SPECIAL AND CLINICAL MICROBIOLOGY

Laboratory workbook

Student name _	 	 	
Faculty	 	 	
Group			

Minsk BSMU 2022

МИНИСТЕРСТВО ЗДРАВООХРАНЕНИЯ РЕСПУБЛИКИ БЕЛАРУСЬ

БЕЛОРУССКИЙ ГОСУДАРСТВЕННЫЙ МЕДИЦИНСКИЙ УНИВЕРСИТЕТ КАФЕДРА МИКРОБИОЛОГИИ, ВИРУСОЛОГИИ, ИММУНОЛОГИИ

ЧАСТНАЯ И КЛИНИЧЕСКАЯ МИКРОБИОЛОГИЯ SPECIAL AND CLINICAL MICROBIOLOGY

Лабораторный практикум

7-е издание



Минск БГМУ 2022

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Содержит информацию для подготовки к практическим занятиям по разделам частной и клинической микробиологии. Приведены схемы, алгоритмы, справочные сведения, методики выполнения лабораторных работ. Первое издание вышло в 2015 году.

Предназначен для студентов 2-го курса медицинского факультета иностранных учащихся, обучающихся на английском языке.

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<u>Class № 1.</u> Microbiological diagnostics of diseases caused by staphylococci, streptococci, neisseria

Date

The subject to study:

Staphylococci, systematics, general characteristics. Methods of microbiological diagnostics of staphylococcal infections. The material for the research depending on the form of the infection. Scheme of pure culture isolation (from pus, mucus, blood, etc.). Identification methods, phage typing. Specific prevention and treatment of staphylococcal infections.

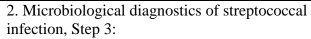
Streptococci. Systematics. Pyogenic streptococci. Pneumococci. General characteristics. Antigenic structure. Acute and chronic diseases, pathogenesis, immunity. Specific antibodies to streptococcal antigens, diagnostic value. Methods for streptococcal infections diagnosis. Bacteriological method, study design. Material for studies depending on the form of the infection, the rules and methods for taking material. Principles of therapy and prevention pro-streptococcal infections.

Neisseria. Systematics, general characteristics.

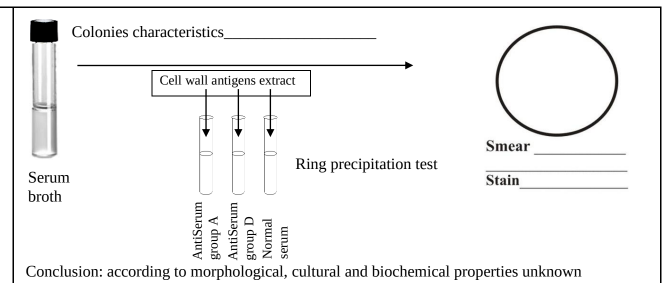
Characteristics of the causative agent, mechanisms of pathogenesis, immunity, methods of microbiological diagnosis of acute and chronic gonorrhea. Characteristics of the causative agent, mechanisms of pathogenesis, immunity, diagnosis and prevention of meningococcal infection-howl. sources:

Laboratory work

Tasks	Methods, results	
Microbiological diagnostics of staphylococcal infection, Step 2 a) macro- and microscopic examination of the colonies on YSA; b) plasmacoagulase test.	Smear MSA (YSA) Sh Siz Su Ed Co Cid Tr:	Feature Staphylococcal colonies hape ze urface dige color donsistency ransparency ecithinase
	Stabilized rabbit plasma, 37 °C, 2–4–24 h. Conclusion: according to morphological, cultural and biochemic bacterium is identified as	

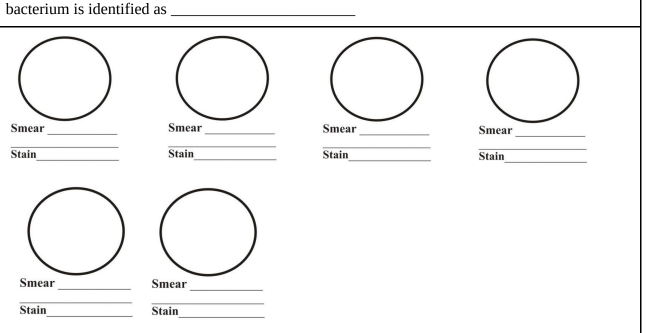


- a) a description of Streptococci growth in serum broth;
- b) determining the morphology of streptococci, Gram staining;
- c) determination of streptococcus serogroup by ring precipitation test.



Demonstration.

- 1. Staphylococcus aureus in pus, Gram staining.
- 2. Streptococcus pneumoniae, pure culture, Gram staining.
- 3. Streptococcus pneumoniae, white mice, Gram staining.
- 4. Gonococcus in pus, Gram staining.
- 5. Meningococcus in cerebrospinal fluid, methylene blue.
- 6. The growth of staphylococci on YSA, blood agar, broth.
- 7. The growth of streptococci on blood agar and broth.
- 8. Plasmacoagulase test
- 9. Anaerobic mannitol fermentation.
- 10. Phage typing of staphylococci.



Signature of the tutor_____

Complementary materials to class 5.

Main pathogenic species		Materia	al for investigation			Staphyloc infectio	
Morphology (size, shape, relative positions of cells) Spores development Capsule Flagella (motility) Gram staining Catalase activity Main pathogenicity factors			pure culture isolation				
	afection diagnostics		Staphylo	ococcus indentifica	tion		
Methods for staphylococcal in							
Method Microscopic	Usage (+/-)	Species	Plasmacoagulation test	Anaerobic mannitol fermentation	DNA-se	Lecithinase	Protein-A
Method Microscopic Cultural	Usage	Species S. aureus	_	mannitol	DNA-se	Lecithinase	Protein-A
Method	Usage		_	mannitol	DNA-se	Lecithinase	Protein

Streptococcus genus characteristics

Main pathogenic	S. pyogenes	S. pneumoniae
species		
Morphology		
Spores development		
Capsule		
Flagella (motility)		
Gram staining		
Group antigen		
Type-specific antigen		
(M-protein)		
Capsule		
polysaccharide		
Catalase activity		

Methods for streptococcal infections diagnostics

Mathada	Usage (+/-)				
Methods	S. pyogenes	S. pneumoniae			
Microscopic					
Cultural					
Biological					
Serological					
Allergic					
Molecular-genetic					

Bacteriological diagnostics of streptococcal infection S. pyogenes infections **Material for investigation** Media for pure culture isolation S. pneumoniae infection **Medium for pure culture accumulation** Other important Str. species **Streptococci identification** Inulin fermentatio Hemolysis (α, β, γ) Growth in nutrition broth Precipitati on nest Optochin test Capsule swelling Bile test Str. species

S. pyogenes

S. pneumoniae

E. faecalis

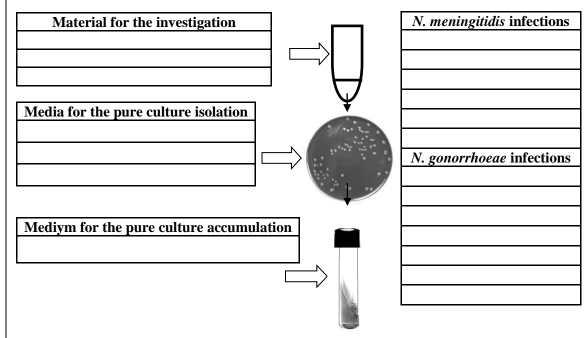
Neisseria genus characteristics

Features	N. meningitidis	N. gonorrhoeae
Morphology (size,		
shape, relative		
positions of cells)		
Spores development		
Capsule		
Flagella (motility)		
Gram staining		
Oxidase activity		
Pathogenicity factors		

Methods for neisserial infections diagnostics

Mathada	Usage (+/-)				
Methods	N. meningitidis	N. gonorrhoeae			
Microscopic					
Cultural					
Biological					
Serological					
Allergic					
Molecular-genetic					

