytokines, adhesion molecules etc.). Immunity, types of immunity. Innate immunity. Immune and not-immune factors. Complement system: composition, way of activation, functions. Methods for estimation of complement system activity. Lysozyme, b-lysins. Polynuclear and mononuclear phagocytes systems. Phagocytosis: phases, intracellular killing mechanisms, outcomes. Laboratory Individual Oral quiz Tests Total results work work Dendritic cells. Methods for estimation of phagocytosis. Natural killer cells. Antigen-presenting cells. TOLL-like receptors. Laboratory work **Laboratory exercises Laboratory report** 1. Determine phagocytosis parameters in Staphylococci are mixed with leucocytes (50:1) and incubated at 37 °C | Smear\_ Stain for 15-120 min. Then slides are prepared and stained by Gimza method. prepared slides stained by Gimza Under oil immersion the phagocyting leucocytes and phagocyted method. staphylococci are counted and phagocytosis parameters calculated. 2. Complete the drawings of slides seen in demonstration room: PI (Phagocytosis index) = Number of phagocyting leucocytes / All incomplete phagocytosis of N. <del>(+++++++++++++++++++</del> <del>(++++1++++++++++++++</del> leucocytes counted gonorrhoea. Norma\* - 40-60 %. phagocytosis of incomplete PN (Phagocytosis number) = Number of phagocyted rhinoscleromatis. staphylococci / Number of phagocyting leucocytes Norma\* - 4-7. 3. Register the complement system activity by 50% hemolysis method. Volume of diluted (1:10) serum, ml 0,05 0,1 0,15 0,25 0,3 0,35 0,45 **50% hemolysis** $1 \text{ CH}_{50}$ – in ml serum Serum is diluted and added in wells from 0,05 to X CH<sub>50</sub> – in 1 ml serum 0,5 ml. Then saline solution is added to the final volume of 1,5 ml. 1,5 ml of hemolytic system is added to each well. Reaction is incubated at 37oC for 45 min, cooled at 4 °C $N 40 - 60 CH_{50}$ and centrifuged at 1500 rpm for 5 min. The well in which 50% hemolysis occurred is determined visually. This means the volume of patient's serum that contains one unit of **Results:** CH50. Then the CH50 for the whole serum is calculated.

		INDIVIDUAL WORK		
Fill cells with types of immunity			Fill with sample of	
immunity, adoptive, passive, natural, artificial, im factors, humoral, cellular, non-immune factors, a	mune	s of immune system	·	
		Write in cells ligand of receptor	s Associate t	he scientist and his discovery
	Pattern Recognition Receptors	<b>Ligand</b> pathogen-associated molecul	Edward Anthony	Phagocytosis, Cell-mediated immunity
	TLR1		Élie Metchnikoff	Chemical structure of antibodies
INNA	TLR2		Polly Celine Eveline Matzinger	Smallpox vaccine, vaccination
IININA	TLR3		Charles Alderson Janeway	side chains, humoral immune response
	TLR4	,0,	Rodney Robert Porter Gerald M. Edelman	Diphtheria antitoxin
	TLR5		Karl Landsteiner	Danger model, danger theory
	TLR6	1	Paul Ehrlich	Immune tolerance
	TLR7	7	Jules Jean- Baptiste Vincent Bordet	pattern recognition theory
active	TLR8		Emil Adolf von Behring	complement
	TLR9		Frank Macfarlane Burnet	blood group system, Rh factor, poliovirus

Compare  INNATE IMMUNITY  ADOPTIVE/ACQUIRED IMMUNITY  1 - 2 - 3 - 4 - 5 - 5 -  Complement system  Activation pathway activators  C3-convertase  C5-convertase  C5-convertase	ŀ
Complement system  Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	ssue
Complement system  Complement system  Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Complement system  Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Complement system  Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Complement system  Activation pathway activators  C3-convertase	
Complement system  Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Complement system  Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Complement system Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Complement system Phases of phagocytosis (write in cells)  Activation pathway activators  C3-convertase	
Activation pathway activators  C3-convertase	
Activation pathway activators  C3-convertase	
Activation pathway activators  C3-convertase	
Activation pathway activators  C3-convertase	
Activation pathway activators  C3-convertase	
C3-convertase	
C3-convertase	
C5-convertase	
MAC development	
The illustration shows the process of phagocytosis.  Granules	
Draw a picture of the possible outcomes of the Invagination of	
process in adjacent cells and named them.	
Bacterium	
Nucleus	
V 112 12 2	

# Practical class 11. Antigens. Antibodies. Immune response

20

30

determine an IgG, A, M concentration in serum by Manchini method (simple

radial gel immunodiffusion).

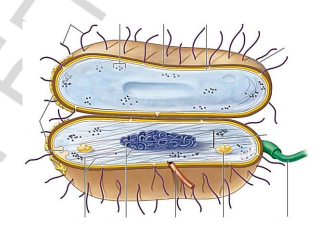
#### Suggested reading for self-study: Signature of the tutor Immune response, definition, main factors. Antigens: definition, main features, classification. B-lymphocytes system. B cells genesis. B cell receptor (BCR). B-cell activation, proliferation, differentiation to plasmocyte, immunoglobulin production. Humoral immune response. Primary and secondary humoral response. Immunoglobulins: structure, functions. Classes and subclasses of immunoglobulins. Monoclonal immunoglobulins. Laboratory Individual Total Oral quiz Tests work work results Methods of B-lymphocytes evaluation: quantitative and functional tests. T lymphocyte system. T-cell markers. TCR. Genetic control of TCR diversity. T-lymphocytes subpopulations: helpers, killers, DTH-effectors, regulators. T helpers of 1, 2, 3 and 17 types. Cellular immune response and its phenomena. Interaction and control of the immune system. Methods for evaluation of T- and B-lymphocytes system: quantitative and functional tests. Laboratory work **Laboratory exercises Laboratory report** 1. Determine the quantity of B-cells by Ν Count Ν Count Ν Count The method reveals CD20 antigen on B-Smear Smear Stain Stain cell surface: immune rosettes methods in ready-11 21 Normal B-cells count by CD20 = 8-20% made slides. 2 12 22 total blood lymphocytes. 3 13 23 2. Complete the drawings of slides seen 14 24 B<sub>CD20</sub>= rosette's Cell/30= in demonstration room: 15 25 - immune rosettes method for T-cell quantity 6 16 26 (Romanowsky-Giemsa determination stain); 27 7 17 Conclusion: blast transformation of lymphocytes 28 18 (Romanowsky-Giemsa stain); 19 29

#### **INDIVIDUAL WORK**

### Write figures for elements of an immunoglobulin molecule indicated on scheme

 <b>6</b>
Light chain (L)
Variable domen of the light chain
Constant domen of the light chain
Heavy chain (H)
Variable domen of the heavy chain
Constant domen of the heavy chain
Hinge fragment
Fc- fragment
Fab- fragment
Active center
Fc-receptor ligand

# Enter the names of structures of bacteria, which are antigens



Write the main cells and molecules that are involved in the humoral immune response molecules cells

structure
chain
a ±
Hinge region

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write nown the character	STICS OF IMMUNOVIONIUM ACCORDING TO CIASS AND MOJECUJE STRUCTURE
Wille down the character	sties of infinianoglobalin according to class and inforcedic structure
write abwii the character	stics of immunoglobulin according to class and molecule structure

	s contained on the same and a second of the sa	
structure	characteristics	class
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		lg D
		lg E
Hinge region		lg G
		lg M

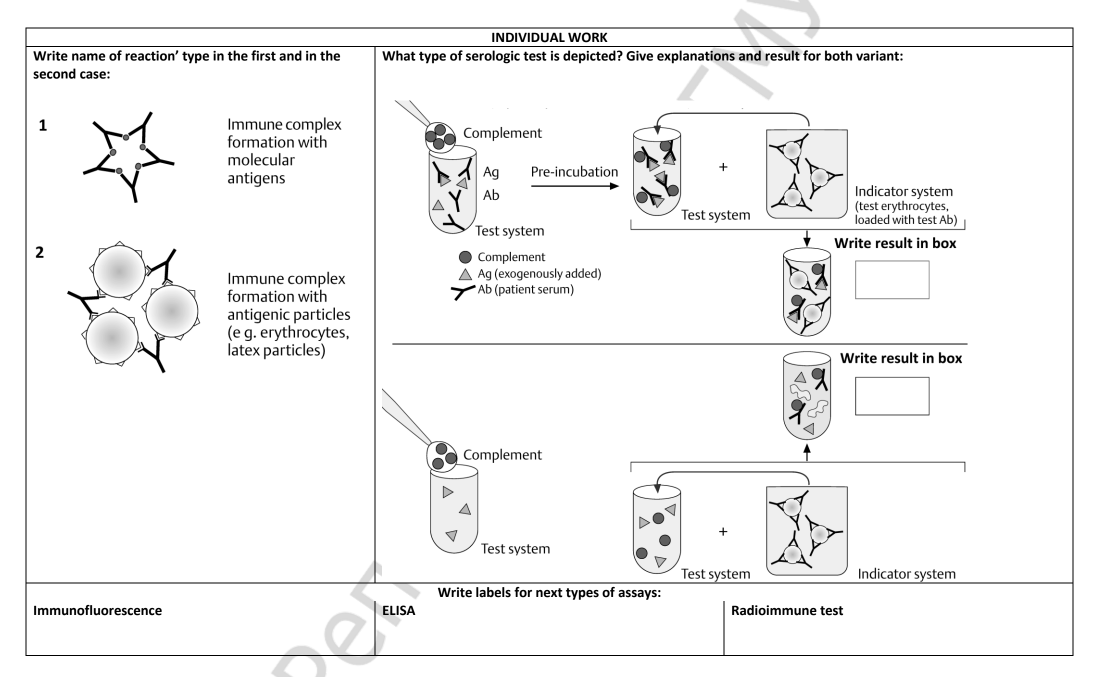
#### **INDIVIDUAL WORK** According to the Write methods of the study of cellular immunity Primary Response Secondary Response following diagram, draw Antibody concentration in serum (log scale) a graph of dynamics of immunoglobulins G and M classes for primary and secondary immune responses. 10 days 15 days 5 days 10 days First exposure Second exposure to antigen gG Plasma cells Plasma cells B cells Memory cells Source: Ryan KJ, Ray CG: Sherris Medical Microbiology, 5th Edition: www.accessmedicine.com Copyright © The McGraw-Hill Companies, Inc. All rights reserved. **Draw the B-lymphocyte** CD4 CD8 CD40b **BCR** CD3 TCR ΤCRα,β slgM slgD **CD19** IL4r **ACR CD52 CD20** ILR HLA **CD45 CD23 CD37** CD11C CD79a CD79b **CD38**

# Practical class 12. Serological method

#### Suggested reading for self-study: Signature of the tutor Serological method, characteristics. Antibody titre. Diagnostic titre. Diagnosticum. Diagnostic serum. Agglutination, passive agglutination, reversed passive agglutination, latex agglutination. Precipitation. Ring precipitation test, double immunodiffusion in a gel (by Ouchterlony), simple radial Laboratory Individual Oral quiz Total results immunodiffusion in a gel (by Mancini), immunoelectrophoresis, electroimmunodiffusion. Tests work work Immune lysis reactions. Complement fixation test: ingredients, implementation, characteristics. Immunofluorescence test: direct and indirect variants. Immunoenzyme test. ELISA. Radioimmune test. Laboratory work **Laboratory exercises** Laboratory report 1. Perform slide agglutination test to 1. antiserum 2. antiserum 3. Saline X-bacteria identify an X-bacteria. S.Typhi E.coli Conclusion: X-microbe is Determine the result **CFT** 1:20 1:40 1:320 SC AC of 1:80 1:160 complement fixation test. Key Assess: **Conclusion: PASSIVE BLOOD AGGLUTINATION TEST** 1/20 Key 1/10 1/40 1/80 1/160 1/320 1/640 SC AC 3. Determine the result of passive hemagglutination reaction. Assess: Conclusion:

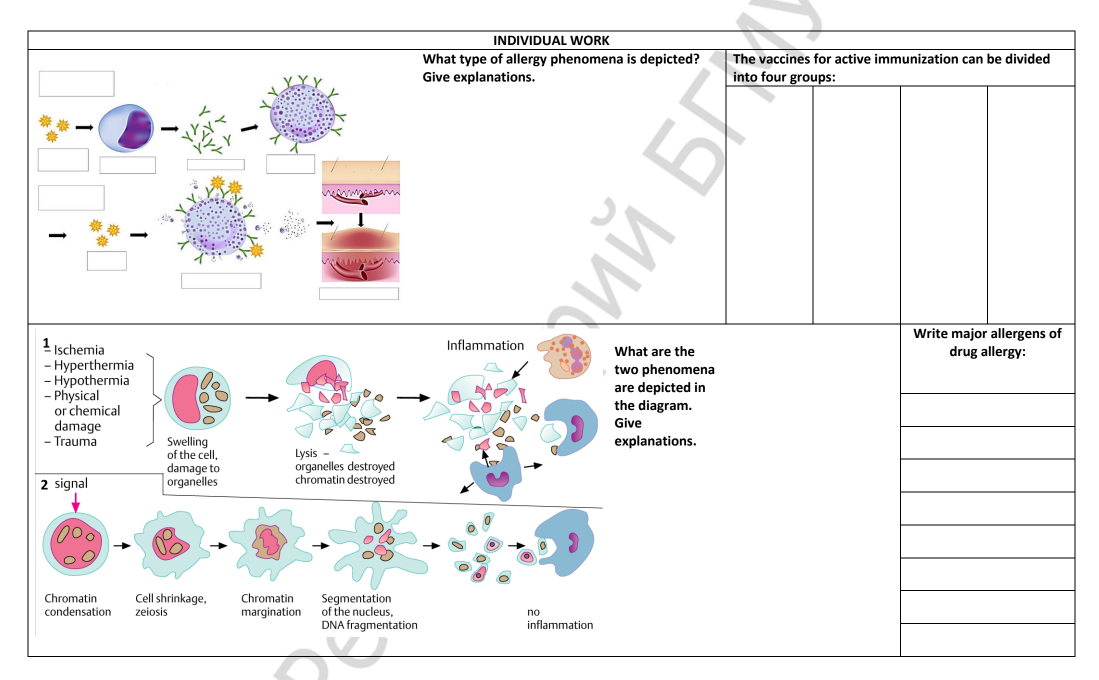
Laboratory exercises		Labora	itory report
<ul> <li>4. Perform ELISA for HBs antigen detection in donor serum:</li> <li>a) put 100 mcl of control serum and samples according to test scheme;</li> <li>b) put 50 mcl of conjugate in each well;</li> <li>c) incubate for 1 hour at 37°C;</li> <li>d) wash the strip 5 times;</li> <li>e) put 100 mcl of chromogen in each well;</li> <li>f) incubate for 30 min at 37°C;</li> <li>g) put 50 mcl of stop-reagent in each</li> </ul>	detecti on in the serum  E  G	Negative control Negative control Low positive control High positive control Sample 1 Sample 2 Sample 3 Sample 4	Test validity: - average OD of negative controls must be < 0,15 OD(NC) (negative controls) = - OD negative controls must range from 0,6 to 1,4 of average OD(NC) - 0,6 OD(NC) = - 1,4 OD(NC) = - average positive controls OD must be more than four times as much as OD(NC): average OD(PC)/ OD(NC) = - Low positive control OD must be higher than cut-off level Cut-off calculation:
well; h) measure the strip on ELISA reader and print out the results;	OD Sample 1		Cut-off = OD(NC) + 0,04  Conclusion:
i) fill in the report: check the test validity and make the final	Sample 2		
conclusion about results.	Sample 3 Sample 4	.07	

		INDIVIDUAL WORK	
Write down the fol	llowing definitions:		
Titer	-		
Diagnostic titer	-		
Diagnosticum	-		
Diagnostic serum	-		
	Direct variant	Draw the scheme of ELISA	Indirect variant
		Antigen – 🛆	
		Antibody –	
		Anti-Ig antibody – <b>G</b>	
		Anti-lg antibody – Genzyme - Cenzyme	



# Practical class 13. Immunoprophylaxis and immunotherapy. Immunopathology and clinical immunology Suggested reading for self-study:

Immunoprophylaxis and immunotherapy.	ina Manaimal immunitus fantamant	Cf = at: = = :t= al = al = = =		f.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		Signatur	e of the tut	or		
Vaccines, classification, essential characterist Passive immunoprophylaxis. Immune sera an				r vaccinal immunity	evaluation.					
Allergy, periods, types. Immediate type of hy				une complex type (I	III). Delayed type					1
of hypersensitivity mechanism (IV). Drug allergy. Aller				une complex type (i	iii). Delayed type	Oral quiz	Laboratory	Individual	Tests	Total
Clinic immunology: definition. Immune status		illergic conditions di	agriostics.		1	Oral quiz	work	work	rests	results
Primary and secondary immunodeficiency.	s. mmanogram.									
Autoimmune disease. Causes, manifestation.	Autoantihodies diagnostic value	e methods of deter	mination Antitu	mor immunity Met	hods of immune					
status correction. Immunosuppression. Immunostimi	· · ·									
	,		tory work							
Laboratory exercises				Laboratory repo	ort					
1. Perform the passive hemagglutination test	1. Saline 2. Patient's	3. ER	1. Saline	2.Patient's	3. Latex		Smear _			
for the detection of rheumatoid factor.	serum	Diagnosticum		serum	Diagnosticum		Stain			
Diagnosticum = armed bull erythrocytes			4 1							
coated with human IgG.										
Rheumatoid factor is an autological antibody										
(IgM) to IgG. It is found in certain		4					/		1	
autoimmune diseases (SLE, RA etc.) and							4		ш)	
is useful for diagnostics.					( )		<i>\\</i>	,	)	
2. Perform the LA test to detect							\		/	
autoantibodies to thyreoglobulin										
Latex diagnosticum = latex microsphera		7	/		7					
coated with thyreoglobulin molecules			/		/					
	/()(		/ (		/					
3. Demonstration:			/		/					
- degranulation of mast cells, Romanowsky-					/					
Giemsa stain;										
- Allergens;	Conclusion:		Conclusion:							
- Medicine for correction.										
							-			
			UAL WORK	10040	(4054)					
	Write down the typ	pes of allergy by	P.G.H.Gell ar	nd P.R.A.Coomb	s (1964):		_			
	0:									



## Practical class 14. Test "Immunology. Immunity. Allergy"

List of questions	Oral quiz Scr	ript	Tests	Total results
List of questions				

- 1. Immunology. Definition, tasks, methods. History of immunology.
- Immune system. Characteristics. Organs, cells, molecules of the immune system.
- Cytokines. Definition, classification. Biological importance.
- Immunity: definition, classification. Characteristics of anti-infection immunity.
- Innate immunity: definition, immune and non-immune factors, characteristics.
- Complement system: definition, ways of activation, functions. Medical importance. 32. Hypersensitivity of delayed type (IY): definition, classification, clinical phenomena. Methods of complement activity evaluation.
- 7. Phagocytosis. Phagocytosis phases. Phagocytosis outcome (complete, 34. Methods for DTH diagnostics (in vivo and in vitro). incomplete). Chemotaxins, opsonins: origin and medical importance.
- 8. Phagocytosis evaluation methods.
- 9. Immune response and factors influencing its strength.
- 10. B-lymphocytes, characteristics, main markers. Humoral immune response, periods.
- 11. Methods for B-lymphocytes quantity and functional activity evaluation.
- 12. Antigens: structure, classification, characteristics.
- 13. Bacteria antigenic structure. Cross-reacting antigens.
- 14. Antibodies, structure-functional organization of immunoglobulin molecule, characteristics. Antiidiotypic and monoclonal antibodies.
- 15. Classes of immunoglobulins, characteristics.
- 16. Mechanisms of antigens and antibodies interactions. Specificity. Phases. Affinity. Avidity.
- 17. Serology reactions, characteristics. Tasks, periods, clinical importance.
- 18. Agglutination reaction. Methods of conduction and result registration. Medical 43. Vaccinal immunity. Factors influencing vaccinal immunity. importance.
- 19. Passive hemagglutination, ingredients. Methods of conduction and result registration. 45. Immunocorrection. Methods for suppression and stimulation of the immune response, Medical importance. Reversed passive agglutination test. Latex agglutination.
- 20. Precipitation reaction. Methods of conduction and result registration. Medical importance.
- 21. Immunofluorescence test. Medical importance.
- 22. Immunoenzyme analysis. ELISA. Ingredients, methods of conduction, results registration, 2. characteristics. Medical importance.
- 23. Immune lysis reactions. Hemolysis.
- 24. Complement fixation test. Ingredients, methods of conduction, results registration, 5. Perform the slide agglutination test characteristics. Medical importance.
- 25. T-lymphocytes system, characteristics. Cellular immune response, dynamics.
- 26. Methods for T-lymphocytes quantity and functional activity evaluation.
- 27. Allergy: definition, classification. Allergy phases and types.
- 28. Allergens: definition, classification, characteristics.

- 29. Allergic reaction of immediate type, clinical phenomena.
- 30. Mediator type of ITH: definition, mechanisms, clinical phenomena, approaches for prophylaxis.
- 31. Cytotoxic (II) and immunocomplex (III) ITH types: definitions, mechanisms, clinical phenomena.
- 33. Methods for ITH diagnostics (in vivo and in vitro).
- 35. Immune tolerance: definition, mechanisms, medical importance.
- 36. Transplantation immunity. MHC antigens of I, II, III types, role for an immune response development. Transplantological reactions. Mechanisms of transplant rejection. Prophylaxis.
- 37. Clinical immunology: definition, aims.
- 38. Primary and secondary immunodeficiencies: definitions, classification, medical importance.
- 39. Immune status: definition, methods for evaluation. Influence of life way on the immune system function.
- 40. Autoimmune diseases, classification. Autoantigens. Mechanisms of autoimmunity.
- 41. Immunoprophylaxis and immunotherapy of infections. Achievements and problems.
- 42. Vaccines, main demands. Classification, characteristics, approaches to development. New vaccines.
- 44. Passive immunoprophylaxis. Antisera for therapy and prophylaxis, medical importance.
- drugs for immunocorrection.

### List of practice.

- 1. Register the result of agglutination test.
- Register the result of gel immunoprecipitation test.
- Register the result of complement fixation test.
- 4. Register the result of passive hemagglutination test.
- Determine the immunoglobulins concentration.
- Determine T-lymphocytes quantity in ready slide by immune rosettes method.
- 8. Determine phagocytosis indices in ready slides

## Practical class 15. Microbiological diagnostics of diseases caused by Staphylococci, Streptococci, Neisseria

### Suggested reading for self-study:

Staphylococci, general characteristics. Pathogenicity factors. Staphylococcal infection, including dentistry. Staphylococci as Signature of the tutor causative agents of nosocomial infections. Methods of staphylococcal infections microbiological diagnostics. The material for the research depending on the infection form. Scheme of pure culture isolation (from pus, mucus, blood, etc.). Identification methods, phagetyping of Staphylococci. Specific prevention and treatment of staphylococcal infections.

Streptococci, systematics, general characteristics. Antigenic structure. S.pyogenes, S.pneumoniae, S.mutans and other spp of the oral cavity. The role in the health and pathology of the oral cavity. Acute and chronic diseases, pathogenesis, immunity. Methods for streptococcal infections diagnosis. Bacteriological method, study design. Material for studies depending on the form of the infection, the rules and methods for taking material. Principles of therapy and prevention streptococcal infections.

Neisseria. Systematics, general characteristics. The role in the health and pathology of the oral cavity. Meningococcus, gonococcus. Pathogenicity factors. Pathogenesis and immunity. Microbiological diagnostics, material for studies. Specific prevention and treatment.

f	Oral quiz	Laboratory work	Individual work	Tests	Total results
١,					
ŝ.					

Laboratory v	work - practical class' duration in second semester is 2 academic hours 15 minute	es
Laboratory exercises	Laboratory report	
1. Microbiological diagnostics of staphylococcal		Staphylococcal colonies
infection, 2 <sup>nd</sup> period:	Stain	shape (form)
- macro- and microscopic examination of the		size/elevation
colonies on YSA;	(++++++++++++++++++++++++++++++++++++++	surface (appearance)
- plasmacoagulase test (stabilized rabbit plasma,		edge (margin)
37°C, 2-4-24 h).		pigmentation
		consistency
	Conclusion: according to morphological, cultural and biochemical properties unknown	transparency
	bacterium is identified as	lecithinase
<ul> <li>2. Microbiological diagnostics of streptococcal infection, 3<sup>rd</sup> period:</li> <li>the description of Streptococci growth in serum broth;</li> <li>determining the morphology of streptococci, Gram staining;</li> <li>determination of streptococcus serogroups by ring precipitation test.</li> </ul>	(++++++++++++++++++++++++++++++++++++++	

Laboratory exercises		Labo	ratory report	
3.Demonstration: - Staphylococcus aureus in pus, Gram staining;	SmearStain	Smear Stain	Smear Stain	Smear Stain
<ul> <li>Streptococcus pneumonia, pure culture, Gram staining;</li> <li>S.pneumoniae, white mice, Gram staining;</li> <li>Neisseria gonorrhoeae in pus, Gram staining;</li> <li>Neisseria meningitidis in cerebrospinal fluid, methylene blue;</li> <li>the growth of staphylococci on YSA, blood agar, broth;</li> <li>the growth of streptococci on blood agar and broth;</li> <li>coagulase test (plasma);</li> <li>anaerobic mannitol fermentation;</li> <li>phage typing of staphylococci.</li> </ul>	Smear Smear			

										II	۷DI	<b>VIDU</b>	AL۱	WOR	K									
							-	7		-														
									Ψ.															

# **Practical class 16.** Microbiological diagnostics of acute enteric infections caused by Enterobacteria. Methods for food poisoning diagnostics

Suggested reading for self-study:

General characteristics of Enterobacteriaces Escherichia, general characteristics. The bio Salmonella, classification and general chara	logical role of Escherichia coli i			of the tutor			
manifestations in the oral cavity.							
Shigella, classification, general characteristi	cs. The role in pathology.		Oral quiz	Laboratory work	Individual work	Tests	Total result
Common principle of microbiological diagno	osis of acute intestinal infection	i.		WOLK	WOIK		
Etiology of food poisoning. Principles of mic	robiological diagnostics.						
	Lab	oratory work					
Laboratory exercises		Laboratory	report				
1. Demonstration:	Smear	Sm	ear				
- <i>E. coli</i> , pure culture, Gram staining;	Stain	Sta	iin				
- Salmonella typhi pure culture, Gram staining;							
- Shigella flexneri pure culture, Gram staining;							
- clean media: Endo, Levin, Ploskirev, bismuth							
sulfite agar, Rapoport, magnesium, Kliglera;	\(\(\frac{1}{1+1+1+1+1+1+1+1+1+1+1+1+1+1+1+1+1+1+	4	+++++++	·······)			
- the same media with the growth of E. coli,	[[]	(					
Salmonella, Shigella;							
- biochemical activity of E. coli and Salmonella;							
2. Slide agglutination test with diagnostic O	Smear	Cli	de agglutin	ation tost			
and H-serum for identification of Salmonella.	Stain	311	de aggiutiin	ation test			
	l/ \				Conclusion	on:	
	<del>(++++++++++++++++++++++++++++++++++++</del>	V					
	/						
	7 .						

## **Practical class 17.** Microbiological diagnostics of diseases caused by Klebsiella, Campylobacter, Helicobacter and Pseudomonada

Signature of the tutor

Suggested reading for self-study:

Klebsiella, classification and general characteristics, main diseases caused.

#### Campylobacter, general characteristics, role in human pathology. Mechanisms of pathogenesis. Diagnosis of campylobacteriosis. Helicobacter. Laboratory Individual Pseudomonas aeruginosa, general characteristics, role in human pathology. Total Oral quiz Tests work work results Laboratory work **Laboratory exercises** Laboratory report 1. Microbiological diagnostics of Klebsiellosis, 3rd Smear Stain period: determine the biochemical properties of Klebsiella: perform slide agglutination test with anticapsule diagnostic sera and determine the K-antigen; determine the titer of CFT for serological Russell diagnosis of Scleroma. Slide agglutination test with anti-capsule serum **Biochemical** K. pneumoniae properties s. rhinoscleromatis s. ozaenae s. pneumoniae 1, 2 Glucose (A+G) +/-1, 3 Lactose 4 Saccharose (4th day) +/-5 Citrate 6 Urea -/+ 7 Malonate + O2b:K4 O1,3-5:K1-3 8 Antigens O2a:K3 К3 CA Conclusion:

Laboratory exercises				l	Laborat	ory re	port					
	1:5	1:10	1:20	SA	AC			C	OMPLEN	/IENT F	IXATIC	ON TEST
	1.5				I I	Var	Sei	rum dilu	tions	sc	AC	Result
							1:5	1:10	1:20			
						1	++++	++++	++++	-	-	Very positive
						2	++++	++++	-	-	-	Positive
				The state of the s	The state of the s	3	+++	-	-	-	-	Slight positive
				~ //		4	-	-	-	-	-	Negative
<ul> <li>2. Demonstration:</li> <li>- K. pneumonia s. rhinoscleromatis capsule (Hins-Burri staining);</li> <li>- K. pneumonia s. rhinoscleromatis, pure culture, Gram staining;</li> <li>- Pseudomonas aeruginosa, pure culture, Gram staining;</li> <li>- C. jejuni, pure culture, Gram staining;</li> <li>- Klebsiella growth on differential diagnostic media;</li> <li>- oxidase test.</li> </ul>	Stain	<del>        </del>		SmearStain	+++		ar			<u>S</u>	Stain	

## Practical class 18. Final test "General microbiology. Immunology"

List of auestions	Oral quiz	Script	Tests	Total results
List of questions				

- objectives, role in the dentist's practice.
- Milestones (periods) in microbiology. Work of Louis Pasteur, Robert Koch, Ilya Mechnikov. Evolution of microorganisms and infectious diseases.
- Common with other organisms and the unique features of microorganisms. Principles of microorganisms 20. The biological (experimental) method of diagnosing the infectious diseases: definition of the term, aim, tasks, systematics . Classification and nomenclature of microorganisms. The term of "species" in bacteria: group of traits for species identification (criteria for speciation).
- Morphology of bacteria. Basic morphological forms of bacteria. The bacterial cell structure. Functions of the surface and cytoplasmic structures of the bacterial cell. Mechanism of Gram staining. Forms of bacteria with the cell wall defects.
- Unique features of metabolism in prokaryotes. Nutrition of bacteria: types, requirements of bacteria, nutrients and pathways of nutrients penetration into the bacterial cell. Nutrient media: specification (what they should be | 23. The antisepsis: definition of the term, types, categories, methods of application. Antiseptic agents: classification, to provide the best growth of bacteria), classification.
- Respiration of microorganisms: types, pathways of energy production. Enzymes and cell structures involved into 24. The antibiotics: characteristics, classification, mechanisms of action. The rational antibiotic therapy principles. the process of respiration. Classification of bacteria regarding their oxygen requirements.
- Growth and reproduction of bacteria. The mechanism of simple division and its phases. Dormant forms of 25. Natural and acquired resistance of microorganisms to antibiotics. The genetic and biochemical mechanisms of microorganisms: general characteristics, factors inducing their formation, medical importance.
- Sampling for microbiological studies: types of samples, the rules of sampling, storage, transportation. Principles of organization, equipment and levels of biosafety in microbiological laboratories.
- Microscopic (bacterioscopic) method of diagnosing the infectious diseases: definition, aim and tasks, steps and evaluation of specificity, sensitivity, disadvantages of the method. Types of microscopic preparations. Staining of 27. The immune system. Central and peripheral organs of the immune system. Immunocompetent cells: microorganisms: methods. Types of microscopes.
- 10. The bacteriological method of the infectious diseases diagnosing: aim, tasks, phases, and evaluation of specificity, 28. Innate immunity. Innate immunity versus acquired immunity. Immune and non-immune factors of innate sensitivity, disadvantages of the method.
- 11. Methods for isolation identification of aerobic and anaerobic bacteria pure cultures. Identification of 29. The complement system: definition, main components, activators and activation pathways, functions of microorganisms without pure culture isolation.
- 12. Genetic apparatus of bacteria (nucleoid, plasmids, transposons, IS-elements) characteristics, functions, effect and 30. Natural killer cells and mechanisms of cytotoxicity. Phagocytes, classification. Phagocytosis reaction: phases, importance. The concept of genetic engineering and biotechnology.
- 13. Inheritance and variability of microorganisms. Types of variability. Mutations. The genetic recombination of bacteria. Phenotypic variability. The practical significance of the variability of microorganisms in the diagnosis, | 31. Antigens: structure, properties, classification. T-dependent and T- independent antigens. Superantigens. treatment and prevention of infectious diseases.
- 14. Molecular biological method of diagnosing the infectious diseases (molecular hybridization, polymerase chain reaction): definition, the principle of the methods, application in dentistry.
- 15. Effect of physical and chemical factors on microorganisms. Disinfection: definition of the term, aim and tasks, types, disinfectants, methods of disinfection quality control.
- Consequences of sterilization errors.
- 17. Infection (infection process): the term definition, causes and conditions of infectious diseases emergence. 35. Antibodies (immunoglobulins): structure, properties, classification, immunoglobulins biosynthesis. The Differences in communicable and non-communicable diseases. Periods of infectious diseases. Infectious diseases classification and outcomes.

- Microbiology: definition, area and fields of microbiology, methods of investigation. Dental microbiology: goals, 18. The role of microorganisms in the infectious process. Pathogenicity and virulence. Factors of pathogenicity of microorganisms. Pathogenicity Island. Microbial toxins. Types of exotoxins and their biological properties Mechanisms of microbial persistence and latency in host's organisms.
  - 19. The role of host, social, environmental factors in the infectious process.
  - phases, evaluation. Disbiosis: causes, consequences, prevention. Gnotobiology.
  - 21. The ecology of microorganisms. Types of ecological relationships in microorganisms. The role of microorganisms in the genesis and development of the Biosphere (the concept of the microbial dominance). Spread of microorganisms in the nature.
  - 22. The chemotherapy and chemoprophylaxis of infectious diseases. Groups of antimicrobial chemotherapeutic agents, mechanisms and spectrum of action on microbial cells. Chemotherapeutic index.
  - mechanism of action, side effects. Principles of rational antisepsis in dental practice.
  - Antibiotics for bacterial complications prophylaxis. Side effects of antibiotics.
  - microorganisms resistance. Genotypic and phenotypic methods for determining the microorganisms susceptibility to antibiotics.
  - 26. Immunology: definition of the term, aim and task, methods, history of development, branches. Immunity: definition, types of immunity.
  - classification, function, molecules.
  - immunity. Mechanisms of recognition in the innate immune system.
  - components and their fragments. Methods of the complement system activity evaluation.
  - mechanisms of intracellular microorganisms killing, outcomes. Methods of phagocytosis evaluation. Phagocytic reaction indexes, definition and importance in clinical practice.

  - 32. Antigens of microorganisms. Antigenic structure of bacteria. Type, species, group antigens. Protective antigens. Cross- reactive antigens, medical importance.
  - 33. Immune response: definition, conditions for development. Humoral immune response: definition, development Activation, proliferation, differentiation and interactions of cells involved. T-dependent and T- independent response. Primary and secondary humoral immune response characteristics.
- 16. Sterilization: the term definition, methods, quality control. Sterilization of instruments and medical devices. 34. B cells: development, markers, antigen-specific B cell receptor. Methods for B-lymphocytes quantity and functional activity assaying.
  - mechanism of interaction of antibodies with antigens: specificity, phases, manifestations. Affinity and avidity.
  - 36. Methods for the immunoglobulins concentration detection: simple radial immunodiffusion, ELISA, nephelometry. Monoclonal antibody: principles of production, application.