Bacteriological method for the *Neisseria* infections diagnostics



Neisseria differentiation

Species	nutrition	Growth at 20 °C	Colonies color	Fermentation			
				Glucose	Maltose		
N. mening	itidis						
N. gonorrh	поеае						
Opportuni species	stic						

Class № 2. Microbiological diagnostics of acute enteric infections caused by enterobacteria

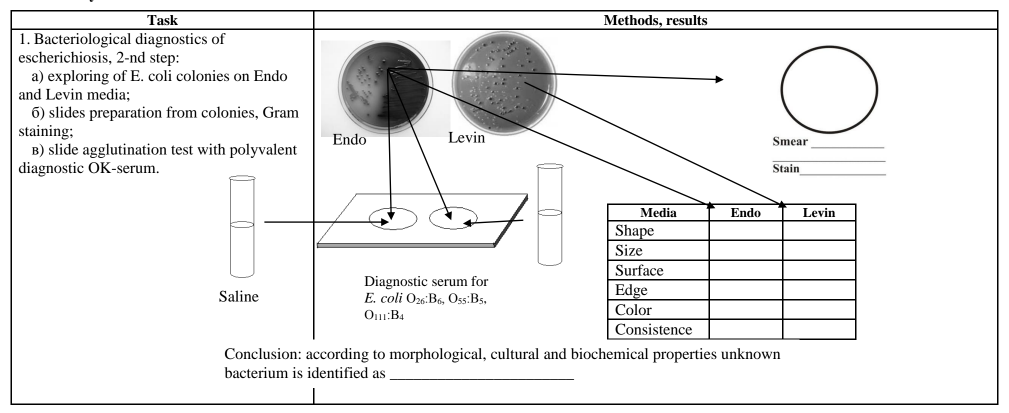
Questions to study: General characteristics of Enterobacteriaceae family. Differences between genera. General principles of diagnostics of acute enteric infections caused by pathogenic enterobacteria. Differential diagnostic media, composition, plinciple of work.

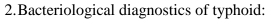
Escherichia, systematic position, general characteristics. The biological role of Escherichia coli. Molecular mechanisms of escherihiosis pathogenesis. Enteropathogenic, enterotoxigenic, enteroinvasive and enterohaemorrhagic Escherichia coli. Escherihisis diagnostics. Antibiotic treatment.

Salmonella, classification and general characteristics. Serological classification of Salmonella. Identification of Salmonella. Molecular biological typing.

Causative agents of typhoid and paratyphoid. The pathogenesis of typhoid. Microbiological diagnostics of typhoid fever, depending on the stage of pathogenesis.

Laboratory work



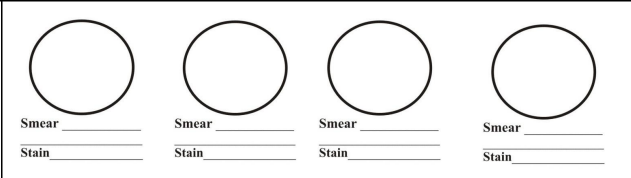


- 2-nd step of coproculture isolation:
 - a) describe colonies on Levin medium;
- b) prepare slide from colonies, Gram staining;
 - c) inoculate Kligler medium.

		SI (Kligler) edium	Smear	
Feature	Levin medium	•	4	
Shape	_		Stain	
Size				
Surface		4		
Edge				
Color				
Consistence				

Demonstration.

- 1. Clean media: Endo, Levin Ploskirev, bismuth sulfite agar, Rapoport, magnesium, Kliglera.
- 2. The same media with the growth of E. coli, Salmonella, Shigella.
- 3. Biochemical Activity of E. coli and Salmonella.
- 4. Dendrograms of Salmonella molecular typing.
- 5. Tube agglutination test with killed E. coli culture.
- 6. The morphology of E. coli, Salmonella, Shigella (Gram staining).



Teacher signature_____

Complementary materials to class 2.

Enterobact	teriaceae genera of n	nedical importance	Methods for diagno	stics of escherichios	is and salmonellosis
			- $	Usage (+/-)	
			Methods	Escherichiosis	Typhoidand paratyphoid
			Microscopic		
			Cultural		
General cha	aracteristics of <i>Enter</i>	obacteriaceae family	Biological		
Characteristic		Enterobacteriaceae	Serological		
Morphology			Allergic		
Spores development			Molecular-genetic		
Capsule			Bacteriological diagr	nostics of escherichi	osis
Flagella (motility)			Material for the	e investigation	
Gram staining				, , , , , , , , , , , , , , , , , , ,	└ / \
Antigens					↓
Exotoxins					
Endotoxins			Media for pure o		
E	Escherichia coli chard	acteristics	Media for pure o	culture isolation	
Characteristics		Escherichia coli			
Morphology					
Spores development				1, 1,	
Capsule			Medium for the pure	culture accumulation	
Flagella (motility)					
Gram staining			Biol	logical properties <i>E. c</i>	eoli,
Antigens			as norm	al microflora represe	
Number of serovars			Positive		Negative
E. coli classification	1.				
according to	2. 3.				
pathogenicity factors	4.				
Diseases caused by E. coli					

Characteristics of certain species from $\it Escherichia$ and $\it Salmonella$ genera

			Fermentat	ion		Indol	H ₂ S production	Catalase activity	Antigenic formula (O, H, K)	
Species	Glucose	Lactose	Mannitol	Maltose	Saccharose	production				
E. coli										
S. typhi										
S. paratyphi A										
S. schottmuelleri										
S. typhimurium										

Methods of microbiological typhoid diagnostics depending on the pathogenesis phase

Dathogonosis phase			Bacteriologi	cal method	Serolog	gical method	
	Pathogenesis phase		Urinoculture	Coproculture	Bileculture	Vidal test	BPAT with Vi-Ag
Incubation	period						
Prodromal period							
midst of	Bacteremia and intoxication						
midst of	Parenchymal diffusion						
illness Allergic-secretory							
Reconvalescence							
Bacteria ca	arrier state				·		

<u>Class № 3.</u> Microbiological diagnostics of acute enteric diseases caused by enterobacteria

Date

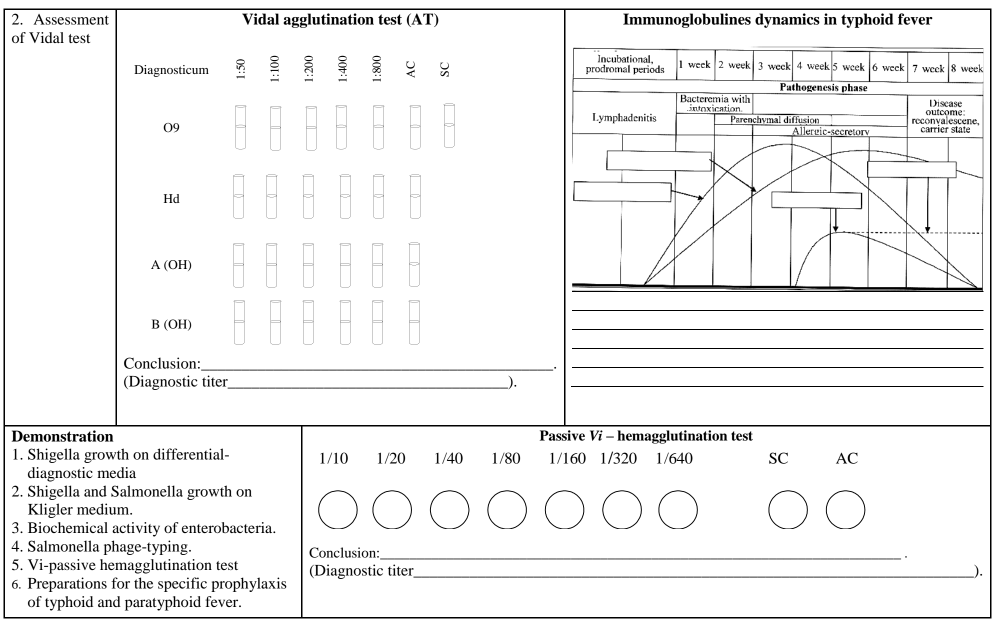
The list of questions to study: Characteristics of immunity in typhoid and paratyphoid fever. Serological diagnosis of typhoid and paratyphoid fever. Formulation and analysis of Vidal reaction. Methods for distinguishing infection, anamnestic and postvaccinal titer. Diagnosis of bacteria carrier state in typhoid fever.