- 37. Serological method of investigation: general definition of the term, objectives, basic concepts (diagnosticum, diagnostic serum, titer, diagnostic titer, paired sera). Samples for serological examination. General characteristics of the method. Use of serological method for infectious and non-infectious diseases diagnostics.
- 38. Agglutination: ingredients, main variants of performance, registration, evaluation, application. Indirect (passive) and reverse passive agglutination: ingredients, mechanism, methodology, registration of results, practical use.
- 39. Immunoprecipitation reaction: ingredients, mechanism, main methods of performance, application. Reaction of the immune lysis. Complement fixation test: ingredients, mechanism, registration of results.
- 40. Solid phase immunoassay reactions. Immunofluorescence (fluorescent antibodies test, FAT), main variants, ingredients, mechanisms, registration of results, practical use. ELISA: ingredients, mechanisms, registration of results, practical use. Immunoblotting (IB). Radioimmunoassay (RIA).
- 41. T cells: development, markers, subpopulations. Helper T-cells, main types (Th1, Th2, Th3, Th17), spectrum of cytokines produced. Control of the immune response of T lymphocytes (Th3, T- regulators, CD4+CD25+Tcells). Methods for assaying of the amount and functional activity of T lymphocytes.
- 42. T-cell receptor: structure, types, genetic control, variety. T-dependent antigens. T- cell epitopes. T-cell restriction.
- 43. Cellular immune response: definition, development, main periods, manifestation. The model of two (three) signals: the response, anergy, apoptosis. Manifestation of cellular immune response. Immunological memory.
- 44. Anti-infection immunity and its types depending on pathogen nature. Innate and acquired defines mechanisms. Protective immunity. Mechanisms of antitoxic, antibacterial, antifungal, antiparasite immunity. Maternal immunity: mechanisms, significance.
- 45. Immunoprophylaxis and immunotherapy for infectious diseases. Active immunoprophylaxis. Vaccines: requirements, characteristics of main vaccines types (live, inactivated (corpuscular, chemical, conjugated, split, subunit), toxoids, genetic engineered). The concept of "ideal vaccine." Adjuvants mechanisms of action. New approaches for the vaccine development. Side effects of vaccination: sever vaccination complications.
- 46. Post-vaccination immunity: mechanisms and factors influencing its development. Indications and contraindications to vaccination. Immunization schedule. Expanded Programme on immunization. Collective immunity to infectious diseases, importance.
- 47. Passive immunoprophylaxis and immunotherapy of infectious diseases: indications, principles, complications. Classification of serum preparations (specificity, the manufacturing method, object of the antibodies action, purpose).
- 48. Allergology: the definition, objectives. Allergens. Allergy: the periods, types of reactions.
- 49. Allergic reaction in the oral cavity. Allergic method of investigation: definition, objectives, general characteristics, periods, evaluation.
- 50. Immediate type hypersensitivity (ITH). Mediator type (I) ITH: allergens, mechanism, development, manifestation, prevention of anaphylaxis. Cytotoxic (II) type ITH: allergens, development, mechanisms, manifestations. Immunocomplex (III) type ITH: allergens, development, mechanisms, manifestations.
- 51. Delayed type of hypersensitivity (IV): allergens, development, mechanism, manifestation (infection and contact allergy), importance in oral cavity.
- 52. Drug allergy: major allergens, the mechanisms and types of allergic reactions, methods for diagnostics and prevention.
- 53. Food allergy. Main allergens. Prevention of food allergy. Paraallergy. Idiosyncrasy.
- 54. Autoantibodies: origin, role in the pathology. Autoimmune diseases: definition, classification, etiology, mechanisms of tissue damage, manifestations. Principles of treatment. Prophylaxis.
- 55. Transplantation immunity. Histocompatibility antigens. Graft reaction types, mechanisms of development, prevention. Immunological tolerance: mechanisms, significance.
- 56. Clinical Immunology: definition, objectives, main concepts. Immune status: principle and methods of examination. Immunogram. Immunodeficiency conditions: classification, causes of development, methods for detection, principles for correction. Antitumor immunity. The concept of immune surveillance. Mechanisms of tumour escape from immune surveillance.

	MICROBIOLOGY	IMMUNOLOGY
INDIVIDUAL WORK		
TEST		
PRACTICAL SKILLS		
AVERAGE GRADE		
ABSENCE FROM PLACTICAL CLASS		
ABSENCE FROM LECTURE		
RATING		
Cred	lit (CROSS ) «PASSED»	«NOT PASSED»

## Practical class 1 (19). Microbiological diagnosis methods of diseases caused by Corynebacteria, Bordetella

Suggested reading for sel	f-study
Corynebacterium	diphth

neria, general characteristics of the pathogen. Types of Corynebacterium diphtheria, their distinctive features. Diphtheria toxin and antitoxic serum. The pathogenesis of diphtheria. Diphtheria in the oral cavity. Methods of diphtheria microbiological and molecular biological diagnosis. Principles of diphtheria therapy and prevention.

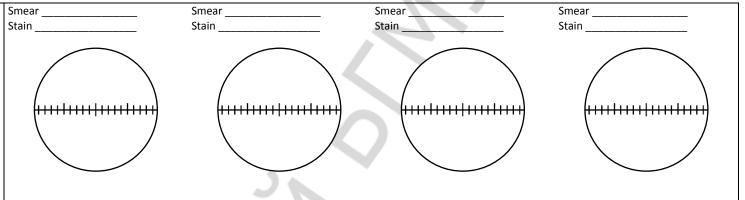
Bordetella pertussis and parapertussis. Characteristics of the pathogen, pathogenicity factors. The pathogenesis of pertussis, manifestation in the oral cavity, immunity, diagnostics. Principles of pertussis therapy and prevention.

	Oral quiz	Laborator	Individual	Tests	Total
r		y work	work	resis	results

### Laboratory work **Laboratory** exercises **Laboratory report** Colonies on serum tellurite agar 1. Bacteriological diagnosis of diphtheria, the 2<sup>nd</sup> Feature period: Shape describe the colonies Corynebacterium on potassium tellurite serum agar; Size - seed bacteria from typical colonies into Hiss Surface media (glucose, sucrose, starch). Edge Starch Color Consistency Biochemical properties of sertain corynobacteria Enzymatic activity Corynobacteria spp. with Acid production Cysteinase Ureasa Glucose Sucrose Starch C. diphtheriae gravis + C. diphtheriae mitis C. pseudodiphtheriae (hofmani) C. xerosis C. ulcerans X-microbe **Conclusion:** according to morphological, cultural and biochemical properties unknown bacterium is identified as

#### 2. Demonstration:

- *Corynebacterium diphtheria* stained by Neisser; *C.diphtheria* stained by Leffler;
- Bordetella pertussis, Gram staining;
- test for Corynebacterium diphtheria toxigenicity;
- preparations for specific prevention and treatment of diphtheria and pertussis;
- Growth of Bordetella pertussis and parapertussis on CCA, NA with tyrosine, urease test;
- assessment of antidiphtheria immunity intensity.



## Practical class 2 (20). Microbiological diagnosis methods of diseases caused by Mycobacteria and Actinomycetes

### Suggested reading for self-study:

Actinomycetes, systematic position, general characteristics, prevalence, role in the oral cavity pathology. Etiology, pathogenesis, microbiological diagnostics principles of the head and neck tissues actinomycosis.

Mycobacteria, general characteristics, resistance to acids. The causative agents of tuberculosis, species composition, morphology, nutritional needs, pathogenicity factors, differences from non-tuberculosis mycobacteria. The pathogenesis of tuberculosis, infectious granuloma, immunity, allergy, anergy. Principles of microbiological diagnostics of tuberculosis, immunoprophylaxis. TB chemotherapeutic drugs. TB symptoms in the oral cavity.

, , , , , , , , , , , , , , , , , , ,									
Oral quiz	Laborator y work	Individual work	Tests	Total results					
Signature of the tutor									

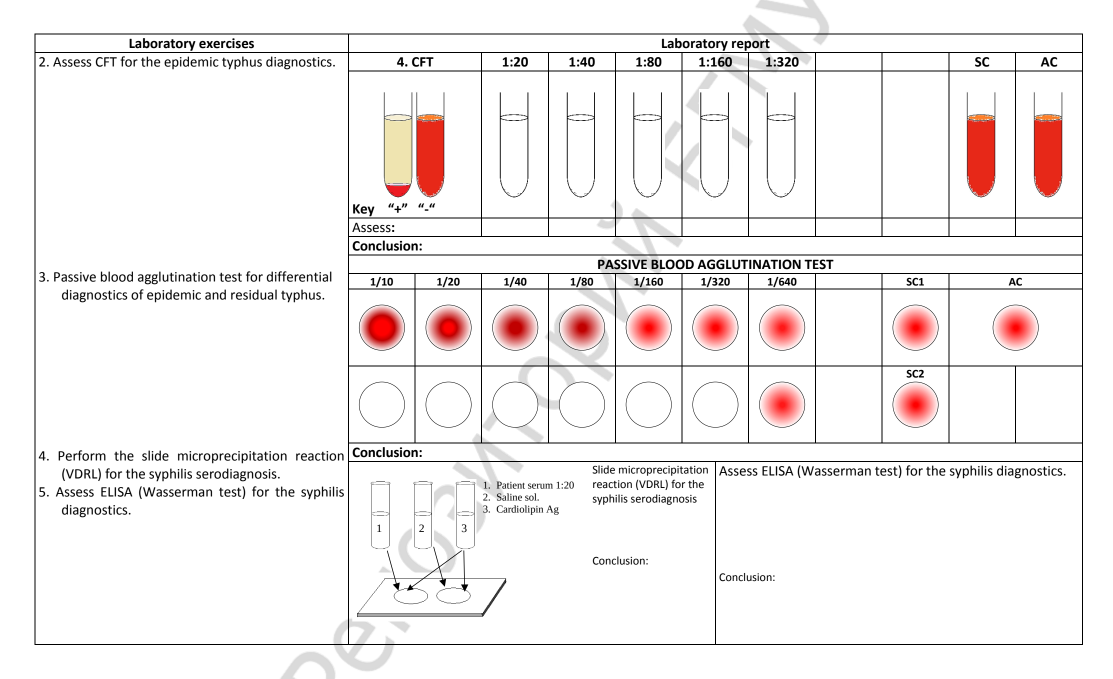
#### Laboratory work **Laboratory exercises** Laboratory report 1. Bacteriological diagnosis of diphtheria, the 3<sup>rd</sup> period: Smear\_ Smear \_\_\_\_\_ Smear Smear \_\_\_\_\_ - the assessment of Corynobacteria enzymatic activity, Stain \_\_\_ Stain Stain identification, conclusion. 2. Demonstration: Cord factor of *M.tuberculosis*, Ziehl-Neelsen staining; Actinomycetes spp., pure culture, Gram staining; M. leprae, Ziehl-Neelsen staining; - M.tuberculosis in sputum, Ziehl-Neelsen staining; Mycobacteria growth on nutrient media; Flotation method; determination of M. tuberculosis drug resistance.

## Practical class 3 (21). Methods of anaerobic infections microbiological diagnostics

#### Suggested reading for self-study: Laboratory Individual Total Oral quiz Tests work work results Anaerobes, classification, general characteristics. Non-spore anaerobes of the oral cavity (streptococci, bacteroides, fusobacteria, peptococci, peptostreptococci, veillonella, fusobacterial, leptotrichi, prevotella, bilophila), role in pathology. Causative agents of gas gangrene, tetanus, botulism, general characteristics. Pathogenicity factors, exotoxins. Signature of the tutor Clostridium role in dentistry. General principles and methods for anaerobic infections diagnosis. Molecular biological diagnostics - PCR. Principles of anaerobic infections therapy and prevention. Laboratory work **Laboratory exercises** Laboratory report 1. Bacteriological diagnosis of diphtheria, the 3<sup>rd</sup> Smear\_\_\_\_\_ Smear \_ Smear Smear \_\_\_\_\_ Stain Stain Stain Stain period: - the assessment of Corynobacteria enzymatic activity, identification, conclusion. 2. Demonstration: - Clostridium, Gram staining; - Bacteroides, Gram staining; veillonella spp., Gram staining; fusobacterial spp., Gram staining; - anaerobes growth on nutrient media.

## Practical class 4 (22). Microbiological diagnostics of diseases caused by Spirochetes, Rickettsia, Chlamydia, Mycoplasma

#### Suggested reading for self-study: Laboratory Individual Total Oral quiz Tests work work results Spirochetes, classification, general characteristics. Treponema. Systematics and general characteristics. Pathogenesis and immunity in syphilis, manifestations in the oral cavity. Methods of syphilis microbiological diagnosis. Principles of syphilis therapy and prevention. Fusospirochetosis pathogens. Leptospira, Borrelia. Role in human pathology. The causative agent of Lyme borreliosis. Signature of the tutor Rickettsiae, systematic position, classification, general characteristics, role in human pathology. Rickettsia typhii, pathogenesis, immunity and methods of microbiological diagnostics. Other pathogenic rickettsia. Chlamydia, systematics and general characteristics, role in human pathology. Mycoplasma, systematics and general characteristics, role in human pathology. Laboratory work **Laboratory exercises Laboratory report** 1. Demonstration: Smear \_\_\_\_\_\_ Smear 🔈 Smear \_\_\_\_\_ Smear Stain Stain Stain Stain Leptospires spp., dark field microscopy; - Borrelia recurentis in blood, Romanovsky-Giemsa staining; - Treponema spp. in dental plaque, Gram staining; - Treponema pallidum, pure culture; Romanovsky-Giemsa staining; - Chlamydia spp. in cell culture, Romanovsky-Giemsa staining; - R.prowazeki, pure culture, Zdrodovski staining; Wasserman test (ELISA). **Smear** Stain



## Practical class 5 (23). Test "Special bacteriology"

List of questions	~	Oral quiz	Script	Tests	Total results
List of questions					

- 1. Staphylococci, classification, general characteristics. Staphylococcal infections, pathogenesis and immunity. Role in in oral cavity pathology. Microbiological diagnosis. Principles of staphylococcal infections treatment and prevention.
- 2. Streptococci, classification, general characteristics, antigenic structure. Acute and chronic streptococcal 21. The causative agent of botulism, general characteristic. Pathogenesis, principles of botulism infections. Oral streptococci. The role of streptococci in oral pathology. Methods of streptococcal infections diagnostics. Principles of therapy and prophylaxis.
- 3. Classification of Neisseria. Meningococcus, general characteristics. Meningococcal infections, mechanisms of pathogenesis, immunity, methods of diagnosis, prevention.
- 4. Gonococci, general characteristics. Mechanisms of pathogenesis and immunity. Microbiological diagnosis of acute and chronic gonorrhea. Principles of therapy and prophylaxis. Gonorrheal stomatitis.
- 5. General characteristics of the family. Enterobacteriaceae.
- 6. General Principles of acute intestinal infections (AII) bacteriological diagnosis. E. coli, common characteristic. The biological role of Escherichia coli. Diseases caused by Escherichia.
- 7. Salmonella. General characteristics. Members of the genus. Diseases caused by Salmonella.
- 8. Pathogens of typhoid, paratyphoid A and B, general characteristic. Pathogenesis, immunity, prophylaxis. and methods of microbiological diagnosis of typhoid and paratyphoid.
- 9. The etiology of bacterial origin food poisoning and intoxication. Materials and methods of diagnosis.
- 10. Shigella. Classification. Characteristics. Pathogenesis, immunity of dysentery.
- 11. Klebsiella, general characteristics. Role in human pathology. Methods of klebsiellosis microbiological diagnostics.
- 12. Pseudomonas aeruginosa, general characteristics, pathogenicity factors. Role in human pathology.
- 13. C.diphtheria, general characteristics. Pathogenesis of diphtheria. Manifestation of diphtheria in oral cavity. Immunity in diphtheria. Methods of microbiological diagnostics, principles of diphtheria therapy and prevention.
- 14. The causative agent of whooping cough, general characteristics. Differentiation with parapertussis agent. Pathogenesis, immunity. Microbiological diagnosis, principles of pertussis treatment and prevention.
- 15. Actinomycetes, general characteristics. Role in the oral cavity pathology. Actinomycosis, characteristic of pathogen diagnostic techniques.
- 16. Classification of Mycobacteria. General characteristics of the tuberculosis causative agents. Pathogenesis, immunity, diagnostic, principles of tuberculosis therapy and prophylaxis. Manifestations of tuberculosis in the oral cavity.
- 17. Quarantine infection. Classification mode. Basic rules of infectious material sampling, sending and transportation. General principles of diagnosis.
- 18. V.cholera, general characteristics. Pathogenesis, immunity, principles of treatment and prevention.
- 19. Classification and general characteristics of anaerobes. Clostridia. Nonspore anaerobes. Role in the oral cavity pathology.

- 20. The causative agent of tetanus, general characteristics. Pathogenesis, immunity, principles of tetanus treatment and prevention. Gas gangrene pathogens, general characteristics. Pathogenesis, principles of gas gangrene treatment and prevention.
- prevention and therapy.
- 22. Methods of anaerobic infections diagnosis.
- 23. Classification and general characteristics of spirochetes. Borreliosis and leptospirosis agents.
- 24. Classification of treponemes and treponemal diseases. Characteristics of syphilis causative agent. Pathogenesis, immunity, principles of syphilis therapy and prophylaxis, manifestations in the oral cavity. Methods of syphilis diagnosis.
- 25. Oral spirochetes. Fusospirochaetosis.
- 26. Rickettsia. Role in human pathology. Pathogenesis, immunity, methods of typhus diagnosis.
- 27. Chlamydia. Role in human pathology. Pathogenesis, immunity, methods of diagnosis.
- 28. Mycoplasma. Role in human pathology. Pathogenesis, immunity, methods of diagnosis.

#### Practical skills:

- 1. Determine the morphology of Staphylococcus, pure culture, Gram stain.
- 2. Determine the morphology of Streptococcus, pure culture, Gram stain.
- 3. Determine the morphology of Gonococci in pus, Gram stain.
- 4. Determine the morphology of Enterobacteria, pure culture, Gram stain.
- 5. Determine the morphology of the mixture of S.aureus and Escherichia coli, Gram stain.
- 6. Determine the morphology of B.anthracis, pure culture, Gram stain.
- 7. Determine the morphology Vibrio, pure culture, Gram stain.
- 8. Determine the morphology of Brucella, a pure culture, Gram stain.
- 9. Determine the morphology Corynebacteria, pure culture, Leffler stain.
- 10. Determine the morphology of Klebsiella, pure culture, Hins-Burri stain.
- 11. Determine the morphology of Mycobacteria in sputum, Ziehl-Neelsen stain.
- 12. Determine the biochemical properties of enterobacteria on Kligler iron agar medium.