Salmonella - causative agents of acute gastroenteritis. Salmonella phage typing and indication.

Shigella. Causative agents of dysentery, classification, general characteristics. Molecular mechanisms of pathogenesis, immunity, methods of laboratory diagnosis of acute and chronic dysentery. Approaches to the prevention of dysentery. Antibiotic treatment.

Laboratory work

Task	Methods, results
 Microbiological diagnostics of typhoid fever: 3-rd step. a) Describe the growth on the Kligler medium; b) prepare the slide from the colonies, Gram staining; a) check the culture for motility and indol production; d) determination of the antigenic structure of the culture isolated in slide agglutination test. 	Biochemical properties assessment: Lactose Glucose H ₂ S production Motility test



Supporting materials to class 3.

	Shigella clas	ssification		GI . II		Main salmonellosis pathogens
Shigella species	3	Serovarient	s number	Shigella	Salmonella րապատարապ	
				000	9 00	
Cultural method for t			s			Bacteriological method for salmonellosis diagnostics
Materials for			$\Rightarrow \bigcup$			Material for the research
Media for the pur	e culture iso	lation		5.4		Medium for the material enrichment
Medium for the pure	culture accu	mulation	\Rightarrow			Medium for pure culture isolation
Shi	gella chara	cteristics		@ @ @@		
Feature Glucose (A+G) Lactose	S. sonnei	S. flexneri	S. dysenteriae	Abscesses in	Lymph. nodes	Medium for pure culture accumulation
Mannitol				mucosa	1	\Rightarrow
Serogroup				Shigellosis and spathogenesis	Blood almonellosis	Methods for salmonella identification

<u>Class № 4.</u> Microbiological diagnostics of diseases caused by Klebsiella, Iersinia, Campylobacter and pseudomonada. <u>Methods for food poisoning diagnostics</u> <u>Data</u>

List of questions to study: Klebsiella, classification and general characteristics, main diseases caused. Pathogenesis, immunity, methods of microbiological diagnosis of acute and chronic klebsiellosis.

The causative agent of intestinal yersiniosis, general characteristics. Pathogenesis, immunity, methods of microbiological diagnostics.

Campylobacter, general characteristics, role in human pathology. Mechanisms of pathogenesis. Diagnosis of campylobacteriosis. Helicobacter.

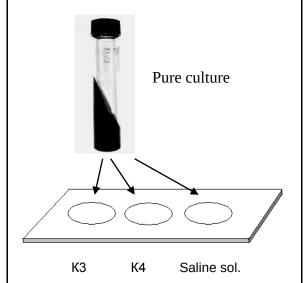
Pseudomonas aeruginosa, general characteristics, pathogenicity factors, role in human pathology. Methods of microbiological diagnostics Pseudomonas infection.

Classification, etiology of food poisoning. Principles of microbiological diagnostics.

Laboratory work

Tasks Methods, results 1. Independent work "Microbiological klebsiellosis diagnostics": A. Examine the growth of Klebsiella on differential-diagnostic media. B. Determine the capsule presence. C. Determine the biochemical properties of Klebsiella. D. Perform slide agglutination test with Smear anti-capsule diagnostic sera and determine the K-antigen. Saccharose Citrate Urea Malonate Stain Russell medium E. Determine the titer of CFT for (Simmonds medium) serological diagnosis of scleroma. Klebsiella characteristics K. pneumoniae K. pneumoniae K. pneumoniae Biochemical properties s. rhinoscleromatis s. pneumoniae s. ozaenae Glucose (A+G) +/-+/-Lactose + Saccharose (4th day) +/-Citrate +/-Urea **-/**+ Malonate +

Slide agglutination test with anti-capsule serum

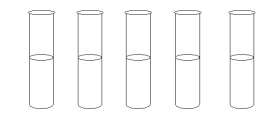


Conclusion:____

Complement fixation test

Variant	S	erum dilution	ıs	CS	CA	Result
Variant	1:5	1:10	1:20	CS		
1	++++	++++	++++	-	-	Very positive
2	++++	++++	-	-	-	Positive
3	+++	-	-	-	-	Weak positive
4	-	-	-	-	-	Negative

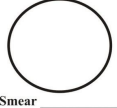
CA 1:10 1:20 CS



Conclusion:

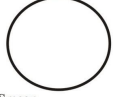
Demonstration.

- 1. Klebsiella growth on differential diagnostic media.
- 2. Klebsiella scleroma capsule (Hins-Burri staining).
- 3. Pseudomonas aeruginosa, pure culture, Gram staining.
- 4. Oxidase test.



Smear

Stain



Smear

Stain

Signature of the tutor_____

Additional materials for class 4

Causative agents	Disease	Materials for bacteriological diagnostics
K. pneumoniae s. rhinoscleromatis		
K. pneumoniae s. ozaenae		
K. pneumoniae s. pneumoniae		
Y. enterocolitica		
C. jejuni		
H. pylori		
P. aeruginosa		

Methods of laboratory diagnostics

	Usage (+/-)						
Method	Klebsiella	Campylo- bacter	Iersinia	Pseudomonas aeruginosa			
Microscopic							
Cultural							
Biological							
Serological							
Allergic							
Molecular-							
genetic							

Diagnosis of bacterial food poisoning

Food poisoning - acute systemic diseases resulting from ingestion of food, massively contaminated with microorganisms or microbial exotoxins. Food poisoning is divided into bacterial foodborne diseases and food poisoning (toxicosis), as well as poisoning of mixed etiology.

Foodborne diseases (FBD): FBDs result from ingestion of products massively colonized by certain bacteria. Pathogens: opportunistic members of the family Enterobacteriaceae - E. coli, Proteus (P. vulgaris, P. mirabilis), Morganella morganii, Citrobacter, Enterobacter, Hafnia, Klebsiella pneumoniae; Sem. Vibrionaceae - V. parahaemolyticus; Sem. Bacillaceae - B. cereus, C. perfringens serovar A; Sem. Streptococcaceae - S. faecalis; Sem. Pseudomonadaceae - P. aeruginosa, and others.

Microbial food toxicosis (intoxication): acute illness arising from eating food, which containes a large amount of exotoxin (as a result of massive reproduction of microbes). These include botulism, toxicosis caused by staphylococcal enterotoxin, toxins from microscopic fungi and others.

Pathogenesis. Pathogen replicates in the intestine, penetrates into lymphoid tissue, where it is killed with the release of endotoxin, which causes damage to the intramural bowel NS, CNS and blood vessels. Bacteria cause inflammation of the intestinal wall.

Pathogenesis is based on the microbial exotoxin, which is not destroyed by food processing, digestive enzymes and acidic stomach contents.

Materials for the research: vomit, stomach washings, feces, urine, blood, sectional material (in the case of death), the remains of the suspected food, raw and semi-finished products used, daily samples of food, swabs and scrapings from kitchen utensils.

Lab. diagnosis: isolation of obligate pathogenic or opportunistic enterobacteria and Vibrio, staphylococci and their toxins, streptococci, bacillus, as well as (if indicated) - botulism pathogens and toxins.

To evaluate the etiologic role of opportunistic bacteria (OB) certain criteria are used.

Main criterion is quantitative: Etiologically significant number of OM is 10⁵-10⁶ or more CFU per 1g of material. The diagnosis is more reliable while simultaneous detecting same germs or toxins in suspected food. Other criteria are: repeated isolation of same germs from the material of the patient, the identity of the pathogen strains (serovars and phage-vars) in a large number of patients in group food poisoning, as well as the increase in antibody titer in the dynamics of the disease.

Class № 5. Microbiological diagnostics of diseases caused by Corynebacteria, bordetella, haemophilus, legionella, listeria Date_____

List of questions to study:

Corynebacterium diphtheria. Systematics, general characteristics of the pathogen. Types of Corynebacterium diphtheria, their distinctive features. Diphtheria toxin and antitoxic serum. The pathogenesis of diphtheria. Methods of microbiological and molecular biological diagnosis of diphtheria. Principles of therapy and prevention of diphtheria. Determination of the effectiveness of post-vaccinal immunity.

Bordetella pertussis. Characteristics of the pathogen, pathogenicity factors. Differentiation with parapertussis agent. The pathogenesis of pertussis, immunity, diagnostics. Principles of therapy and prevention of pertussis.

Haemophilus, general characteristics, role in human pathology.

Legionella, general characteristics, role in human pathology.

Listeria, general characteristics, role in human pathology.

Laboratory work

Tasks Methods. results 1. Bacteriological diagnosis of diphtheria, 2nd step: a) Describe the colonies Corynebacterium on potassium tellurite serum agar b) Seed bacteria from typical colonies onto Hiss media (glucose, sucrose, starch). Smear **Smear** Stain Tinsdale medium Stain Glucose Sucrose Starch Demonstration. 1. Corynebacterium diphtheriastained by: a) Neisser: b) Leffler. Colonies on serum tellurite agar **Feature** 2. Test for Corynebacterium diphtheria Shape toxigenicity. Size 3. Preparations for specific prevention and Surface treatment of diphtheria and pertussis. 4. Growth of Bordetella pertussis and parapertussis Edge Smear Smear on CCA, NA with tyrosine, urease test. Color 5. Bordetella pertussis, Gram staining Consistency Stain Stain 6. Assessment of antidiphtheria immunity intensity

Additional materials and independent work for Class N_2 5.

Corynebacterium char	racteristics
Properties	C. diphtheriae
Morphology (size, shape, relative	
positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Pathogenicity factors	

Medically important corynebacteria

Species	Diseases
C. diphtheriae	Diphtheria
C. ulcerans, C. minutissimum, C. xerosis, C. pseudodiphtheriticum	Opportunistic infections

C. diphtheriae pathogenicity factors

Pathogenicity factors	Biological effect		
Protein exotoxin (includes	Protein synthesis arrest, specific damage		
A and B subunits)	of the myocardium, adrenal glands and		
	nerve ganglia		
Glycolipid (6-6'-diester-	Phagocytosis impairment		
trehalose)	i nagocytosis impairment		
Hyaluronidase	Darmachility of tissues violation		
Neuraminidase	Permeability of tissues violation		

Laboratory diagnostics and specific prophylaxis of diphtheria

Method	Properties
Microscopic	
Cultural	
Molecular-genetic	
Specific prophylaxis	

Bordetella pertussis characteristics		
Properties	B. pertussis	
Morphology (size, shape, relative positions of cells)		
Spores development		
Capsule		
Flagella (motility)		
Gram staining		

Bordetella differentiation

Feature	B. pertussis	B. parapertussis

B. pertussis pathogenicity factors

Pathogenicity factors	Biological effect
Filamentous hemagglutinin	Binds cell membrane glycolipid of ciliated airway epithelium, binds surface R3 - glycoprotein receptor and initiates phagocytosis
Pertussis toxin (Pertussin)	S1 Pertussin subunit ribosylates membrane protein Gi; toxin inhibits the activity of phagocytes and monocyte migration. S2 - subunit binds to the respiratory tract cell surface glycolipid; S3 - subunit binds to phagocytes surface gangliosides
Pili	Adhesion to the ciliated epithelium of the respiratory tract
Pertactin	Adhesion to the ciliated epithelium of the respiratory tract
Adenylate cyclase	Suppresses killing- activity of phagocytes and monocytes migration
Dermatonekrotoksin	Damages the skin and is lethal to laboratory animals
Tracheal toxin	peptidoglycan fragment - destroys ciliated cells of the respiratory tract; stimulates interleukin-1 secretion (fever)
Endotoxin (LPS)	Activates complement and stimulate the production of cytokines

Laboratory diagnostics and specific prophylaxis of pertussis

Method	Properties
Bacteriological	
Serological	
Specific prophylaxis	

Haemophilus genus representatives and respective diseases		
Diseases		

Haemophilus genus characteristics

Properties	H. influenzae
Morphology	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Antigens	

H. influenzae pathogenicity factors

Pathogenicity factors	Biological effect
Polysaccharide capsule	Inhibition of phagocytosis
Pili and other adhesins	Attaching to epithelial cells
Lipopolysaccharide and glycopeptide	Epithelium surface and cilia damage
Ig A protease	Suppression of local immunity

Laboratory diagnostics and specific prophylaxis of infections caused by Haemophilus

Method	Properties
Microscopic	
Cultural	
Serological	
Specific prophylaxis	

Legionella characteristics		
Properties	Legionella pneumophila	
Morphology (size, shape, relative		
positions of cells)		
Spores development		
Capsule		
Flagella (motility)		
Gram staining		

Legionella pneumophila pathogenicity factors

Legionetta pheumophita patnogemeny factors		
Pathogenicity factors	Biological effect	
Optional intracellular parasitism		
Toxin (peptide)	inhibiting the "oxidative burst" during phagocytosis	
Catalase	inactivation of toxic metabolites during macrophage activation	
Factors of unknown nature	inhibit fusion of phagosomes and lysosomes, electron transport	
2. Production of toxins, enzymes		
Labile exotoxin (Cytotoxin and hemolysin)	dysfunction or cell lysis	
Endotoxin	dysfunction or cell lysis	
Proteolytic enzymes: phosphatase, lipase, nuclease	degradation of host cells	
2 Suppression of the expression of MHC class II melacules on meanning as		

3. Suppression of the expression of MHC class II molecules on macrophages, violation of Ag-presenting functions - the suppression of cellular immune response

Laboratory diagnostics and specific prophylaxis of legionellosis

Method	Properties
Microscopic	
Cultural	
Serological	
Molecular-genetic	
Specific prophylaxis	

Listeria characteristics

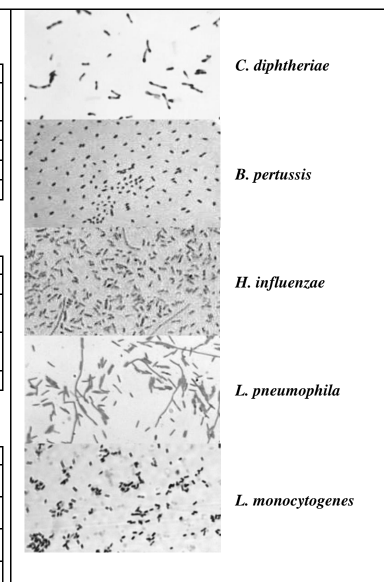
Properties	L. monocytogenes
Morphology (size, shape, relative positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	

Listeria pathogenicity factors

Pathogenicity factors	Biological effect
Endotoxin	Toxic effects -
Internalin - membrane protein	Listeria entry into macrophages and endothelial cells, (from phagosome into the cytoplasm)
listeriolysin O	Hemolysin, cause phagolysosomes membrane disruption
Phospholipase	Membrane damage and penetration into the cell

Laboratory diagnostics and specific prophylaxis of listeriosis

Method	Properties
Microscopic	
Cultural	
Serological	
Molecular-genetic	
Specific prophylaxis	



<u>Class № 6.</u> Methods of microbiological diagnosis of diseases caused by mycobacteria and actinomycetes. Methods of microbiological diagnostics anaerobic infections

Data
Data

The list of questions to study:

Actinomycetes, systematic position, general characteristics, role in human pathology.

Mycobacteria classification. TB germs, general characteristics. Pathogenesis, immunity, methods of microbiological diagnostics, principles of treatment and prevention of tuberculosis. Mantoux test.

The causative agent of leprosy, general characteristics, role in human pathology.

Mycobacteriosis. Nocardia.

Anaerobes, classification, general characteristics. Causative agents of gas gangrene, tetanus, botulism. Systematics and general characteristics. Exotoxins.properties Principles of therapy and prevention of anaerobic infections.

Clostridial gastroenteritis. Clostridium difitsile role in human pathology.

Nonspore anaerobes. Bacteroides. Peptococci. General characteristics, pathogenicity factors, role in human pathology.