## **Practical class 6 (24).** Methods of investigations in virology. Bacteriophages

## Suggested reading for self-study:

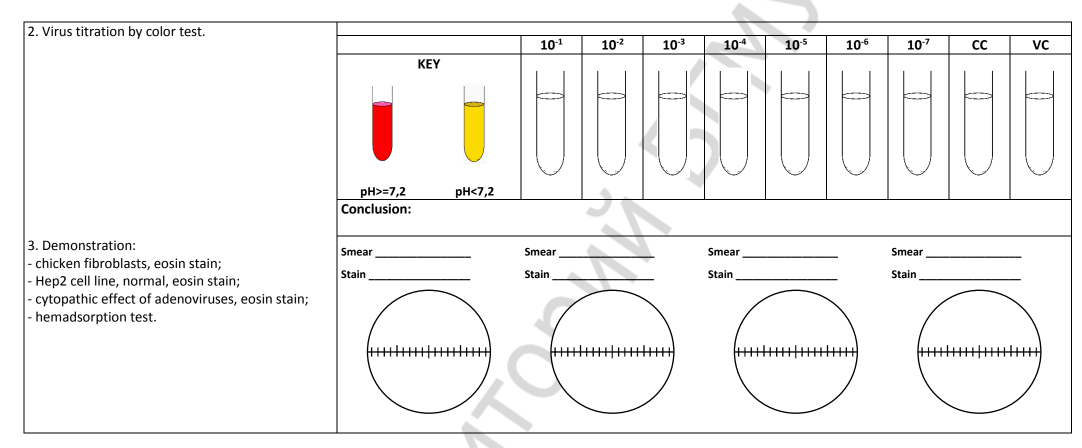
Viruses. Taxonomy and morphology of viruses. Mechanisms of reproduction. Strict parasitism and cytotropism of viruses.

The types of viral infection. The mechanisms of antiviral immunity. Principles for the prevention of viral infections in the dental practice. Methods of viral infections diagnostics. Culturing of viruses.

Viruses of bacteria (bacteriophages), characteristics of bacteriophages. Use of bacteriophages in medical Signature of the tutor practice.

Oral quiz	Laboratory work	Individual work	Tests	Total results	

	Laboratory work	
Laboratory exercises	Laboratory report	
1. Chicken embryo inoculation with influenza virus in allantois cavity.	1.Study the structure of hen embryo (8-11 days) 2. Examine hen embryo in ovoscope and determine the vitality signs: a) the dimensions of the embryo shape b) presence of the developed blood vessels pattern c) active mobility of the embryo d) mark the air cavity border 3. Set embryo on the egg rack and work with the shell as follows: a) 70% alcohol b) 5% iodine 4. Inoculate embryo as follows: a) flame scissors b) carefully pierce the shell for 3-5 mm above the air cavity border c) introduce 0,2 ml of viral material (live influenza vaccine) into the syringe d) put the needle into the embryo (25 mm) vertically and introduce the material. 5. Repeat the shell manipulations according to p.3. 6. Seal the shell with tape or melted wax. Mark the embryo (group number).  Inoculation of the Allantois cavity: 1. Use cotton wool and 70 percent alcohol to swab the eggs end to be inoculated. Allow the alcohol to evaporate. 2. Swab the eggshell punch with 70 percent of alcohol solution. Place used cotton wool in discard tray. 3. Pierce a hole in the end of the egg at the marked inoculation site. 4. Attach needle to 1 mL syringe. 5. Draw inoculum into 1 mL syringe. 6. Keeping the needle and syringe vertically, run through the eggshell hole approximately for 16 mm into the egg to reach the allantois cavity. 7. Inject 0.1 mL of inoculum into the egg. 8. Take the needle out from the egg. 9. Seal the hole in the shell with stationery tape or melted wax. 10. Discard the used needles and syringes. 11. Put the inoculated eggs into an incubator.	1 2  3 7  1. Shell membrane 2. Air sac 3. Chorioallantoic membrane 4. Allantois cavity 5. Amnion cavity 6. Yolk sac 7. Albumin 8. Extraembryonic cavity 9. Embryo



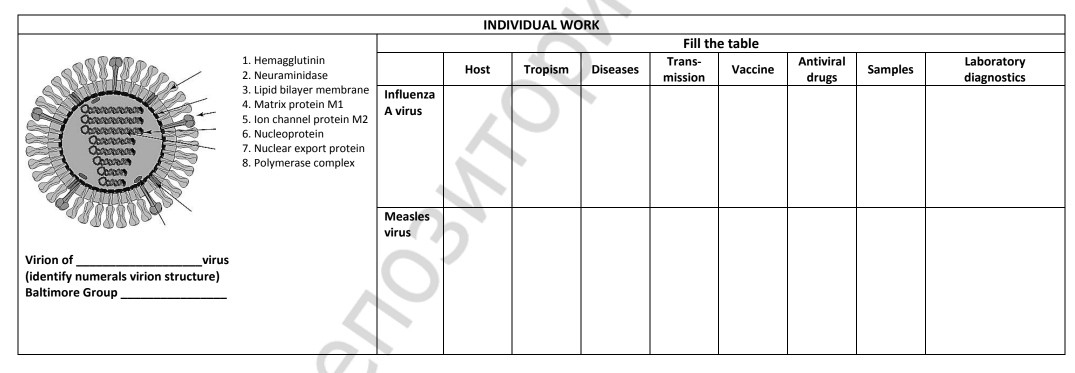
			INDIVIDU	IAL WORK						
	According to Baltimore classification, viruses are divided into the following seven classes (fill table)									
class	I	II (	III	IV	V	VI	VII			
Description of genome and replication strategy										
tip	T-C-A-G A-G-T-C	T-C-A-G	U-C-A-G A-G-U-C	U-C-A-G	U-C-A-G-	U-C-A-G↓↑	T-C-A-G ↓↑ A-G-T-C			

## **Practical class 7 (25).** Virology diagnostics of diseases caused by Orthomyxoviruses, Paramyxoviruses. Togaviruses

#### Suggested reading for self-study: Laboratory Individual Total Oral quiz Tests work work results Orthomyxoviruses. Taxonomy and characteristics of the family. Influenza viruses, morphology, antigenic structure and antigenic diversity (shift and drift) and its consequences. Methods for influenza diagnostics. Principles of therapy and prophylaxis. Paramyxoviruses. Taxonomy and characteristics of the family. Differentiation with Orthomyxoviruses, Signature of the tutor Parainfluenza viruses, Mumps virus, Morbilivirus, HRSV. Pathogenesis, immunity, specific prophylaxis. Rubella virus. General characteristics. Role in pathology. Manifestations of rubella in the maxillofacial region. Prevention of rubella. Laboratory work **Laboratory exercises** Laboratory report 1. Before autopsy embryo should be cooled for 2-3 hours at 4-6° C for blood vessels constriction. 1. Chicken embryo autopsy. 2. Virus indication by slide HT. 2. Treat the eggshell with 70%-alcohol and flamed. Repeat it once more. 3. Evaluation of HIT for influenzavirus 3. Open the shell by sterile scissors 2-3 mm above air sack border. Remove shell membrane and aspirate 1 ml of allantois identification. cavity liquid. 4. Amnion cavity liquid can also be taken (0,5-1,5 ml). 5. Remove an embryo on the Petri plate. Allantois membrane should be carefully examined by eyes. Usually influenza viruses produce no CPE. 6. Perform slide HT for virus indication 3 1 2 **SLIDE HT** Put two drops of 5% chicken erythrocytes | Smear | suspension onto glass slide. Add and mix one Stain\_ drop of allantois liquid (experiment) and saline (negative control) with each drop. The test is positive if flakes of erythrocytes are developed. The test is negative if erythrocytes remain in suspension after 5-7 min. 1. Allantois liquid. 2. Saline.

3. 5% chicken erythrocytes.

Laboratory work									
Laboratory exercises	Laboratory report								
4. Evaluation of HIT for influenza virus	L patient's virus	Anti H <sub>1</sub> N <sub>1</sub>	Anti H <sub>3</sub> N <sub>2</sub>	Anti H₅N₁	EC	VC	КантиС1	К <sub>анти</sub> С2	К <sub>анти</sub> СЗ
identification									
	D patient's virus								
	Conclusion:			1					



## Practical class 8 (26). Virologic diagnostics of diseases caused by picornaviruses and hepatitis viruses

#### Suggested reading for self-study: Laboratory Individual Total Oral quiz Tests work work results Picornaviruses. Characteristics of the family, importance for human pathology. Etiology, pathogenesis, immunity, diagnostics and immunoprophylaxis of poliomyelitis. Coxsackieviruses and ECHOviruses. Stomatitis in diseases caused by RNA-viruses. Hepatitis viruses A, B, C, D, E. Taxonomy and characteristics, role in human pathology. Pathogenesis and immunity in Signature of the tutor hepatitis B. Laboratory diagnostics. Specific and non-specific prophylaxis in dentistry. Laboratory work **Laboratory exercises** Laboratory report 1. Performance of ELISA for VHC diagnostics. Antibodies from patients' serum bind to b) put 100 µl of control sera and samples C- - negative control; 2 1 C+ - positive control; recombinant antigens adsorbed on the well according to the plate layout's) close strip Core C- $X_1$ X<sub>1</sub>- serum patient 1: of a plate. Specific immune complexes then with adhesive tape and incubate for 1 hour $X_2$ – serum patient 2; C-Χı NS₃ detected by conjugate antibody-enzyme and at 37°C; The protocol is based on the commercial ELISA kit for VHC «1», «2» - plate vertical rows: respective enzymatic reaction. Colored d) wash wells 5 times; diagnostics "RecombiBest anti-HCV" by VectorBest, RF. $NS_4$ C C- $X_1$ A-H - plate horizontal rows; product developed is measured by ELISA e) put 100 μл of conjugate in each well; The method reveals antibodies (IgG and IgM) to HCV $NS_5$ D C- $X_1$ reader. f) seal strip with tape and incubate for 30 antigens. Reaction scheme: min at 37°C; Core Ε C+ $X_2$ a) HCV antigens are adsorbed on the strip g) wash 5 times; F C+ $X_2$ NS₃ **Card STATEMENT** wells as follows: rows A, E - core h) put 100 µl of substrate in each well; $NS_4$ G C+ $X_2$ rows B,F-NS3 i) incubate for 30 min at 37°C; rows C,G - NS4 i) put 50 µl of stop solution in each well; н C+ $X_2$ NS<sub>5</sub> rows D. H - NS5 k) measure the plate by ELISA reader: I) evaluate results. **Antigens** Row OD OD **Cut-off** 1. Test results validation: PI(core-Ag) = OD sample(core)/ Cut-off(core-Ag) = Results control probe PI(NS3-Ag) = OD sample (NS3)/Cut-off(NS3-Ag) = Negative control OD < 0,2 Core Α Mean negative control OD = PI(NS4-Ag) = OD sample (NS3)/Cut-off(NS4-Ag) = В NS₃ Mean positive control OD >0,8 PI(NS5-Ag) = OD sample (NS3)/Cut-off(NS5-Ag) = $NS_4$ С Mean positive control OD = 4. Results evaluation: D $NS_5$ 2. Cut-off level for each antigen: a) If PI less than 1, sample is considered negative; E Core Cut-off (core-Ag) = NC ODO(core) + 0.2 =б) the results are considered positive if IP exceeds 1 for: F NS₃ Cut-off(NS3-Ag) = NCOD(NS3) + 0.2 =core-Ag G NS₄ Cut-off(NS4-Ag) = NCOD(NS4) + 0.2 =any two antigens NS<sub>5</sub> Н Cut-off(NS5-Ag) = NCOD(NS5) + 0.2 =B) result is considered uncertain if IP exceeds 1 for one 3. Positivity index determination for each antigen: nonstructural protein only.

2. Neutralization test on cell culture in paired sera for poliomyelitis serodiagnostics accounting of reaction.

NT IN PAIRED SERA FOR POLIOMYELITIS

SERODIAGNOSTICS

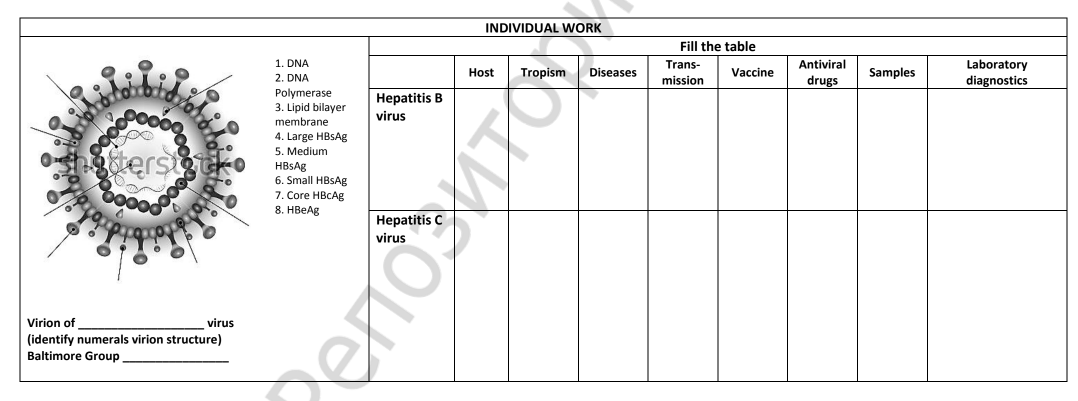
1/10 1/20 1/40 1/80 1/160 SC1 VC CC

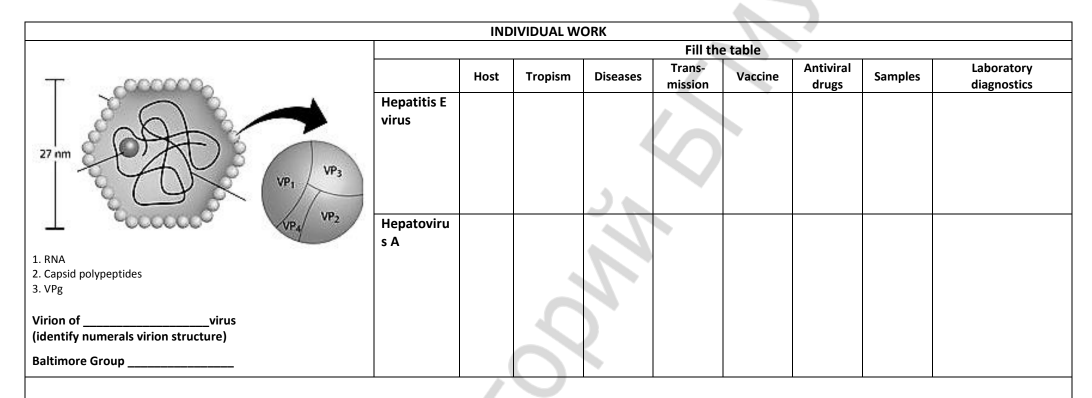
Patient Z' serum

Patient X' serum

Conclusion:

Signature of the tutor\_\_\_\_\_\_\_





Virus	Family-Genus-Species	Genome	The structure, size of the virion, nm	High-risk group
HAV	Picornaviridae — Hepatovirus - Hepatitis A virus			
HBV	Hepadnaviridae – Orthohepadnavirus - Hepatitis B virus			
HCV	Flaviviridae – Hepacivirus - Hepatitis C virus			
HDV	Unassigned - Deltavirus - Hepatitis delta virus			
HEV	Hepeviridae- Hepevirus - Hepatitis E virus			

## Practical class 9 (27). Methods of diagnostics for diseases caused by Retroviruses and Rabdoviruses

Retroviruses. Taxonomy and characteristics of the family. Human immunodeficiency virus (HIV-1, HIV-2). Pathogenesis.

AIDS-associated diseases. Manifestations in the oral cavity. HIV diagnostics, prophylaxis, treatment. HIV in Belarus.

Suggested reading for self-study:

	cteristics of rabdoviruses. Pathogenesis, immunity and specific prophylaxis of rab	ies. Sign	nature of t	he tutor					
	Laboratory work								
Laboratory exercises Laboratory report									
1. Demonstration:  - Negry bodies in mouse brain homogenate, Muromtcev stain.	Smear Stain								
Practical class 10 (28). Method: Suggested reading for self-study:	s of diagnostics for diseases caused by herpes- and adend		es disea	SES IN O		ty <sub>Total</sub>			
Herpes viruses. Taxonomy and family immunity, diagnostics, chemo and immunother of chicken pox and herpes zoster. Cytomegal properties, role in human pathology. Infection Immunity, diagnosis, chemotherapy and immunity.	or characteristics. HSV-1, HSV-2, properties, role in human pathology, pathogenesis, papy. Herpetic stomatitis, keratoconjunctivitis, facial skin lesions and red lip rims. A virus ovirus, properties, forms of infection. Cytomegalovirus parotitis. Epstein-Barr virus, pus mononucleosis. Herpesviruses of human 6, 7, 8 types, role in human pathology. sotherapy of herpetic infections.	Oral quiz Signature	of the tute	work  or	Tests	results			
Adenoviruses. Characteristics. Human a	adenoviruses. Virions structures, pathogenesis, immunity, laboratory diagnostics.								
Laboratory exercises	Laboratory work  Laboratory report								
1. Demonstration: - CPE of adenoviruses.	SmearStain	111111							

Oral

quiz

Laboratory

work

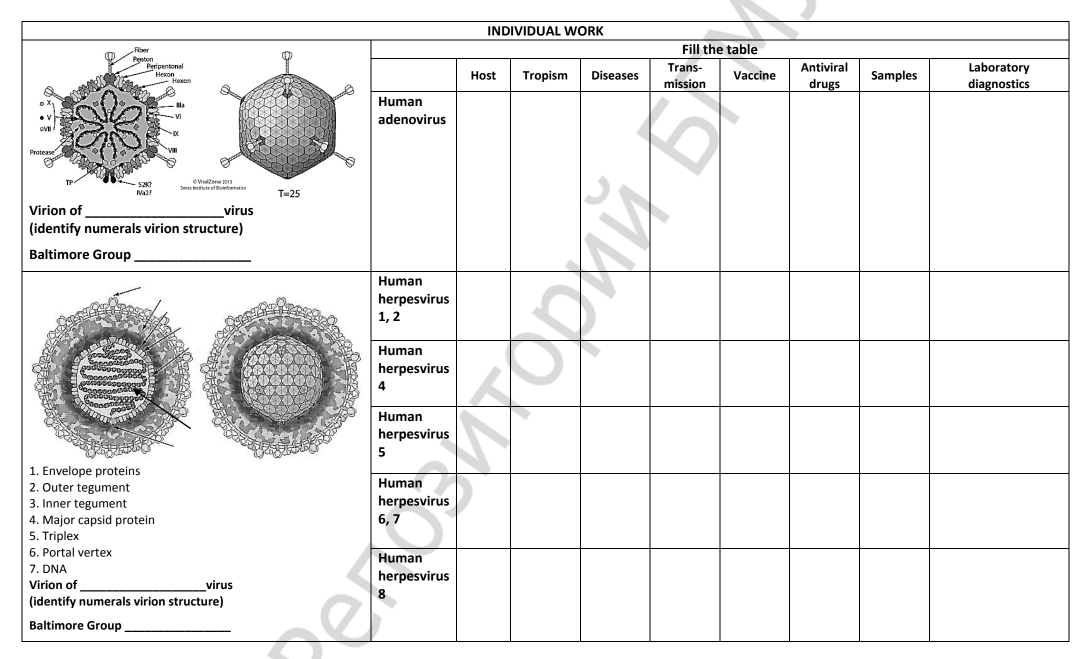
Individual

work

Total

results

Tests



Practical class 11 (29). Dental microbiology. Methods of oral cavity normal flora investigation. Etiology and pathogenesis of caries