General principles and methods for anaerobic infections diagnosis. Molecular biological diagnostics - PCR.

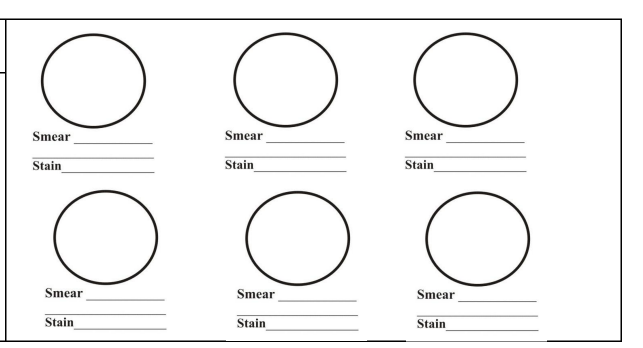
Laboratory work

Tasks	Methods, results							
1. The assessment of enzymatic activity of corynobacteria,		Bioche	emical pr	operties	of sert	ain coryno	bacteria	ı
identification		Corynobacteria	Enzymatic activity				Nitrate	
		spp.	Glucose	Sucrose	Starch	Cysteinase	Urease	reduction
		C. diphtheriae						
		gravis mitis	+ +	-	+			+
		C.	+	-	-			+
		pseudodiphtheriae (hofmani) C. xerosis C. ulcerans	- + +	- + -	- - +	- - +	+ + +	+ + -
	Conclusion: according to morp bacterium is identified as		and bioc	nemical j	properti	es unknowi	1	

2.	Microscopy	of ready	smear	of	tuberculosis	patient
sp	utum, Ziehl-l	Neelsen st	aining.			

Demonstration.

- 1 Mycobacteria growth on nutrient media.
- 2. Flotation method
- 3. Determination of M. tuberculosis drug resistance
- 4. Cord factor of M.tuberculosis, Ziehl-Neelsen staining.
- 5. Actinomycetes, pure culture, Gram staining.
- 6. M. leprae, Ziehl-Neelsen staining.
- 7. M.tuberculosis in sputum, Ziehl-Neelsen staining.
- 8. Anaerobes growth on nutrient media.
- 9. Clostridium, Gram staining.
- 10. Bacteroides, Gram staining.



Signature of the tutor_

Description

Materials for independent work for class N_2 6

Microbiological diagnostics and specific prophylaxis of **Actinomyces characteristics** actinomycosis Characteristics Actinomyces israelii Morphology (size, Method shape, relative positions Microscopic of cells) Spores development Cultural Capsule Flagella (motility) Specific Gram staining prophylaxis Pathogenicity factors

Classification of medically important culturable mycobacteria

	Slowly growing		Fast gre	owing
Tuberculosis agents	Non chromogenic	Chromogenic	Non chromogenic	Chromogenic
M. tuberculosis M. bovis M. africanum	M. avium complex M. xenopi M. haemophilum et al.	M. kansasii M. marinum M. simae et al.	M. fortuitum M. chelonae M. smegmatis et al.	M. phlei M. vaccae

Myvobacteria characteristics

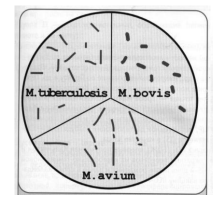
Characteristics	M. tuberculosis	M. leprae
Morphology (size,		
shape, relative positions		
of cells)		
Spores development		
Capsule		
Flagella (motility)		
Gram staining		
Pathogenicity factors		

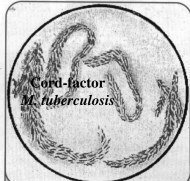
M. tuberculosis pathogenicity factors

Pathogenicity factors	Biological effects
Cord-factor	
(trehalose-6,6-dimycolate)	
Sulphatides (sulfur-containing	
glycolipids)	
Antigens	

Microbiological diagnostics and specific prophylaxis of tuberculosis

Methods	Remarks
Microscopic	
Cultural	
Serological	
Biological	
Molecular-genetic	
Allergic	
Specific prophylaxis	





Microbiological diagnostics and specific prophylaxis of leprosy

Methods	Renarks
Microscopic	
Allergic	
Biological	
G 131	
Specific prophylaxis	
prophylaxis	

Ecological group of anaerobic bacteria				Clostridia characteristics						
Gran	n-negative			Characteristics		C. perfringens	C. tetani	C. botulinum		
	oreing rods	Di	seases induced	Morphology (size, shape,						
Bacteroide				Π		ative positions				
Fusobacter	rium species			T	Spo	ores developme	ent			
Leptotrichi				П		psule				
Prevotella	species			Π	_	gella (motility)				
Porphyron	onas species			П	_	am staining				
Bilophila v	vadsworthia			П	Pat	hogenicity fact	ors			
Gramposi	tive spore form	ning rods								
	Clostridium tei	tani	Tetanus (Lockjaw)							
		erfringens, C. novyi,	Gas gangrene, necrotizing			Clostri	dium per	fringens pathog	genicity fact	tors
C. ramosum, C. histolyticum, enteritis C. septicum C. septicum C. septicum C. septicum C. septicum C. septicum		athogenicity factors		Biolog	ical effects					
	Clostridium bo	otulinum	Botulism			alpha-toxin	cleaves lecithin in cell membranes; increase		ncreases	
	Clostridium dij	fficile	Pseudomembranous colitis, antibiotic-associated diarrhe	ous colitis, (Lecithinase) vascular permeability destroying		stroying ery	oying erythrocytes;			
Gramnegative cocci		Ħ	☐ ≦ beta-toxin		necrotizing activity; induction of hypertension as a					
Veillonella Septic infections		Main toxins		octa-toxiii		formation of catecholamines				
Grampositive cocci				epsilon toxin	increases vascular permeability of the		ie			
Enterococo	cus species				Maj	- F	_	testinal tract		
Peptococcus species Septic infections Peptostreptococcus spp.					Iota toxin	necrotizing activity and increased vascular permeability				
Bacteroides pathogenicity factors		\parallel		enterotoxin	violates the permeability of the mucosa of intestine		osa of the small			
Pathogenicity factors Biological effect				delta-toxin	hemolysis					
toxins	endotoxin	general toxic effect				theta toxin	hemolysis, cytolysis			
LOAIIIS	leukocidin	damages leukocytes			in	kappa toxin	collagenase, gelatinase, necrotizing activity			activity
	collagenase		gen fibers of the connective		tox	lambda-toxin	protease		<u>U</u>	-
		tissue (spread of p	urulent process)		Minor toxin	mu-toxin	hyaluror	nidase: increases	the permea	bility of tissues
enzymes	DNAse, heparinase	cause intravascula	r blood clotting		Mi	nu-toxin		hemolytic, necres gangliosides ce		
	fibrinolysin	dissolves blood cle	ots			neuraminidase	_	s gangnosides ce osis in capillaries		promotes

	beta-lactamase	destroys the be	ta-lactam antibiotics					
surface cell	pili	adhesion to the	substrate	Clos	tridium botulinum pathogenicity factors			
structure	capsule	protects the bact	eria from phagocytosis					
Metabolites	fatty acid	inhibit the chem leukocyte	otaxis and cytotoxicity of	Pathogenicity factors	Biological effects			
Microbiolo	ogical diagnosti		ctions caused by bacteroides	Botulinum	Blocks the transmission of nerve impulses in the peripheral cholinergic synapses, providing			
N	lethod	Remarks		exotoxin	neurotoxic effects (lethal dose for humans is			
Microscopi	c			CAOLOAIII	about 0.3 g)			
Cultural								
Serological								
Molecular-	genetic							
			prophylaxis of gas gangrene	Microbiologica	al diagnostics and specific prophylaxis of botulisn			
Method		R	emarks	Methods	Remarks			
Microscopi	c			Serological				
Cultural				Biological				
Biological				Cultural				
Specific prophylaxis				Specific	Botulinum toxoids A, B, E, are used according to			
	•	n tetani pathoge	enicity factors	prophylaxis	indications. For urgent passive prophylaxis specif antitoxic serum is used.			
Pa	thogenicity facto		Biological effects	7				
Tetanus tox	in		<u> </u>]				
Microb	iological diagno	ostics and speci	fic prophylaxis of tetanus	_				
Method		R	emarks	<u> </u>				
Microscopi	c			<u> </u>				
Biological				7				

<u>Class № 7.</u> Microbiological diagnostics of especially dangerous infections

Date

The list of questions to study:

Classification and general characteristics of the especially dangerous infections. Demands to collection and transportation of biological material. Principles of diagnostics.

Vibrio cholerae, the systematic position. Classification and general characteristics, pathogenicity factors. Biovars. Differentiation from non-cholera vibrio. Pathogenesis of cholera. Methods of microbiological diagnostics. Rapid methods. Principles of treatment and prevention.

The causative agent of plague, systematic position, characteristics, pathogenicity factors. Differences from other Yersinia. Pathogenesis, principles of treatment and prevention of plague.

The causative agent of tularemia, systematics, general characteristics. Pathogenesis, principles of treatment and prevention.

Causative agents of brucellosis. Systematics and general characteristics, pathogenicity factors, pathogenesis. Microbiological diagnosis of brucellosis. Principles of treatment and prevention.

Anthrax. Systematics and general characteristics, pathogenicity factors. Differences from non-pathogenic bacilli. Pathogenesis. Microbiological diagnosis of anthrax. Principles of treatment and prevention.

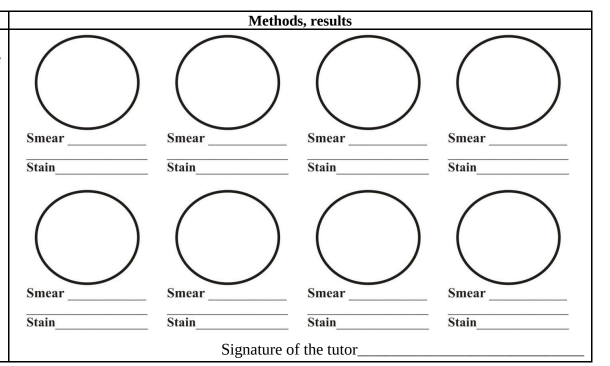
Laboratory work

Demonstration.

1. Growth of vibrio cholera on alkaline agar, TCBS, peptone water.

Tasks

- 2. Phage lysability of vibrio cholera classica and El Tor.
- 3. Tube agglutination test.
- 4. Biochemical properties of V. cholerae.
- 5. Mobility of Vibrio spp.
- 6. V.cholera, pure culture, Gram staining.
- 7. I.pestis in the organs, Leffler staining.
- 8. The causative agent of tularemia (pure culture), Gram staining.
- 9. Preparations for specific prophylaxis of especially dangerous infections.
- 10. The causative agent of brucellosis, Gram staining.
- 11. The growth of Bacillus spp. on nutrient media.
- 12. B.anthracis in organs, Gram staining.
- 13. B.anthracis, pure culture, Gram staining.
- 14. B.anthracis spores, Ozheshko staining.



Additional materials for independent study for class №7

V. cholera characteristics

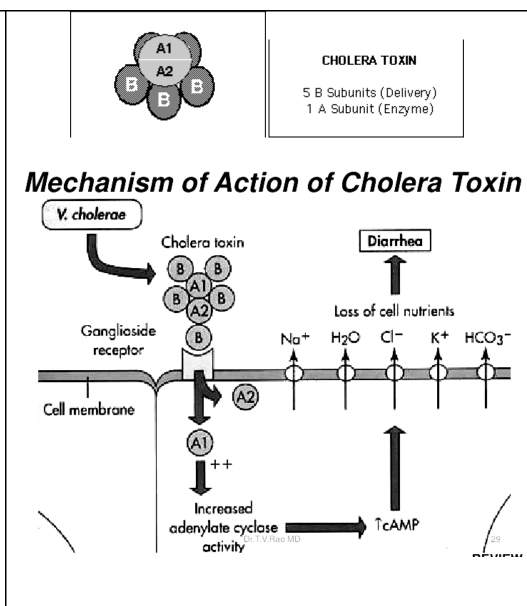
Characteristics	Vibrio cholerae
Morphology (size, shape, relative positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Pathogenicity factors	

Vibrio cholerae pathogenicity factors

Pathogenicity factors	Biological effects
Exotoxin (choleragen)	Violation of water-salt metabolism, the cytotoxic effect on the epithelium of the small intestine
Endotoxin	Inhibition of phagocytosis, drop in blood pressure; infectious-toxic shock
Pili	Adhesion to mucosal cells
Fibrinolysin hyaluronidase	Enzymes invasion (aggression)

Microbiological diagnostics and specific prophylaxis of cholera

Method	Remarks
Microscopic	
Cultural	
Serological	
Molecular-genetic	
Specific prophylaxis	



I. pestis characteristics

Characteristics	Yersinia pestis
Morphology (size, shape,	
relative positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Pathogenicity factors	

Y. pestis pathogenicity factors

Pathogenicity factors	Biological effects
Capsular Ag, F1-Ag,	protection against the absorption of
fraction 1)	phagocytes, non-toxic, the immunogen
Plasminogen	activates lysis of fibrin clots, and inactivates
activator - protease	C5a
V/W(Vi)-Ag	Includes protein (V-phase) and LPS (W-phase); exhibits antiphagocytic properties, promotes intracellular bacterial growth
Murine toxin	adrenergic receptor antagonist, proteinaceous substance, localizes intracellularly
Bacteriocins (pestitsiny)	Immunogenic properties

Microbiological diagnostics and specific prophylaxis of plague

Method	Remarks
Microscopic	
Cultural	
Molecular-genetic	
Biological	
Specific prophylaxis	

F. tularensis characteristics

Characteristics	Francisella tularensis
Morphology (size, shape,	
relative positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Pathogenicity factors	

F. tularensis pathogenicity factors

Pathogenicity factors	Biological effects
Intracellular parasitism	Inhibition of phagocytes lysosomal function,
Capsule	Protection from phagocytosis
Endotoxin	Less active than other Gram-negative rods endotoxin (e.g., E. coli)

Microbiological diagnostics and specific prophylaxis of tularemia

Method	Remarks
Microscopic	
Cultural	
Serological	
Molecular-genetic	
Allergic	
Biological	
Specific	
Specific prophylaxis	

Brucellosis agents characteristics

Characteristics	Brucella spp.
Morphology (size, shape,	
relative positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Pathogenicity factors	

Brucella pathogenicity factors

Pathogenicity factors	Biological effects
Endotoxin	Systemic toxic effect
Hyaluronidase	Breaks down hyaluronic acid
Outer Membrane Proteins	Adhesion

Microbiological diagnostics and prophylaxis of brucellosis

Method	Remarks
Microscopic	
Cultural	
Serological	
Allergic	
Molecular-	
genetic	
Biological	
Specific	
prophylaxis	

Anthracs pathogen characteristics

Characteristics	B. anthracis
Morphology (size, shape,	
relative positions of cells)	
Spores development	
Capsule	
Flagella (motility)	
Gram staining	
Pathogenicity factors	

Bacillus anthracis pathogenicity factors

Pathogenicity factors	Biological effects	
Protein	Exotoxin contains three factors:	
exotoxin	lethal factor - the cytotoxic effect, pulmonary edema,	
(synthesis is	protective Ag - interacts with cell membranes mediates the	
controlled	activity of others. components, edematous factor - the increase	
plasmid)	in the concentration of cAMP, the development of edema	
Capsule	Antiphagocytic activity	

Microbiological diagnostics and specific prophylaxis of anthrax

Method	Remarks
Microscopic	
Cultural	
Serological	
Allergic	
Molecular-genetic	
Biological	
Specific prophylaxis	

Class № 8 Microbiological diagmostics0f diseases caused by spirochetes

Date

List of questions to study:

Spirochetes, classification, general characteristics.

Treponema. Systematics and general characteristics. Pathogenesis and immunity in syphilis. Material for the study. Methods of microbiological diagnosis of syphilis. Principles of therapy and prevention of syphilis.