Individual

Total

Tests

Laboratory

Oral quiz

Suggested reading for self-study:

Dental microbiology, goals and objective	ves. Normal microflora of the oral cavity, characteris	stic. Ontogeny	of normal microflora.	Oral quiz	work	work	rests	results
	n the composition of the oral cavity microflora (which							
soft tissue, contact with alien microorganisms,	, diet and oral hygiene). The value of normal microflo	ra. Methods of	study.					
Dysbacteriosis of the mouth, causes, di	agnostic methods.	4		ianatura	of the tut	or		
The etiology of caries. Causal important	ce of microorganisms. S. mutans, properties. Subsidi	ary germs. Path	nogenesis. Conditions	ignature	or the tu	.01		
	laxis and therapy of caries. Rules and methods of sa	mpling for the	study of caries ogenic					
microflora. Criteria for assessment of the isola	ted microorganisms etiological significance.							
	Laboratory	work						
Laboratory exercises		Lab	oratory report					
1. Perform isolation of normal flora from	- Divide agar plates into four sections with a marking pen	or pencil. Mark e	each section with 1, 2, 3, 4.			Blood aga	MacCo	nkey agai
mucus of oral cavity membrane surfaces	- Mark each plate with group number and your name.		*		1.			
to gain the microorganisms diversity	- Add sterile isotonic solution to the Petri dish with sterile			·	. 2			
understanding at these body locations	<ul> <li>Use flamed forceps to cover the squares of the various (saliva, lips, gum, mucus membranes of tong, cheeks) witl</li> </ul>			investigate	° 3●		1 /	
and exclude/confirm dysbacteriosis.	- Put the squares of filter paper for 60 sec on the surface					1		
	- Fill in the table with the sites in which the microbial flora			C for 24-48				
	hours.		1					
2. Register the results of experiment on	Results of registration of dysbacteriosis:	Body site	1 -	2 -		3	•	
normal flora isolation from mucus		Amount of						
membrane surfaces, Gram stain different		colonies						
types of colonies, explore under	Conclusion:	and their						
microscope, complete the report. (The		description						
task will be given at the next lesson).		description						
	3 Smear 1 -	Ct.:	Smear	Smear		C m	near	
3. Prepare heat-fixed smear from dental	3 Smear 1 - Stain 2 -	Gram stain	Stain	Stain			ieai ain	
plaque, Gram stain, explore under	3 -			.				
microscope, complete the report.	4-							
	5-							
4. Demonstration:	6-		/	\		\	•	\
- slide with dental plaque, Gram stain;	<del>                                    </del>		<del>                                   </del>	+)   (++++	<del>                                      </del>	<del>                                      </del>	+++++++++	<del>+++++++</del> )
- methods for detection of pathogenicity	8-			<i> </i>		<i> </i>		
factors (capsule, hemolysins, lecithinase,	9-			′   \				
cougulase).	10-			\		/		
<b>0</b> /								

## Practical class 12 (30). Dental microbiology. Methods of oral cavity immunity factors investigation

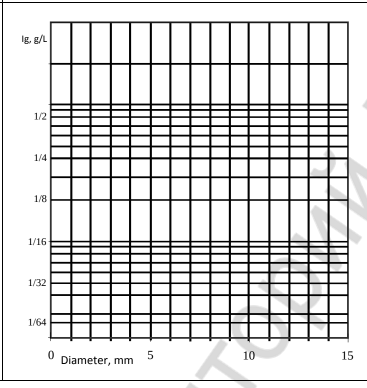
#### Suggested reading for self-study: Laboratory Individual Total Oral quiz Tests work work results Immune and non-immune mechanisms in the oral cavity (natural and acquired). Protective mechanisms of saliva, mucous membranes of the oral cavity, enamel, dentin and pulp of the teeth. Importance of phagocytosis. Immunoglobulins of the oral cavity. Secretory immunoglobulin A. Signature of the tutor Cell-mediated immunity. Mechanisms of antibacterial and antiviral immunity in the oral cavity. Laboratory work **Laboratory exercises Laboratory report** 1. Determine the content of lysozyme in Smear 2 4 Saliva, 1-1,5 ml Stain saliva. - collect 1-1,5 ml saliva in a tube. - mark the Petri dish with the ready-hole seeded Micrococcus lysodeikticus, according to the scheme. - pipette in the wells of the lysozyme appropriate dilutions 50 µl (from low 6.25 12.50 25.00 50.00 to high concentration). mcg/ml mcg/ml mcg/ml mcg/ml - in the central well of the test add 50 µl Standard curve Standard of Zone of inhibition, of saliva. Lysozyme, mcg/ml diameter in mm - incubate the plate for 24 hours. 6,25 (1/8) construct a calibration curve and 12,50 (1/4) determine the concentration of 25,00 (1/2) lysozyme in your sample. - compare with the standard and make a 50,00 (1) conclusion. X sample **Conclusion:** <sup>0</sup> Diameter, mm <sup>5</sup> 10 15

# Laboratory exercises 2. Determine the IgA concentration in saliva by Manchini method (simple radial

sIgA standard – 2,0 g per liter.

gel immunodiffusion).

3. Register the experiment results on normal flora isolation from mucus membrane surfaces, Gram stain different types of colonies, explore under the microscope, complete the report.



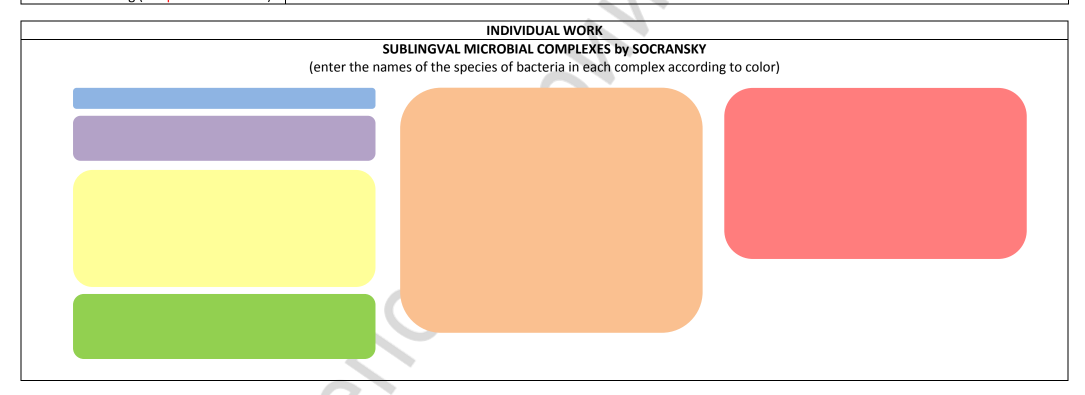
	Standa	rt curve				
Standard sIgA = 2 g/l						
	titer	concentrtion,	Diameter, mm			
	-	g/l				
Point 1	1	2,000				
Point 2	1/2	1,000				
Point 3	1/4	0,500				
Point 4	1/8	0,250				
Point 5	1/16	0,125				
X-sample						

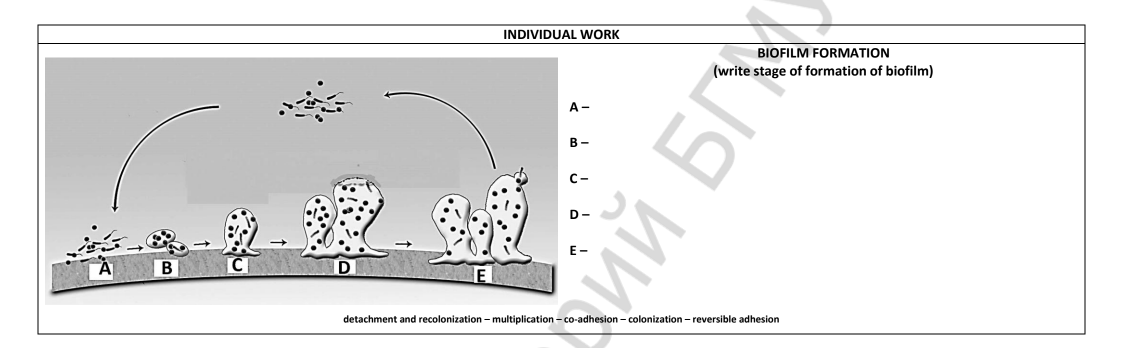
As a normal slgA ranger is 0,3-0,4 g/l

Conclusion:

## Practical class 13 (31). Dental microbiology. Microbiology of periodontal and peri-implantitis diseases

Suggested reading for self-study:	sted reading for self-study:		Laboratory work	Individual	Tests	Total
Plaque: stages of formation, microo	Plaque: stages of formation, microorganisms-colonizers. Plaque as a biofilm. Periodontal diseases: classification,			work		results
etiology, risk factors. Theories of the pathog						
mechanisms of invasion and persistence. Microbial complexes (Socransky, 1998). Immune mechanisms in diseases of the tissues of the periodioth. Principles of prevention and treatment of periodontitis. Dynamics of microflora with successful						
and complicated dental implantation.						
Laboratory work						
Laboratory exercises	Laboratory report					
1. Determine the content of lysozyme in						
saliva – ending (see practical class 12).						





## **Practical class 14 (32).** Dental microbiology. Methods of microbiological diagnostics of stomatitis. Microbiological diagnostics of fungal infections

## Suggested reading for self-study:

Inflammatory diseases of the oral mucosa. Bacterial stomatitis: specific (gonococcal, typhoid fever, antrax stomatitis, manifestations in the oral cavity of syphilis, tuberculosis, actinomycosis, scarlet fever) and nonspecific. Viral stomatitis.

× /	Oral quiz	Laboratory work	Individual work	Tests	Total results	
` .						

Signature of the tutor

Classification and general characteristics of fungi. Classification of mycosis. Candida, general characteristics. Role in human pathology. Soor. General principles of fungal infections diagnostics.

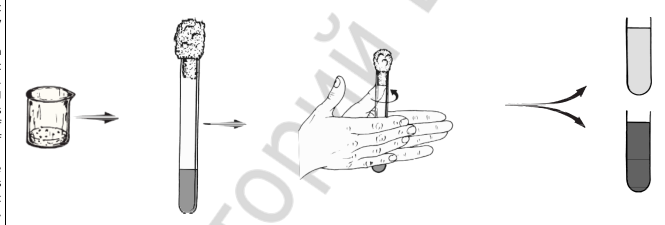
#### Laboratory work **Laboratory exercises** Laboratory report 1. Research of the sample of the patient's Sample of pus Serial dilution of the sample Blood sample examination, Smear 2.1 pus with an abscess subcutaneous tissue from 1<sup>st</sup> period Stain of maxillofacial area, the 1<sup>st</sup> period: microscopy of pus (smear, Gram stain); preparation of inverse hundredfold dilutions material in sterile saline (1:100; 1:10000; 1:1000000); quantitative (50 mcl) streak respective **Burnt wound** sectors dilutions of pus on solid nutrient 10-2 10-4 10-6 media (MSA, Endo, blood agar, NA with 0,1 ml 0,1 ml furagin) depending on the results of 10 ml:60 ml 9.9 ml ≻9.9 ml microscopy. Saline sol. Saline sol. Saline sol. 2. Research of the blood sample from the patient with stomatogenic sepsis, the 1st period: Steak respective sector with 0,05 ml (1 drop) Smear 3.1 microscopy of blood, smear "thick drop". Stain Medium methylene blue stain or Romanovsky; - Crop material in the liquid medium of 10-4 the primary crop (enrichment) in a ratio 10<sup>-6</sup> of 1: 10-60: Incubation of cultivation in an incubator at 37 $^{\circ}$ C - 18-24 hours and up to 14 days. All inoculations are placed in a incubator for 24 hours, then transferred Blood agar Levin Nutrient agar with to a refrigerator for 14 days.

#### **Laboratory exercises**

- 3. Snyder's caries susceptibility test
- formation of tooth decay (dental caries) acid by bacteria (Streptococcus mutans and Of the various methods that have been devised to determine one's susceptibility to tooth decay, M. L. Snyder's caries susceptibility test is a relatively simple test that has been shown to have a high reliability correlation.
- This method relies on the rapidity of organisms in saliva to lower the pH in the medium that contains 2% dextrose (Snyder test agar). Since decalcification of enamel begins at pH of 5.5, and progresses rapidly as the pH is lowered to 4.4 and less, the demonstration of pH lowering becomes evidence of susceptibility to caries.
- To indicate the presence of acid production in the medium, the indicator bromcresol green is incorporated in it. This indicator is green at pH 4.8 and becomes yellow at pH 4.4, remaining yellow below 4.4.
- Figure illustrates the procedure that is used in the Snyder caries susceptibility test. Note that 0.2ml of saliva is added to the tube of liquefied Snyder test agar (50° C) and mixed well by rotating the tube between the palms of both hands. After the medium has solidified, the tube is incubated at 37° C for a 1 ml pipette period of 24-72 hours. If the medium turns vellow in 24-48 hours, the individual is said to be susceptible to caries.
- Although we will be performing this test only once, it should be noted that test reliability is enhanced by performing the test on three consecutive days at the same time each day. If the test is performed correctly after tooth

- 1. Liquefy a tube of Snyder test agar and cool it to 50° C.
- The degradation of enamel and dentin in the | 2. After allowing a piece of paraffin to soften under the tongue for a few | tube vigorously between the palms of the hands. minutes, start chewing it. Chew it for 3 minutes, moving it from one side of the the small sterile beaker.
  - others) in the presence of sucrose high levels. 3. Vigorously shake the sample in the beaker from side to side for 30 seconds The degree of caries susceptibility is determined from the table below. to disperse the organisms.
    - 4. With a 1 ml pipette transfer 0.2 ml of saliva to the tube of agar. Do not allow the pipette to touch the side of the tube or agar.

- **Laboratory report** 5. Before the medium solidifies, mix the contents of the tube by rotating the
  - 6. Write your name on a gummed label and attach it to the tube.
- occurs as a result of the production of lactic mouth to the other. Do not swallow the saliva. As it accumulates, deposit it in 7. Incubate the tube at 37° C. Examine the tube every 24 hours to see if the bromcresol green indicator has changed to yellow. If it has, the test is positive.
  - 8. Record your results on the Laboratory Report.



#### Materials:

- 1 tube of Snyder test agar (5 ml in 15 mm dia tube)
- 1 30 ml sterile beaker
- 1 piece of paraffin (1/4" 1/4" 1/8")
- 1 gummed label

CARIES SUSCEPTIBILITY	MEDIUM TURNS YELLOW IN:				
	24 HOURS	48 HOURS	72 HOURS		
Marked	Positive				
Moderate	Negative	Positive			
Slight	Negative	Negative	Positive		
Negative	Negative	Negative	Negative		

## Practical class 15 (33). Test "General and special virology. Dental microbiology"

List of questions	Oral quiz	Script	Tests	Total results

- 1. Virology, tasks and methodologies. The systematic position and classification of viruses.
- 2. Forms of viruses existence. The morphology of virions. The interaction of viruses with susceptible cells.
- 3. Features of infection and immunity in viral infections.
- 4. Methods of virus cultivation (cell culture, chicken embryo, laboratory animals).
- 5. General principles of viral infections diagnostics.
- 6. Influenza viruses. General characteristics. Pathogenesis, specific and non-specific treatment and prevention, influenza laboratory diagnosis. Manifestations in the oral cavity.
- 7. Paramyxoviruses, general characteristics. Mumps virus, respiratory-syncytial virus, measles virus, parainfluenza viruses. Manifestations in the oral cavity.
- 8. Enteroviruses, general characteristics, role in human pathology. Poliovirus, pathogenesis and laboratory diagnostics, specific prevention. Manifestations of enteroviruses infection in oral cavity.
- 9. Classification of hepatitis viruses. Characterization of hepatitis A, B, C virus. Pathogenesis, immunity, laboratory diagnosis, prevention.
- 10. Retroviruses. Human immunodeficiency virus (HIV-1, HIV-2). Pathogenesis. AIDS-associated diseases in dentistry. HIV diagnostics, prophylaxis.
- 11. Adenoviruses, general characteristics. Pathogenesis, laboratory diagnostics of adenoviral infections. Manifestations in oral cavity.
- 12. Herpes viruses. Classification. General characteristics, disease. Herpetic stomatitis.
- 13. Bacterial viruses (bacteriophages), properties, classification. The practical use of bacteriophages.
- 14. The microflora of the oral cavity (indigenous, transient). Ontogeny of normal oral flora.
- 15. Representatives of the normal oral flora: Gram-positive and Gram -negative cocci (streptococci, peptostreptococci, staphylococci, veillonella, Neisseria), their role.
- 16. Representatives of the normal oral flora: Gram-positive (propionjbacterium, lactobacillus, actinomyces, corynebacterium) and Gram-negative rods (bacteroides, prevotella, porphyromonas, fusobacterium, leptotrichia), their role
- 17. Representatives of the normal oral flora spiralshaped bacteria (vibrio, wolinella, centipedia, selenomonas, campylobacter, spirochetes), mycoplasma, protozoa, fungi, and their role.
- 18. Microflora of specific areas of the mouth: saliva, dorsum of the tongue, dental pocket, mucous membranes Methods of study of oral microflora.
- 19. Influence of environmental factors and physiological features on oral flora. The role of the oral cavity normal microflora (positive and negative). Disbacteriosis of the oral cavity: causes, outcome, prevention, principles of correction.
- 20. Antigens and the immune system of the oral cavity. Citrullinated antigens. Immune mechanisms in the oral cavity. Antimicrobial factors of saliva: defensins, cathelicidin, mucins, histatin, statherin, cystatins. Proinflammatory cytokines.
- 21. Nonspecific mechanisms of defense of the mucous membranes, saliva, gingival fluid, tooth enamel, normal microflora's.
- 22. Factors and mechanisms of acquired immunity of oral cavity. Local Immunity of the oral cavity. Immunological aspects of relationship of inflammatory periodontal diseases, cardiovascular and rheumatic diseases.
- 23. Types of inflammatory processes of the oral cavity, their characteristics. Cytokines of early and late phase of inflammation: cell producers, properties. Methods of cytokines detection: obtaining of specimens, storage, methods of determination (ELISA, genetic).

- 24. The etiology of dental caries. Features of cariogenic microorganisms. Cariogenic streptococci. Characteristics of *S.mutans*. Characteristics of lactobacilli. Associative (additional) microorganisms. The role of the microorganism in the development of caries.
- 25. Cariogenesis: mechanisms of streptococci adhesion to teeth and their role in dental plaque formation. Role of glucans and their characteristics. Factors responsible for caries development. Resistance to caries. Prevention of dental caries.
- 26. Odontogenic infections: etiology, types. The role of microorganisms in the etiology and pathogenesis of gingivitis. Dynamics of the microflora of implants in case of successful implantation and complicated.
- 27. The role of microorganisms in the pathogenesis of pulpitis, acute and chronic periodontitis ray, periostitis, osteomyelitis, abscesses and soft tissue abscesses.
- 28. Periodontal diseases: classification, risk factors. General properties of periodontopathogenic microorganisms. Red complex microorganisms: *Porphyromonas gingivalis, Tannerella forsythia, Treponema denticola*. Characterization, pathogenicity factors and their role in the pathogenesis of periodontitis. Characteristics of *Aggregatibacter actinomycetemcomitans* and role in the development of aggressive periodontitis.
- 29. Dental Plaque: microflora, formation stages. The role of dental plaque in the development of periodontitis. Microorganisms of orange and yellow complexes, their role in the development of periodontal disease. Plaque as a biofilm. The role of quorum sensing factors in the formation of plaque. New approaches to reduce the bioburden of plaque.
- 30. Immune mechanisms in the development of periodontal diseases. Factors contributinging invasion of microorganisms. Mechanisms to protect tissues from microbial invasion. Principles of prevention and treatment of periodontitis
- 31. The role of microorganisms in the formation of dental calculus. Pathogenesis of the carie dental calculus formation.
- 32. Inflammatory diseases of the oral mucosa: classification, the role of microorganisms in their development. Specific and nonspecific stomatitis.
- 33. Stomatitis caused by obligate pathogens and opportunistic bacteria.
- 34. Fusospirochetal diseases: etiology, characteristics of pathogens, pathogenesis, clinical forms.
- 35. Actinomyces spp.: systematics, classification, characteristics, antigenic structure, factors of pathogeneity. Cervico-maxillo-facial actinomycosis: pathogenesis, immunity, microbiological diagnosis, prevention.
- 36. Viral stomatitis.
- 37. Candida: systematics, properties, pathogenicity factors. Candidosis: factors responsible for the development, methods of diagnosis and prevention.
- 38. Methods of studying the normal oral flora. Methods of sampling for dental diseases diagnosis.
- 39. Manifestations of allergic and immunodeficiency conditions in the oral cavity. Recurrent viral aphthous stomatitis.
- 40. Types and etiology of stomatogenic infections.
- 41. Dental Clinical Microbiology. Opportunistic pathogens. Specific features opportunistic pathogens and infections caused by them. Specific features of pathogenesis and diagnosis of opportunistic diseases. Criteria of Etiological significance of isolated bacteria from a specimen.

**Practical class 16 (34).** Dental microbiology. Method of microflora investigation in diseases of the teeth and oral cavity soft tissues

#### Suggested reading for self-study: Individual Laboratory Total Oral quiz Tests work work results Odontogenic inflammation. Microflora, pathogenesis, microbiological diagnosis of pulpitis, periodontitis, periodottitis, osteomyelitis, odontogenic abscesses and phlegmon. Purulent-inflammatory dental diseases of soft tissues and bones of the maxillofacial area. Pathogens, pathogenesis, Signature of the tutor methods of microbiological diagnostics (material for research, rules and methods of sampling, a scheme for bacteriological examination of pus, criteria for the etiological role of isolated microorganisms). Determination of sensitivity to antibiotics, Dental sepsis. Pathogens, methods of microbiological diagnosis. Laboratory work Laboratory report **Laboratory exercises** 1. Research of the sample of the Colonies Smear Medium Medium characteristics Stain patient's pus with an abscess Shape subcutaneous tissue of maxillofacial Size area, 2<sup>nd</sup> period: Surface microscopy of slides prepared from all Edge types of colonies; Color the study of microbial growth on the Consistency media; Transparency determination of the pathogen **Determination of CFU** Coagulase test Oxidase test quantity per ml/g (CFU) of the sample Calculation of bacteria quality per ml/g of the Sample control with formula: sample: oxidase test; $N_{(CFU/mI)} = n \times 20 \times 10^x$ coagulase test; n – colonies quantity in respective sector, Sample Conclusion: seeding the culture for pure 20 - conversion factor for 1 ml, accumulation biochemical and 10x – the degree of the sample dilution. identification, incubation in incubator at 37 °C - 18-24 hours. $N_{(CFU/mI)} =$ control Stabilized rabbit plasm: 37 °C - 2, 4, 24 h 2. Research of the blood sample from the patient with stomatogenic sepsis, the 2<sup>nd</sup> period: - the study of microbial growth on the media: - microscopy of slides prepared from the media; - seeding on the blood and Yolk-salt agar for the pure culture.

Laboratory exercises		Lab	oratory report			
3. Research of the sample of the patient's		-	Antibiotic	Diamo	eter of inhibitio	n zones (mm)
pus with an abscess subcutaneous tissue		Smear	- Antibiotic	resistant		susceptible
of maxillofacial area, 3 <sup>rd</sup> period ( <i>The</i>		Stain		Staphylococo	us spp.	
•			Penicillin	≤28		≥29
task will be given at the next lesson):			Oxacillin CNS	≤17		≥18
- microscopy of slides prepared from pure			S.aureus	≤10 ≤13		≥13
culture;			Canamycine Gentamicin	≤13 ≤12		≥18 ≥15
- the study of microbial growth on the media;		/ \	Ciprofloxacin	≤15		≥21
- seeding the pure culture for accumulation		<del>(++++++++++++++++++++++++++++++++++++</del>	Tetracycline	≤14		≥19
and biochemical identification, incubation		\	Erythromycine	≥23		≥23
in an incubator at 37 °C - 18-24 hours;			Lincomycine	≤13		≥21
- seeding the pure culture for determination			Chloramphenicol	<17		≥18
	1			Enterobacteria	acea spp.	
of antibiotic resistance.			Ampicillin	≤13		≥17
			Cefazolin	≤14		≥18
			Cefotaxime	≤14		≥23
			Canamycine	≤13		≥18
			Gentamicin	≤12		≥15
			Ciprofloxacin	≤15		≥21
		4. 1	Lomefloxacin	≤18		≥22
4. Research of the sample of the patient's			Tetracycline	≤14		≥19
pus with an abscess subcutaneous tissue			Doxicycline	≤12		≥16
of maxillofacial area, 4 <sup>th</sup> period ( <b>The</b>			Chloramphenicol	≤12		≥18
task will be given at the next lesson):			A Athair Athai	antibioticgr		
- microscopy of slides prepared from pure		Smear	Antibiotic –	Diameter of inhibiti	on zone, mm	Interpretation of res
culture;		Stain				<del>                                     </del>
- the study of microbial growth on the media;		Stanii	_			
- determination of antibiotic resistance;						
<ul> <li>conclusion: identification and typing results,</li> </ul>	h n n n n n 1					
antibioticgramm.						
antibioticgramm.		\				
		<del>(++++++++++++++++++++++++++++++++++++</del>				
			DD	NA		
			==		tion	
			- make standard inoculum with saline solution			
			(0,5 unit MacFarlane); - microscopy of slides prepared from inoculum			
				repared from mocu	iuiii //	
	Conclusion:	culture)				
			- seeding of 1,0 ml of inoculum on MH agar; - incubation 18-20 hours 35°C.			8 6
			- incubation 18-20 hours	33°C.		8
	- V 1					
<u> </u>						

Practical class 17 (35). Clinical microbiology. Microbiological diagnostics of purulent infections of bronchi and lungs. Hospital-acquired infection Suggested reading for self-study: Laboratory Individual

Suggested reading for self-study:				Oral quiz	Laboratory	Individual	Tests	Total
Dental bronchopulmonary diseases.	Pathogens. Pathogenesis. Conditi	ions of occurrence. Me	ethods of microbiologica	ıl Granqaız	work	work	16363	results
diagnosis (materials for research, rules and	methods of sampling, a scheme for	or bacteriological sputu	m examination, bronchia	ıl				
washings, criteria for the etiological role of iso	lated microorganisms).			Signatura	of the tute	~ u		
Determination of sensitivity to antibio	otics.			Signature	or the tutt	JI		
Nosocomial infections. Pathogens, fe	·	, principles of diagnosis	. Anti-epidemic regime i	n				
dental practice. Principles of microbiological di	lagnosis. Prevention.							
		Laboratory work						
Laboratory exercises			Laboratory report					
1. Research of the blood sample from the	Blood agar Y:	SA MH	agar Co	oagulase test		Gluco	se and man	nitol
patient with stomatogenic sepsis, the 3 <sup>rd</sup>			Ex	p Contro	l	fermen	tation (anae	robic)
period:			0\					
the study of microbial growth on the	1 / /	1 / 0	<u> </u>		>			
medium;		) \	0 )					
microscopy of slides prepared from all types								
of colonies;								
oxidase test;				八八	)	/		
coagulase test;	Hemolyses Lecithinas	se Kirby-Bau	ier method Stabilized rab	bit plasm: 37 °	C -		0 0	
seeding the pure culture for accumulation and biochemical identification, incubation			2, 4, 24 h					
in an incubator at 37 °C - 18-24 hours.		Colonies	8.0 - di	0.012		Conclusion	:	
incubation at 27 °C - 18-24 hours	Smear	characteristics	Medium	Medium				
incubation at 57°C - 10-24 nours.	Stain	Shape						
2. Research of the blood sample from the								
patient with stomatogenic sepsis, the 4 <sup>th</sup>		Size						
period:		Surface						
the study of tests used for identification of		Juliace						
cultures and antimicrobial sensitivity level		Edge						
in DDM.		Color						
		_						
		Consistency						
		Transparency						

## Exam' questions for the dental faculty students

#### List of questions

- 1. Microbiology: definition, area and fields of microbiology. Objects and methods of research. Dental microbiology: goals, objectives, role in the dentist's practice.
- 2. Milestones (periods) in microbiology. Work of L.Pasteur, R.Koh, I.I.Mechnikov. Evolution of microorganisms and infectious diseases.
- 3. Common with other organisms and the unique features of microorganisms. Principles of systematics of microorganisms. Classification and nomenclature of microorganisms. The term of "species" in bacteria: group of traits for species identification (criteria for speciation).
- 4. Morphology of bacteria. Basic morphological forms of bacteria. The structure of a bacterial cell. Functions of the surface and cytoplasmic structures of a bacterial cell. Mechanism of Gram staining. Forms of bacteria with the cell wall defects.
- 5. Unique features of metabolism in prokaryotes. Nutrition of bacteria: types, requirements of bacteria, nutrients and pathways of nutrients penetration into the bacterial cell.
- 6. Respiration of microorganisms: types, pathways of energy production. Enzymes and cell structures involved in the process of respiration. Classification of bacteria regarding their oxygen requirements.
- 7. Growth and reproduction of bacteria. The mechanism of simple division and it's phases. Dormant forms of microorganisms: general characteristics, factors inducing their formation, medical importance.
- 8. Sampling for microbiological studies: types of samples, the rules of sampling, storage, transportation. Principles of organization, equipment and levels of biosafety in microbiological laboratories.
- 9. Microscopic (bacterioscopic) method of diagnosing the infectious diseases: definition, aim and tasks, steps and evaluation of specificity, sensitivity, disadvantages of the method. Types of microscopic preparations. Staining of microorganisms: methods. Types of microscopes.
- 10. The bacteriological method of diagnosing the infectious diseases: aim, tasks, phases, and evaluation of specificity, sensitivity, disadvantages of the method.
- 11. Cultivation of bacteria, nutrient media: requirements, classification. Methods for the isolation of pure cultures of aerobic and anaerobic bacteria.
- 12. Methods of identification of aerobic and anaerobic bacteria pure cultures. Identification of microorganisms without isolation of a pure culture.
- 13. Genetic apparatus of bacteria (nucleoid, plasmids, transposons, IS-elements) characteristics, functions, effect and importance. The concept of genetic engineering and biotechnology.
- 14. Inheritance and variability of microorganisms. Types of variability. Mutations. The genetic recombination of bacteria. Phenotypic variability. The practical significance of the variability of microorganisms in the diagnosis, treatment and prevention of infectious diseases.
- 15. Molecular biological method of diagnosing the infectious diseases (molecular hybridization, polymerase chain reaction): definition, the principle of the methods, application in dentistry.
- 16. Infection (infection process): definition of the term causes and conditions of infectious diseases emergence. Differences in communicable and non-communicable diseases. Periods of infectious diseases. Infectious disease classification and outcomes.
- 17. Classification of infectious processes: the nature of the pathogen, the source of infection, the mechanisms and routes of infection, prevalence, the multiplicity of infection, duration.

- 18. The role of microorganisms in the infectious process. Pathogenicity and virulence. Factors of pathogenicity of microorganisms. Pathogenicity island. Microbial toxins. Types of exotoxins and their biological properties. Mechanisms of microbial persistence and latency in host's organisms.
- 19. The role of host, social, environmental factors in the infectious process.
- 20. The biological (experimental) method of diagnosing the infectious diseases: definition of the term, aim, tasks, phases, evaluation.
- 21. The ecology of microorganisms. Types of ecological relationships in microorganisms. The role of microorganisms in the genesis and development of the Biosphere (the concept of the microbial dominance). Spread of microorganisms in the nature.
- 22. The characteristic of normal human microflora and its biological role. Methods of study. Disbiosis: causes, consequences, prevention. Gnotobiology.
- 23. Sterilization: definition of the term, methods, quality control. Sterilization of instruments and medical devices. Consequences of sterilization errors.
- 24. Disinfection: definition of the concept, types, methods of conducting. Groups of disinfectants used in dentistry.
- 25. The antisepsis: definition of the term, types, categories, methods of application. Antiseptic agents: classification, mechanism of action, side effects. Principles of rational antisepsis in dental practice.
- 26. The chemotherapy and chemoprophylaxis of infectious diseases. Groups of antimicrobial chemotherapeutic agents, mechanisms and spectrum of action on microbial cells. Chemotherapeutic index.
- 27. Antibiotics: characteristic, classification. Requirements for antibiotics. Mechanisms of action of antibiotics.
- 28. Principles of a rational antibiotic therapy in stomatology. Antibiotics for prophylaxis of bacterial complications. Side effects of antibiotics. New approaches to the development of antibiotics.
- 29. Natural and acquired resistance of microorganisms to antibiotics. The genetic and biochemical mechanisms of resistance of microorganisms.
- 30. Genotypic and phenotypic methods for determining the susceptibility of microorganisms to antibiotics. Instruments and test systems for the automated detection of antibiotic susceptibility of microorganisms
- 31. Immunology: definition of the term, aim and task, methods, history of development, branches. Immunity: definition, types of immunity.
- 32. Immune system of the body: organs, cells, molecules of the main histocompatibility complex (structure, distribution on cells, biological role), cytokines (classification, functions).
- 33. Innate immunity. Immune and non-immune factors of innate immunity. Mechanisms of recognition in the innate immune system.
- 34. Phagocytes, classification. Phagocytosis reaction: phases, mechanisms of intracellular microorganisms killing, outcomes. Methods of phagocytosis evaluation. Phagocytic reaction indexes, definition and importance in clinical practice.
- 35. The complement system: definition, main components, activators and activation pathways, functions of components and their fragments. Methods of evaluation of the complement system activity.
- 36. Antigens: structure, properties, classification. T-dependent and T-independent antigens. Superantigens.
- 37. Antigens of microorganisms. Antigenic structure of bacteria. Type, species, group antigens. Protective antigens. Cross- reactive antigens, medical importance.

- 38. Antigen presenting cells: types, characteristics. B-lymphocytes: development, markers, antigen-specific B-cell receptor.
- 39. Humoral immune response: definition, development. Activation, proliferation, differentiation and interactions of cells involved. T-dependent and T-independent response. Primary and secondary humoral immune response characteristics.
- 40. Antibodies (immunoglobulins): structure, properties, classification, Immunoglobulins biosynthesis. The mechanism of interaction of antibodies with antigens: specificity, phases, manifestations. Affinity and avidity. Monoclonal antibody: principles of production, application.
- 41. Serological method of investigation: general definition of the term, objectives, basic concepts (diagnosticum, diagnostic serum, titer, diagnostic titer, paired sera). Samples for serological examination. General characteristics of the method. Use of serological method for infectious and noninfectious diseases diagnostics.
- 42. Agglutination: ingredients, main variants of performance, registration, evaluation, application. Indirect (passive) and reverse passive agglutination: ingredients, mechanism, methodology, registration of results, practical use.
- 43. Immunoprecipitation reaction: ingredients, mechanism, main methods of performance, application.

  Reaction of the immune lysis. Complement fixation test: ingredients, mechanism, registration of results.
- 44. Immunofluorescence (fluorescent antibodies test, FAT), main variants, ingredients, mechanisms, registration of results, practical use. ELISA: ingredients, mechanisms, registration of results, practical use. Immunoblotting (IB). Radioimmunoassay (RIA).
- 45. T cells: development, markers, subpopulations. Helper T-cells, main types (Th1, Th2, Th3, Th17), spectrum of cytokines produced. T-cell receptor: structure, types, genetic control, variety.
- 46. Cellular immune response: definition, development, main stages, manifestation. The model of two (three) signals: the response, anergy, apoptosis. Manifestation of cellular immune response. Immunological memory.
- 47. Anti-infection immunity and its types depending on pathogen nature. Mechanisms of antitoxic, antibacterial, antifungal, antiparasite immunity.
- 48. Immunoprophylaxis and immunotherapy for infectious diseases. Active immunoprophylaxis. Vaccines: requirements, characteristics of main types of vaccines. Adjuvants mechanisms of action. Side effects of vaccination: sever vaccinal reaction, post-vaccination complications.
- 49. Post-vaccination immunity: mechanisms and factors influencing its development. Indications and contraindications to vaccination. Immunization schedule. Expanded Programme on immunization. Collective immunity to infectious diseases, importance.
- 50. Passive immunoprophylaxis and immunotherapy of infectious diseases: indications, principles, complications.
- 51. Allergology: the definition, objectives. Allergens. Allergy: the stages, types of reactions. Classification of allergens. Allergens in dentistry.
- 52. Immediate type hypersensitivity (ITH). Mediator type (I) ITH: allergens, mechanism, development, Manifestations in the oral cavity, ways to prevent anaphylaxis.
- 53. Cytotoxic (II) type ITH: allergens, development, mechanisms, manifestations. Immunocomplex (III) type ITH: allergens, development, mechanisms. Manifestations of allergic reactions II and III types in the oral cavity.
- 54. Delayed type of hypersensitivity (IV): allergens, development, mechanism, manifestation (infection and contact allergy), importance in oral cavity.