Fusospirochetosis pathogens

Leptospira. Systematics and general characteristics. Pathogenesis, methods of microbiological diagnostics, principles of treatment and prevention of leptospirosis.

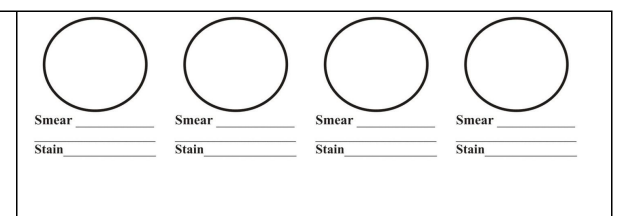
Borrelia. Systematics and general characteristics. Pathogenesis and methods of microbiological diagnosis of relapsing fever. The causative agent of Lyme borreliosis, principles of treatment and prevention.

Laboratory work

Tasks	Methods, results
1. Perform the slide microprecipitation reaction (VDRL) for the syphilis serodiagnosis.	Slide microprecipitation test
2. Assess ELISA (Wasserman test) for the syphilis diagnostics.	1. Patient serum 1:20 2. Saline sol. 3. Cardiolipin Ag
	Conclusion:

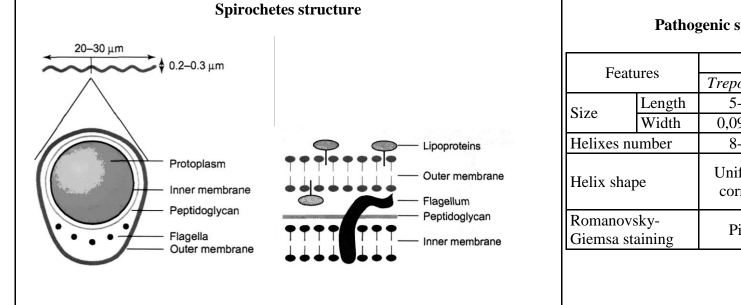
Demonstration.

- 1. Leptospires, dark field microscopy.
- 2. Borrelia in blood, Romanovsky-Giemsa staining.
- 3. Wasserman test (ELISA).
- 4. Treponema in dental plaque, Gram staining.
- 5. Treponema pallidum, pure culture, Romanovsky-Giemsa staining.



Signature of the tutor_____

Additional materials for independent study for class N_2 7.



Pathogenic spirochetes characteristics

Features		Spirochetes genera			
		Treponema	Borrelia	Leptospira	
Size	Length	5-20	3-20	7-14	
Size	Width	0,09-0,5	0,2-0,5	0,1-0,15	
Helixes number		8-12	2-8	12-24	
Helix shap	pe	Uniform, correct,	Uneven, different size	Uniform, correct secondary curls	
Romanovs Giemsa st	•	Pink	Blue Purple	Pink, Red	

Diseases caused by treponema

Treponema spp.	Disease	Morbidity (countries, continents)
T. pallidum, subspecies pallidum		
T. pallidum, subspecies bedjel (endemicum)		
T. pallidum, subspecies pertenue		
T. carateum		
Opportunistic or saprophytic:		
T. vincentii, T. refringens, T. denticola,		
T. minutum, T. scoliodontum		

Methods for spirochetosis diagnostics

		Me	thod usage (+/-)	
Methods	Syphilis	Epidemic relapsing fever	Endemic relapsing fever	Lyme disease	Lepto- spirosis
Microscopic					
Cultural					
Serological					
Allergic					
Molecular-genetic					
Biological					

	Pathogenesis	s of syphilis
Disease stage	Period	Main pathogenetic mechanisms
Primary		
Secondary		
Tertiary		

Serological diagnosis of syphilis:

CFT (Wasserman) with treponemal and cardiolipin antigens in primary syphilis becomes positive in the 6th week of the disease in 25-50% of patients, in 7-8 weeks - 75-90%. In secondary syphilis it is positive in 98-100% cases. In tertiary syphilis CFT is positive in only 60-70% patients. CFT for syphilis diagnostics has unsatisfactory sensitivity and specificity and is replaced now by ELISA.

ELISA is the common used technics for syphilis diagnostics. Confirmatory tests:

- treponema immobilization test is rather specific, but time and labor consuming, subjective, requires treponema culture;
- immunofluorescence (IF) with serum from patients.

Screening tests: slide microprecipitation test, ELISA

Laboratory diagnosis of Lyme disease (Lyme borreliosis):

Microscopic method: dark-field microscopy (scrapings of skin lesions, plasma pellet, CSF, urine), microscopy of smears, impregnated with silver, IFT, and electron microscopy.

Cultural method: B. burgdorferi isolation is possible in 80% cases from skin lesions (1stage) on special nutrient media.

Molecular genetic methods: PCR allows the identification of the pathogen's DNA in the samples of the skin, blood, cerebrospinal fluid.

Serological: ELISA, indirect IFT, Western blot. Sometimes there are false-positive results due to cross-reactions among patients with syphilis, mononucleosis, rheumatoid arthritis and others.

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Class № 8.	Microbiological diagnostics of dise	ases caused by Rickettsia.	Chlamydia and Myconlasma
Clabb t 1= 01	where obtaining teat and should of the	uses caused by inclicible,	Cilium y and and my copiusing

Date

List of questions to study:

Rickettsiae, systematic position, classification, general characteristics, role in human pathology. Rickettsia typhii, pathogenesis, immunity and methods of microbiological diagnostics. Other pathogenic rickettsia.

Chlamydia, general characteristics, role in human pathology. Pathogens of psittacosis, trachoma, respiratory and urogenital chlamydiosis. Methods of microbiological diagnosis of chlamydiosis. PCR in chlamydiosis diagnostics.

Mycoplasma, general characteristics, role in human pathology. Methods of microbiological diagnostics of mycoplasmoses.

Laboratory work

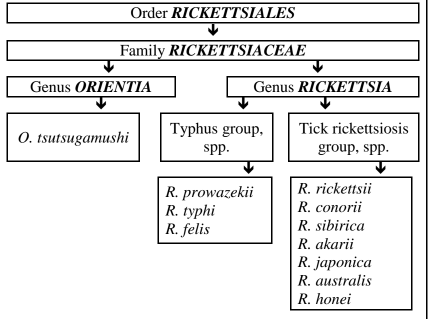
Tasks	Methods, results 0,5										
1. Perform CFT for the typhus diagnostics	Reagents	1	2	3	4	5	6	7	SC	DC	Hemolytic
		1:5	1:10	1:20	1:40	1:80	1:160	1:320			system
	Saline sol.	-	0,5	0.5	0.5	0,5	15	0.5	0,5	0,5	4 ml 3%
	Serum of the patient	0,5	0,5	Ņ	- <u>-</u> -				0,5	-	erythrocytes
1	Diagnosticum	0,5	0,5	0,5	0,5	0,5	0,5	0,5	-	0,5	suspension + 4
	Complement	0,5	0,5	0,5	0,5	0,5	0,5	0,5	0,5	0,5	ml hemolytic
	Incubation 30 minut at 37° C										serum
	Hemolytic system										
Smear	Incubation for 30' at 37 °C										
Stain	Assessment										
	Conclusion:										
Demonstration.	1/10 1/20 1/40	1/80	1/160 1/	320 1/6	40	SC	DC				/
 Passive blood aggl;utination test for differential diagnostics of epidemic and residual typhus Chlamydia spp. in cell culture, Romanovsky-Giemsa staining. R. prowazeki, pure culture, Zdrodovski staining. 	Conclusion:								Smear Stain_		

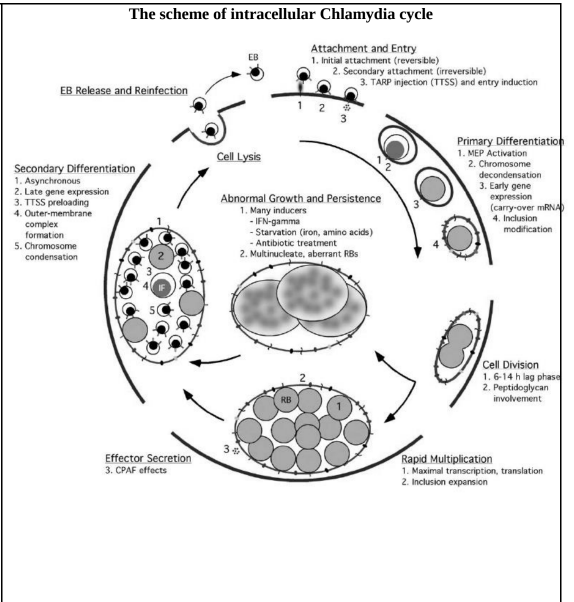
Additional materials for independent work wor class № 8

Actual classification of Rickettsia:

On the basis of a molecular genetic studies (genome sequencing, PCR) classification of microorganisms belonging to the order Rickettsiales has undergone significant changes. The genus Coxiella with C.burnetti was excluded from the family and added in the order Legionellales, family Coxiellaceae. Genus Rochalimaea was removed, and its representatives - R. quintana (Trench fever) and R. henselae (cat scratch disease) were included in the family Bartonellaceae, genus Bartonella.