- 55. Drug allergy: major allergens, the mechanisms and types of allergic reactions, methods for diagnostics and prevention. Food allergy. Main allergens. Prevention of food allergy. Idiosyncrasy.
- 56. Methods of diagnosing allergic diseases. Prevention of allergy.
- 57. Antitumor immunity. The concept of immune surveillance. Mechanisms of tumor escape from immune surveillance.
- 58. Clinical Immunology: definition, objectives, main concepts. Immune status: principle and methods of examination. Methods for determining the amount and functional activity of T-and B-lymphocytes.
- 59. Autoantibodies: origin, role in pathology. Autoimmune diseases: definition, classification, aetiology, mechanisms of tissue damage, manifestations.
- 60. Immunodeficiency conditions: classification, causes of development, methods for detection, principles for correction.
- 61. Staphylococci: classification, characterization, antigenic structure, pathogenicity factors. Staphylococcal infections: pathogenesis, immunity, microbiological diagnosis and principles of prevention, immunotherapy. Staphylococcal carriage: diagnosis, significance. Staphylococcus aureus: MRSA, antibiotics of choice for their therapy.
- 62. Streptococci: classification, characterization, antigenic structure, pathogenicity factors. Streptococcal disease: pathogenesis, immunity, microbiological diagnosis, and prevention. Pneumococci: classification, characterization, antigenic structure, pathogenicity factors. Pneumococcal infections.
- 63. Neisseria meningitidis: systematics, characterization, antigenic structure, pathogenicity factors. Meningococcal infections: pathogenesis, immunity, microbiological diagnosis, prophylaxis.
- 64. Neisseria gonorrhoeae: systematics, characterization, antigenic structure, pathogenicity factors. Pathogenesis, immunity, microbiological diagnosis of acute and chronic gonorrhoea, prophylaxis. Prevention of gonorrhoea and gonorrhoeal conjunctivitis, stomatitis.
- 65. Family of Enterobacteria: classification, characterization, pathogenicity factors. Principle of microbiological diagnosis of GIT diseases caused by Enterobacteria. Principles of identification of enterobacteria.
- 66. Escherichia: systematics, characterization, antigenic structure, pathogenicity factors. Pathogenic and opportunistic Escherichia coli. The biological role of Escherichia coli. Escherichiosis: pathogenesis, immunity, microbiological diagnosis and prevention.
- 67. Salmonella: systematics and classification, characterization, antigenic structure, pathogenicity factors, role Salmonella in pathology. Salmonellosis and Typhoid fever: pathogenesis, immunity, prevention.
- 68. Shigella: classification, characteristics, antigenic structure, pathogenicity factors. Bacterial dysentery: pathogenesis, immunity, microbiological diagnosis, prophylaxis.
- 69. Food poisoning of microbial aetiology: classification, etiology, pathogenesis, principles of microbiological diagnosis, prophylaxis.
- 70. Klebsiella: classification, characteristics, antigenic structure, pathogenicity factors, Klebsiella diseases. Pseudomonas: characteristics, antigenic structure, pathogenicity factors, role in the pathology.
- 71. Campylobacter, Helicobacter: characteristics, role in pathology.
- 72. Corynebacterium: classification, characteristics, antigenic structure, pathogenicity factors. Diphtheria: pathogenesis, immunity, microbiological diagnostics, immunotherapy and aetiological therapy of diphtheria, prophylaxis. Manifestation of diphtheria in oral cavity.
- 73. Bordetella: classification, characteristics, antigenic structure, pathogenicity factors. Whooping cough: pathogenesis, immunity, microbiological diagnosis, prophylaxis. Haemophilus spp.: characteristics, role in pathology, prophylaxis Hib-infections.
- 74. Actinomyces: classification, characterization, antigenic structure, pathogenicity factors. Cervico-maxillofacial actinomycosis: pathogenesis, immunity, microbiological diagnosis, prevention.

- 75. Mycobacteria: classification, characteristics, antigenic structure, pathogenicity factors. Tuberculosis: pathogenesis, immunity, methods of diagnosis, principle of prevention and treatment. Mycobacterioses. Manifestation of tuberculosis in oral cavity.
- 76. Obligate anaerobes. Classification and characteristics. Clinical signs of anaerobic infection. Features of taking the material in case of suspected anaerobic infection.
- 77. Gas gangrene Clostridia spp.: classification, characteristics, antigenic structure, pathogenicity factors.

  Anaerobic myonecrosis: pathogenesis, immunity, microbiological diagnostics and prophylaxis, aetiological treatment.
- 78. Clostridium tetani: systematics, characterization, antigenic structure, pathogenicity factors. Tetanus: pathogenesis, immunity, microbiological diagnosis, prevention, aetiological treatment.
- 79. Nonsporforming anaerobes: classification, characteristics, role in pathology of oral cavity. Principles of sampling in anaerobic bacteriology. Principle of bacteriological diagnosis of infections caused by nonsporforming anaerobes.
- 80. Quarantine diseases: characteristics, classification. Principles of collection, transportation and investigation of specimens with pathogens of 3d and 4th biosafety levels.
- 81. Vibrio: classification, characteristics, antigenic structure, pathogenicity factors. Cholera: pathogenesis, immunity, microbiological diagnosis, prophylaxis.
- 82. Classification and characteristics of causative agents of plague, tularemia, pathogenicity factors, microbiological diagnosis, prophylaxis.
- 83. Classification and characteristics of causative agents of brucellosis, anthrax, pathogenicity factors, microbiological diagnosis, prophylaxis.
- 84. Spirochetes: classification, characteristics, antigenic structure, pathogenicity factors. Role of Borrelia spp. in human pathology. Lyme borreliosis: aetiology, pathogenesis, immunity, microbiological diagnosis, prophylaxis. Role of Leptospira in human pathology, prophylaxis of leptospirosis.
- 85. Treponema: classification, characteristics, antigenic structure, pathogenicity factors. Syphilis: pathogenesis, immunity, microbiological diagnosis, prophylaxis. Manifistation of Syphilis in oral cavity.
- 86. Treponema of oral cavity and their role in pathology. Fusospirochetozes: etiology, characteristics of pathogens, pathogenesis, clinical forms.
- 87. Chlamydia: classification, characterization, development cycle, antigenic structure, pathogenicity factors, role in pathology. Microbiological diagnostics and prevention.
- 88. Mycoplasma spp.; classification, characteristics, role in pathology.
- 89. Rickettsia: classification, characteristics, role in pathology.
- 90. Pathogenic fungi: classification, characteristics. Fungal infections promoting factors and conditions. Role microfungi in human pathology. Prophylaxis of mycoses.
- 91. Virology: definition, objectives, methods. Systematic position and classification of viruses. History. D.Ivanovski works importance. Forms of existence of viruses. Morphology and biochemical structure of virions. Structure, function and properties of virion nucleic acid, proteins, lipids and carbohydrates. Prions, role in human pathology.
- 92. Interaction of the virus and susceptible cell. Strict parasitism and cytotropism of viruses. Cell receptors for viruses. Viral genome organization. Reproduction strategy of DNA and RNA viruses.
- 93. Types of viral infection of cell. Changes in the host cells in the process of a viral infection. Peculiarities of viral infections of an organism. Acute, chronic and slow infection. Local and systemic mechanisms of antiviral immunity. Factors of innate and adaptive antiviral immunity. Interferons: classes, properties, mechanisms of antiviral activity.

- 94. Principles of etiologic diagnostics of viral infections. Rapid methods. Serological diagnostics: principles, criteria for diagnosis. Principles of viral infections chemotherapy. Groups of antiviral drugs.
- 95. Cultivation of viruses. Indication and identification of viruses.
- 96. The aetiology of acute respiratory viral infections. Influenza viruses: classification, characteristics, antigenic properties. Influenza: pathogenesis, immunity, prevention, etiologic diagnostics of influenza, chemotherapy and chemoprophylaxis of influenza.
- 97. Paramyxoviruses: classification, characteristics, role in pathology. Prevention of infection caused by paramyxoviruses
- 98. Rabies virus: classification, characteristics, specific inclusion. Rabies: pathogenesis, etiologic diagnosis, prevention.
- 99. Rubella virus. General characteristics. Role in pathology. Prevention of rubella.
- 100. Enteroviruses: classification, characteristics. Enterovirus infections: pathogenesis, prevention. Role in pathology of oral cavity.
- 101. Viral hepatitis A: pathogenesis, immunity, etiologic diagnosis, prevention.
- 102. Parenteral hepatitis viruses: classification, characteristics. Parenteral hepatitis: pathogenesis, immunity, etiologic diagnostics, prevention.
- 103. Retroviruses. Human immunodeficiency virus (HIV). HIV infection: pathogenesis, immunity, etiologic diagnostics, principles of therapy, prophylaxis. AIDS related illnesses. HIV-associated diseases in oral cavity.
- 104. Herpesviruses: classification, characterization, role in pathology. Herpetic stomatitis. Chickenpox. Herpes viruses of 4-8 types, their role in human pathology.
- 105. Adenoviruses: classification, characteristic. Adenoviral infections: pathogenesis, immunity, etiological diagnosis. Papillomaviruses: characteristics, role in pathology, disease prevention.
- 106. Dental microbiology: definition, goals, objectives. General principles of microbiological diagnosis of dental diseases
- 107. The microflora of the oral cavity (indigenous, transient). Ontogeny of normal oral flora.
- 108. The role of normal microflora of the oral cavity (positive and negative). Dysmicrobiosis of the oral cavity: causes, effects, prevention, principles of correction. Influence of environmental factors, physiological features of the oral cavity and other factors of the microorganism on the microflora of the oral cavity.
- 109. Representatives of the normal microflora of the oral cavity: aerobes and facultative anaerobes (streptococci, corynebacteria, staphylococci, Neisseria), their role. General characteristics of streptococci of the oral cavity.
- 110. Representatives of the normal oral flora: anaerobes (velonella, propionjbacterium, lactobacillus, actinomyces, bacteroides, prevotella, porphyromonas, fusobacterium, leptotrichia), their role.
- 111. Representatives of the normal oral flora spiralshaped bacteria (vibrio, wolinella, centipedia, selenomonas, campylobacter, spirochetes), mycoplasma, protozoa, fungi, and their role.
- 112. Microflora of specific areas of the mouth: saliva, dorsum of the tongue, dental pocket, mucous membranes. Features of these biotopes, affecting microorganisms.
- 113. Methods of study of oral microflora. Methods of sampling material for dental diseases. Environments for the isolation of cariogenic streptococci, lactobacilli.
- 114. Nonspecific mechanisms of defense of the mucous membranes, saliva, gingival fluid, tooth enamel, normal microflora's, system of polymorphonuclear leukocytes.
- 115. Functions of saliva. Antimicrobial factors of saliva: defensins, cathelicidin, mucins, histatin, statherin, cystatins, peroxidase.

- 116. The role of factors and mechanisms of acquired immunity of the oral cavity. Local immunity of the oral cavity. Functions of secretory immunoglobulins A.
- 117. Dental plaque: the stages of formation, microorganisms-colonizers. Plaque as a biofilm. The role of factors in the quorum of sensing in the formation of plaque. New approaches to reducing the bioburden of plaque.
- 118. Etiology of caries. Criteria of cariogenicity. Cariesogenic streptococci. Characteristic of S. mutans et sobrinus. Characteristics of lactobacilli. Associative (auxiliary) microorganisms. The role of the macroorganism in the development of caries.
- 119. Pathogenesis of caries: mechanisms of adhesion (carbohydrate-dependent and carbohydrate-independent) streptococci and mechanisms of destruction of tooth tissues. The role of streptococci in coaggregation. Glukans. Conditions for the development of caries. Caries resistance. Prophylaxis of caries. Fluorides and their influence are microorganisms.
- 120. Odontogenic inflammation: etiology, types and phases of inflammation. Significance in pathology of foci of chronic odontogenic infection. Immunological aspects of the relationship between inflammatory periodontal diseases, cardiovascular and rheumatic diseases.
- 121. Types of microorganisms and their role in the origin and pathogenesis of pulpitis, acute and chronic apical periodontitis, periostitis, osteomyelitis, abscesses and phlegmon soft tissues.
- 122. Periodontal disease: classification, risk factors for development. The role of microorganisms in the etiology and pathogenesis of gingivitis. Dynamics of microflora of implants in case of successful and complicated implantation.
- 123. The role of dental plaque in the development of periodontitis. The role of microorganisms in the formation of dental plaque. Pathogenetic importance of dental plaque.
- 124. General properties of periodontopathogenic microorganisms. Microorganisms of the red complex: Porphyromonas gingivalis, Tannerella forsythia, Treponema denticola. Characteristics, pathogenicity factors, their role in the pathogenesis of periodontitis.

- 125. Microorganisms of orange, green and yellow complexes, their role in the development of periodontal diseases. Characteristics Aggregatibacter actinomycetemcomitans, pathogenicity factors, the mechanism of invasion and persistence, a role in the development of periodontitis.
- 126. Immune mechanisms in diseases of periodontal tissues. Factors contributing to the invasion of microorganisms. Mechanisms of tissue protection from microbial invasion. Principles of prevention and treatment of periodontitis.
- 127. Inflammatory diseases of the oral mucosa: specific and nonspecific bacterial stomatitis.
- 128. Viral stomatitis.
- 129. Candida: systematics, properties, pathogenicity factors. Candidosis: factors responsible for the developement, methods of diagnosis and prevention.
- 130. Manifestations of allergic and immunodeficiency conditions in the oral cavity. Recurrent viral aphthous stomatitis.
- 131. Dental Clinical Microbiology. Opportunistic pathogens. Specific features opportunistic pathogens and infections caused by them. Specific features of pathogenesis and diagnosis of opportunistic diseases. Criteria of Etiological significance of isolated bacteria from a specimen.
- 132. Etiology and principles of microbiological diagnosis of opportunistic diseases of skin and subcutaneous tissue of stomatogenic origin.
- 133. Etiology and principles of microbiological diagnosis of opportunistic diseases of bronchopulmonary tract of stomatogenic origin.
- 134. Etiology and principles of microbiological diagnosis of bacteremia, sepsis of stomatogenic origin.
- 135. Nosocomial infections: definition of the term, etiology, incidence and spread, principles of microbiological diagnosis, prevention. Antiepidemic control in stomatology.

## PRACTICAL SKILLS FOR DEMONSTRATION (PRE-EXAM)

- 1. Prepare a smear from bullion culture of bacteria and stain by Gram method.
- 2. Prepare a smear from agar medium culture of bacteria and stain by Gram method.
- 3. Identify Staphylococcus spp.
- 4. Identify Streptococcus spp.
- 5. Identify Neisseria gonorrhoeae.
- 6. Identify Escherichia coli.
- 7. Identify a mixture of Staphylococcus spp. and Escherichia coli.
- 8. Identify a causative agent of anthrax Bacillus anthracis.
- 9. Identify Vibrio spp.
- 10. Identify Brucella spp.
- 11. Identify Candida spp.
- 12. Identify Corynebacterium diphtheria (Loffler stain).

- 13. Identify capsule of *Klebsiella spp.* (negative contrasting)
- 14. Identify Mycobacterium in sputum (Ziehl–Neelsen stain stain)
- 15. Demonstrate inoculation technique on plated agar medium from slant media.
- 16. Demonstrate inoculation technique on slant agar medium from plated medium.
- 17. Demonstrate inoculation technique on slant medium from slant medium.
- 18. Register and assess the results antibiotic susceptibility testing by disc diffusion method.
- 19. Assess the results of agglutination reaction in tubes.
- 20. Assess the results of Complement fixation test.
- 21. Assess the results of Indirect (passive) agglutination test.
- 22. Assess the results of haemagglutination inhibition test.
- **23.** Demonstrate the technique of slide agglutination testing.

### References

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- 1. Medical microbiology / F. H. Kayser [et al.]. 10<sup>th</sup> ed., 2005. 742 p.
- 2. Manual of Clinical Microbiology / ed. in chief, J. H. Jorgensen. 11<sup>th</sup> ed. American Society for Microbiology, 2015. P. 2892.
- 3. Samaranayake, L. Essential microbiology for dentistry / L. Samaranayake. 3rd ed., 2005. P. 389.
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- 6. Alexander, S. V. Lab Exercises in Organism and Molecular Microbiology / S. V. Alexander, D. Strete, M. J. Niles. 2004. 384 p.
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- 11. Paul, W.,E. Fundamental Immunology / W. E. Paul. 6th ed. Lippincott Williams & Wilkins, 2008. P. 1555.
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- 13. Immunobiology: immune system in health and disease / Ch. A. Janeway [et al.]. 5<sup>th</sup> ed. Garland Publishing, 2001. P. 735.
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#### **INTERNET SOURCE**

http://www.bsmu.by
http://www.ada.org
http://www.asm.org
American Society for Microbiology

The Foreith Institute

http://www.iadr.org International Association for Dental Research

http://www.nidcr.nih.gov The National Institute of Dental and Craniofacial Research

http://www.nih.gov The National Institutes of Health

This site provides important information

This site provides important information about practicing good oral hygiene

This organization provides valuable resources about bacteria and microorganisms.

This institute is a leader in oral biology research.

This association provides valuable resources about oral care and research in dentistry.

This site provides information about dental research funding in America.

The site provides information about grants and research funding in America.

# Appendix 1. Classification of bacteria

# PROCARIOTE by Bergy, 2001 DOMAIN BACTERIA

PHYLUM	CLASS	ORDER	FAMILY	GENUS	SPECIES
	Alphaproteo-	Rickettsiales	Rickettsiaceae	Rickettsia	R.prowazekii, R.typhi, R.felis,R.rickettsii, R.conorii, R.australis,R.akari, R.sibirica, R.japonica, R.honei
	• •			Orientia	O.tsutsugamushi
	bacteria		Ehrlichiaceae	Ehrlichia	E.chaffeensis, E.sennetsu, E.equilike (E.phagocytophila)
		Rhizobiales	Bartonellaceae	Bartonella	B.quintana, B.henselae, B.bacilliformis, B.chlaridgeae, B.elizabethae
			Brucellaceae	Brucella	B.melitensis, B.abortus, B.suis u ∂ρ.
	Betaproteo-	Burkholderiales	Burkholderiaceae	Burkholderia	B.mallei, B.pseudomallei, B.cepacia u ∂p.
			Alcaligenaceae	Alcaligenes	A.faecales и др.
	bacteria			Bordetella	B.pertussis, B.parapertussis, B.bronchiseptica и др.
		Neisseriales	Neisseriaceae	Neisseria	N.gonorrhoeae, N.meningitidis, N.sicca, N.subflava u ∂p.
				Eikenella	E.corrodens E.corrodens
				Kingella	K.kingae и др.
		Nitrozomonadales	Spirillaceae	Spirillum	S.minus и др.
		Thiotrichales	Francisellaceae	Francisella	F.tularensis
		Legionellales	Legionellaceae	Legionella	L.pneumophila и др.
			Coxiellaceae	Coxiella	C.burnetii
0,		Pseudomonadales	Pseudomonadaceae	Pseudomonas	P.aeruginosa u dp.
<b>Z</b>			Moraxellaceae	Moraxella	Подрод Moraxella (M.lacunata и др.); Подрод Branhamella (B.catarralis и др.)
9				Acinetobacter	A.calcoaceticus u ∂p.
Proteobacteria		Vibrionales	Vibrionaceae	Vibrio	V.cholerae (биовары: cholerae, eltor), V.parahaemolyticus, V.vulnificus, V.sputorum и др.
X	ja	Aeromonadales	Aeromonadaceae	Aeromonas	A.hydrophilia
1.2	Gammaproteobacteria	Enterobacteriales	Enterobacteriaceae	Enterobacter	E.cloacae, E.sakazakii, E.agglomerans, E.gergoviae и др.
7				Calymmatobacterium	C.granulomatis
)	ğ			Citrobacter	C.freundii, C.amalonaticus, C.diversus и др.
te	qc			Edwardsiella	E.tarda u др.
0	l e			Erwinia	E.amylovora и др.
<u> </u>	ot			Escherichia	E.coli, E.fergusonii, E.germannii, E.vulneris, E.blattae
<u>a</u>	20		4	Hafnia	H. alvei
	a			Klebsiella	К.pneumoniae (подвиды: ozaenae, rhinoscleromae, pneumoniae), К.oxytoca, К.planticola, К.terrigena
	8			Morganella	M.morganii
	8			Plesiomonas	P.shigelloides
	ja ja			Proteus	P.vulgaris, P.mirabilis, u δp.
	0			Providencia	P.alcallifaciens и др.
				Salmonella	S.enterica, S.bongori. Вид S.enterica состоит из 6 подвидов (subsp.: arizonae, diarizonae, enterica, houtenae,
					indica, salamae). Серовары: S.Typhi, S.Paratyphi A, S.Schottmuelleri, S.Enteritidis, S.Typhimurium, S.Choleraesuis
				0 11	u δρ.
				Serratia	S.marcescens u dp.
				Shigella	S.dysenteriae, S.flexneri, S.boydii, S.sonnei
		De et e cellete	De de de la lace	Yersinia	Y.pestis, Y.enterocolitica, Y.pseudotuberculosis u δp.
		Pasteurellales Control of the state of the s	Pasteurellaceae	Haemophilus	H.influenzae, H.ducreyi u dp.
	Epsilonpro-	Campylobacteriales	Campylobacteriaceae	Campylobacter	C.jejuni, C.fetus, C.coli u dp.
	teobacteria		Helicobacteriaceae	Helicobacter	H.pylori, H.heilmanii u ∂p.
				Wolinella	W.succinogenes

PHYLUM	CLASS	ORDER	FAMILY	GENUS	SPECIES
	Clostridia	Clostridiales	Clostridiaceae	Clostridium	C.botulinum, C.perfringens, C.novyi, C.histolyticum, C.septicum, C.tetani, C.defficile и др.
			Peptostreptococcaceae	Peptostreptococcus	P.anaerobius u ∂p.
			Peptococcaceae	Peptococcus	P.niger
				Centipeda	C.periodontii
				Mitsuokella	M.dentalis
10			Acidaminococcaceae	Selenomonas	S.sputigena
9				Veillonella	V.parvula и др.
Firmicutes	Mollicutes	Mycoplasmatales	Mycoplasmataceae	Mycoplasma	M.pneumoniae, M.hominis, M.fermentans, M.salivarum, M.orale, M.artritidis u ∂p.
75				Ureaplasma	U.urealiticum и др.
	Bacilli	Bacillales	Bacillaceae	Bacillus	B.anthracis, B.cereus u ∂p.
, <b>5</b>	20.0		Listeriaceae	Listeria	L.monocytogenes u др.
<i>i</i> ;			Staphylococcaceae	Staphylococcus	S.aureus, S.epidermidis, S.saprophyticus u ∂p.
		Lactobacillales	Lactobacillaceae	Lactobacillus	L.caseii, L.fermentum, u dp.
			Enterococcaceae	Enterococcus	E.faecalis, E.faecium и др.
			Leuconostoccaceae	Leuconostoc	L.mesenteroides
			Streptococcaceae	Streptococcus	S.pyogenes, S.pneumoniae, S.agalactiae, S.anginosus, S.bovis, S.mutans, S.mitis, S.salivarius, S.sanguis, S.milleri
					u ∂p.
				Lactococcus	L.lactis u ∂p.
Actino-	Actinobacteria	Actinomycetales	Actinomycetaceae	Actinomyces	A.israelii, A.naeslundii, A.viscosus, A.odontolyticus, A.pyogenes,
ACUITO-			Micrococcaceae	Micrococcus	M.lysodeicticum, M.luteus и др.
bacteria			Corynebacteriaceae	Corynebacterium	C.diphtheriae, C.ulcerans, C.urealyticum, C.xerosis u др.
bucteriu			Mycobacteriaceae	Mycobacterium	M.tuberculosis, M.bovis, M.africanum, M.leprae, M.kasasii, M.avium, M.ulcerans, M.fortuitum u др.
			Nocardiaceae	Nocardia	N.asteroides, N.farcinica u ∂p.
			Propionibacteriaceae	Propionibacterium	P.acnes, P.propionicus u dp.
		Bifidobacteriales	Bifidobacteriaceae	Bifidobacterium	B.bifidum u ∂p.
				Gardnerella	G.vaginalis
Chlamydiae	Chlamydiae	Chlamydiales	Chlamydiaceae	Chlamydia	C.trachomatis
omanny and c	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			Chlamydophila	C.psittaci, C.pneumoniae
Spirochaetes	Spirochaetes	Spirochaetales	Spirochaetaceae	Borrelia	B.recurrentis, B.burgdorferi, B.duttoni, B.persica u ∂p.
Cp. Condicios	-			Treponema	Т.pallidum (подвиды – pallidum, endemicum, pertenue), T.carateum, T.denticola, T.minutum, T.refringens,
			4		T.scoliodontum, T.vincentii и др.
			Leptospiraceae	Leptospira	L.interrogans, L.biflexa
<b>Bacteroidetes</b>	Bacteroidetes	Bacteroidales	Bacteroidaceae	Bacteroides	B.fragilis, B.gingivalis u ∂p.
			Porphyromonadaceae	Porphyromonas	P.gingivalis, P.endodontales u ∂p.
			Prevotellaceae	Prevotella	P.melaninogenica, P.denticola и др.
	Flavobacteria	Flavobacteriales	Flavobacteriaceae	Flavobacterium	F.meningosepticum, F.breve и др.
Fusobacteria	Fusobacteria	Fusobacteriales	Fusobacteriaceae	Fusobacterium	F.nucleatum, F.necroforum, F.vincentii и др.
				Leptotrichia	L.buccalis и др.
				Streptobacillus	S.moniliformis

## Appendix 2. Classification of viruses (updates approved during EC 48, Budapest, Hungary, August 2016; Email ratification 2017)

Genome	Order	Family	Subfamily	Genus	Species	
dsDNA	Herpesvirales	Herpesviridae	Alphaherpesvirinae	Simplexvirus	Human alphaherpesvirus 1, 2	
dsDNA	Herpesvirales	Herpesviridae	Alphaherpesvirinae	Varicellovirus	Human alphaherpesvirus 3	
dsDNA	Herpesvirales	Herpesviridae	Betaherpesvirinae	Cytomegalovirus	Human betaherpesvirus 5	
dsDNA	Herpesvirales	Herpesviridae	Betaherpesvirinae	Roseolovirus	Human betaherpesvirus 6A, 6B, 7	
dsDNA	Herpesvirales	Herpesviridae	Gammaherpesvirinae	Lymphocryptovirus	Human gammaherpesvirus 4	
dsDNA	Herpesvirales	Herpesviridae	Gammaherpesvirinae	Rhadinovirus	Human gammaherpesvirus 8	
dsDNA	Unassigned	Adenoviridae		Mastadenovirus	Human mastadenovirus A-F	
dsDNA	Unassigned	Iridoviridae	Alphairidovirinae	Lymphocystivirus	Lymphocystis disease virus 1	
dsDNA	Unassigned	Papillomaviridae		Alphapapillomavirus	Alphapapillomavirus 1-72	
dsDNA	Unassigned	Papillomaviridae		Betapapillomavirus	Betapapillomavirus 1	
dsDNA	Unassigned	Papillomaviridae		Deltapapillomavirus	Deltapapillomavirus 1	
dsDNA	Unassigned	Papillomaviridae		Gammapapillomavirus	Gammapapillomavirus 1	
dsDNA	Unassigned	Polyomaviridae		Alphapolyomavirus	Human polyomavirus 12	
dsDNA	Unassigned	Polyomaviridae		Betapolyomavirus	Human polyomavirus 1	
dsDNA	Unassigned	Polyomaviridae		Deltapolyomavirus	Human polyomavirus 6	
dsDNA	Unassigned	Poxviridae	Chordopoxvirinae	Molluscipoxvirus	Molluscum contagiosum virus	
dsDNA	Unassigned	Poxviridae	Chordopoxvirinae	Orthopoxvirus	Vaccinia virus	
dsDNA	Unassigned	Poxviridae	Chordopoxvirinae	Orthopoxvirus	Variola virus	
dsDNA	Unassigned	Poxviridae	Chordopoxvirinae	Orthopoxvirus	Monkeypox virus	
ssDNA(-)	Unassigned	Anelloviridae		Alphatorquevirus	Torque teno virus 1	
ssDNA(-)	Unassigned	Anelloviridae		Betatorquevirus	Torque teno mini virus 1	
ssDNA(-)	Unassigned	Anelloviridae		Gammatorquevirus	Torque teno midi virus 1	
ssDNA(+/-)	Unassigned	Circoviridae		Circovirus	Human associated circovirus 1	
ssDNA(+/-)	Unassigned	Genomoviridae		Gemykibivirus	Human associated gemykibivirus 1	
ssDNA(+/-)	Unassigned	Genomoviridae		Gemyvongvirus	Human associated gemyvongvirus 1	
ssDNA(+/-)	Unassigned	Parvoviridae	Parvovirinae	Bocaparvovirus	Ungulate bocaparvovirus 1	
dsDNA-RT	Unassigned	Hepadnaviridae		Orthohepadnavirus	Hepatitis B virus	
ssRNA(-)	Bunyavirales	Nairoviridae		Orthonairovirus	Crimean-Congo hemorrhagic fever orthonairovirus	
ssRNA(-)	Bunyavirales	Peribunyaviridae		Orthobunyavirus	Bunyamwera orthobunyavirus	
ssRNA(-)	Bunyavirales	Peribunyaviridae		Orthobunyavirus	California encephalitis orthobunyavirus	
ssRNA(-)	Mononegavirales	Bornaviridae		Bornavirus	Mammalian 1 bornavirus	
ssRNA(-)	Mononegavirales	Filoviridae		Ebolavirus	Bundibugyo/Reston/Sudan/Taï Forest/Zaire ebolavirus	
ssRNA(-)	Mononegavirales	Filoviridae		Marburgvirus	Marburg marburgvirus	
ssRNA(-)	Mononegavirales	Paramyxoviridae		Henipavirus	Hendra henipavirus	
ssRNA(-)	Mononegavirales	Paramyxoviridae		Morbillivirus	Measles morbillivirus	
ssRNA(-)	Mononegavirales	Paramyxoviridae		Respirovirus	Human respirovirus 1, 3	
ssRNA(-)	Mononegavirales	Paramyxoviridae		Rubulavirus	Human rubulavirus 2, 4	
ssRNA(-)	Mononegavirales	Paramyxoviridae		Rubulavirus	Mumps rubulavirus	
ssRNA(-)	Mononegavirales	Pneumoviridae		Metapneumovirus	Human metapneumovirus	
ssRNA(-)	Mononegavirales	Pneumoviridae		Orthopneumovirus	Human orthopneumovirus	
ssRNA(-)	Mononegavirales	Rhabdoviridae		Lyssavirus	Rabies lyssavirus	
ssRNA(-)	Mononegavirales	Rhabdoviridae		Vesiculovirus	Indiana vesiculovirus	
ssRNA(-)	Unassigned	Orthomyxoviridae		Influenzavirus A	Influenza A virus	
ssRNA(-)	Unassigned	Orthomyxoviridae		Influenzavirus B	Influenza B virus	
ssRNA(-)	Unassigned	Orthomyxoviridae		Influenzavirus C	Influenza C virus	
ssRNA(-)	Unassigned	Orthomyxoviridae		Influenzavirus D	Influenza D virus	

Genome	Order	Family	Subfamily	Genus	Species
ssRNA(-)	Unassigned	Orthomyxoviridae	, ·	Quaranjavirus	Quaranfil virus
ssRNA(-)	Unassigned	Orthomyxoviridae		Thogotovirus	Thogoto virus
ssRNA(-)	Unassigned	Unassigned		Deltavirus	Hepatitis delta virus
ssRNA(+/-)	Bunyavirales	Phenuiviridae		Phlebovirus	Rift Valley fever phlebovirus
ssRNA(+/-)	Bunyavirales	Phenuiviridae		Phlebovirus	Uukuniemi phlebovirus
ssRNA(+/-)	Unassigned	Arenaviridae		Mammarenavirus	Junín mammarenavirus
ssRNA(+/-)	Unassigned	Arenaviridae		Mammarenavirus	Lassa mammarenavirus
ssRNA(+/-)	Unassigned	Arenaviridae		Mammarenavirus	Lymphocytic choriomeningitis mammarenavirus
ssRNA(+/-)	Unassigned	Arenaviridae		Mammarenavirus	Machupo mammarenavirus
ssRNA(+)	Nidovirales	Coronaviridae	Coronavirinae	Alphacoronavirus	Human coronavirus 229E, NL63
ssRNA(+)	Nidovirales	Coronaviridae	Coronavirinae	Betacoronavirus	Human coronavirus HKU1
ssRNA(+)	Nidovirales	Coronaviridae	Torovirinae	Torovirus	Human torovirus
ssRNA(+)	Picornavirales	Picornaviridae		Aphthovirus	Foot-and-mouth disease virus
ssRNA(+)	Picornavirales	Picornaviridae		Cardiovirus	Cardiovirus A
ssRNA(+)	Picornavirales	Picornaviridae		Cosavirus	Cosavirus A
ssRNA(+)	Picornavirales	Picornaviridae		Enterovirus	Enterovirus C
ssRNA(+)	Picornavirales	Picornaviridae		Enterovirus	Rhinovirus A
ssRNA(+)	Picornavirales	Picornaviridae		Hepatovirus	Hepatovirus A
ssRNA(+)	Picornavirales	Picornaviridae		Kobuvirus	Aichivirus A
ssRNA(+)	Picornavirales	Picornaviridae		Parechovirus	Parechovirus A, B, C
ssRNA(+)	Picornavirales	Picornaviridae		Rosavirus	Rosavirus A
ssRNA(+)	Picornavirales	Picornaviridae		Salivirus	Salivirus A
ssRNA(+)	Unassigned	Astroviridae		Mamastrovirus	Mamastrovirus 1
ssRNA(+)	Unassigned	Caliciviridae		Norovirus	Norwalk virus
ssRNA(+)	Unassigned	Caliciviridae		Sapovirus	Sapporo virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	Dengue virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	Japanese encephalitis virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	Murray Valley encephalitis virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	Omsk hemorrhagic fever virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	Tick-borne encephalitis virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	West Nile virus
ssRNA(+)	Unassigned	Flaviviridae		Flavivirus	Yellow fever virus
ssRNA(+)	Unassigned	Flaviviridae	4	Flavivirus	Zika virus
ssRNA(+)	Unassigned	Flaviviridae		Hepacivirus	Hepacivirus C
ssRNA(+)	Unassigned	Flaviviridae		Pegivirus	Pegivirus H
ssRNA(+)	Unassigned	Hepeviridae		Orthohepevirus	Orthohepevirus A
ssRNA(+)	Unassigned	Togaviridae		Alphavirus	Chikungunya virus
ssRNA(+)	Unassigned	Togaviridae		Alphavirus	O'nyong-nyong virus
ssRNA(+)	Unassigned	Togaviridae		Alphavirus	Semliki Forest virus
ssRNA(+)	Unassigned	Togaviridae		Alphavirus	Sindbis virus
ssRNA(+)	Unassigned	Togaviridae		Alphavirus	Venezuelan equine encephalitis virus
ssRNA(+)	Unassigned	Togaviridae		Rubivirus	Rubella virus
dsRNA	Unassigned	Picobirnaviridae		Picobirnavirus	Human picobirnavirus
dsRNA	Unassigned	Reoviridae	Sedoreovirinae	Rotavirus	Rotavirus A-G
dsRNA	Unassigned	Reoviridae	Spinareovirinae	Coltivirus	Colorado tick fever virus
ssRNA-RT	Unassigned	Retroviridae	Orthoretrovirinae	Deltaretrovirus	Primate T-lymphotropic virus 1
ssRNA-RT	Unassigned	Retroviridae	Orthoretrovirinae	Lentivirus	Human immunodeficiency virus 1, 2
ssRNA-RT	Unassigned	Retroviridae	Spumaretrovirinae	Spumavirus	Simian foamy virus

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