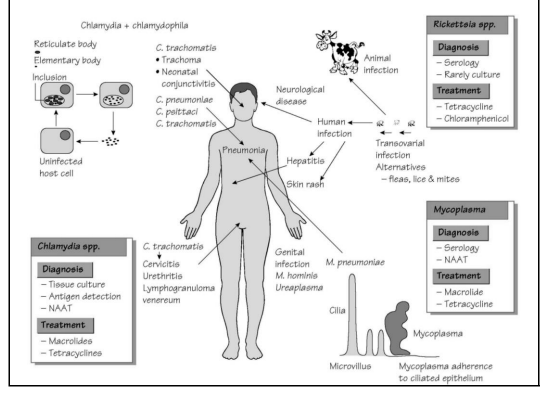
The family Rickettsiaceae now include three genera: Rickettsia, Orientia, Wolbachia. The medical importance of the latter genus is still unclear.





Laboratory diagnostics of diseases caused by Rickettsia, Chlamydia and Mycoplasma

Method		Method usage			
		rickettsiosis	chlamydiosis	mycoplasmosis	
Microscopic					
Cultural	Nutrition media				
	Chicken embryo				
	Cell culture				
	Lab animals				
Biological					
Serological					
Allergic					
Molecular-genetic					



Chlamydiosis characteristics

Disease	Pathogen	Source	Transmission
Trachoma			
Urogenital chlamidiosis			
Veneral lymphogranulomas			
Psittacosis			
Pharyngitis, sinusitis, bronchitis, pneumonia			

Mycoplasma and mycoplasmosis characteristics

Properties -	Mycoplasma spp
Size	
Cell wall, peptidoglican	
Gram staining	
Capsule	
Flagella	
Spore	
Resistance in environment	
Cultural properties	
Reproduction	
Parasitism peculiarities	
Source of infection	
Transmission mechanisms	
Immunity	

<u>Class № 9</u> Concluding test "Special microbiology"

- 1. Staphylococci, general characteristics. Role in human pathology. Pathogenicity factors and mechanisms of pathogenesis of staphylococcal infections. Microbiological diagnosis. Principles of treatment and prevention of staphylococcal infections.
- 2. Streptococci, classification. General characteristics. Pathogenicity factors. Antigenic structure. Pathogenesis, immunity, microbiological diagnosis, principles of treatment and prevention of streptococcal infections.
- 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.
- 5. General characteristics of the family Enterobacteriaceae.
- 6. General Principles of bacteriological diagnosis of acute intestinal infections (AII). The nutrient medium for enterobacteria. Classification principles of application.
- 7. Materials for researches in AII diagnostics.
- 9. E. coli, common characteristic. The biological role of Escherichia coli. Diseases caused by Escherichia.
- 10. Salmonella. General characteristics. Members of the genus. Serological classification by Kaufmann-White. Molecular biological typing.
- 11. Pathogens of typhoid, paratyphoid A and B, general characteristic. Phage typing. Vi-antigen and its value.
- 12. Pathogenesis and methods of microbiological diagnosis of typhoid and paratyphoid.
- 13. Immunity in typhoid fever. Serological diagnosis of typhoid and paratyphoid. Specific prophylaxis.
- 14. The etiology of food poisoning and intoxication of bacterial origin. Materials and methods of diagnosis.
- 15. Salmonellosis. Characteristics of pathogens and diagnostic methods. Nosocomial salmonellosis.
- 16. Shigella. Classification. Characteristics. Pathogenesis, immunity. Methods of microbiological diagnostics of acute and chronic dysentery.
- 17. Klebsiella. Classification, general characteristics. Pathogenesis, immunity, methods of microbiological diagnostics of klebsiellosis.
- 18. Pseudomonas aeruginosa, general characteristics, pathogenicity factors. Role in human pathology.
- 19. Pathogens of intestinal yersiniosis, general characteristics. Pathogenesis. Methods of diagnosis of yersiniosis.
- 20. C.diphtheria, general characteristics. Differences from non-pathogenic corynebacteria. Mechanisms of pathogenesis and microbiological diagnosis of diphtheria.
- 21. Diphtheria toxin and its properties. Toxoid. Immunity in diphtheria and its character. Determination of antitoxic immunity. Principles of therapy and prevention of diphtheria.
- 22. The causative agent of whooping cough, general characteristics. Differentiation with parapertussis agent. Pathogenesis, immunity. Microbiological diagnosis, principles of treatment and prevention of pertussis.
- 23. General characteristics of the causative agents of tuberculosis. Pathogenesis, immunity, diagnosis and specific prevention of tuberculosis. Mycobacteriosis.
- 24. The causative agent of leprosy. Characteristic, pathogenesis, immunity.

- 25. Particularly dangerous infections. classification mode, Basic rules of sampling, sending and transportation of infectious material General principles of diagnosis TELO.
- 26. V. cholera. Systematics. General characteristics. Differentiation of biovars. Pathogenesis, immunity, principles of treatment and prevention. Methods of microbiological diagnostics.
- 27. The causative agent of plague, a general characteristic. The pathogenesis of plague. Immunity, the principles of therapy and prevention of plague.
- 28. B. anthracis characteristic. Pathogenesis, immunity, principles of treatment and prophylaxis of anthrax.
- 29. The causative agent of tularemia, general characteristic. Pathogenesis, immunity, principles of treatment and prophylaxis of tularemia.
- 30. Pathogens of brucellosis, a general characteristic. Differentiation of Brucella species. Pathogenesis, immunity, principles of treatment and prevention of brucellosis.
- 31 Spirillae family. Campylobacter, characteristics, role in human pathology. Helicobacter.
- 32. Classification and general characteristics of anaerobes. Clostridia. Bacteroides, Peptococci and other nonspore anaerobes. Pathogenicity factors. Role in human pathology.
- 33. The causative agent of tetanus, general characteristics. Pathogenesis, immunity, principles of treatment and prevention of tetanus.
- 34. Gas gangrene pathogens, general characteristics. Pathogenesis, principles of treatment and prevention of gas gangrene.
- 35. The causative agent of botulism, general characteristic. Pathogenesis, principles of botulism prevention and therapy. Clostridial gastroenteritis.
- 36. Methods of diagnosis of anaerobic infections.
- 37. Classification and general characteristics of spirochetes.
- 38. Classification of treponemes and treponemal diseases. Characteristics of syphilis causative agent. Pathogenesis, immunity, diagnostic tests for syphilis.
- 39. Leptospires. General characteristics. The pathogenesis of leptospirosis, immunity, specific prevention. Microbiological diagnosis of leptospirosis.
- 40. Borrelia, general characteristics. Recurrent fever pathogenesis, immunity. Microbiological diagnosis. The causative agent of Lyme borreliosis.
- 41. Systematic position and characterization of Rickettsia. Pathogenesis, immunity, methods of diagnosis of typhus.
- 42. Characteristics of chlamydia. Causative agents of trachoma, psittacosis, respiratory and urogenital chlamydiosis. Pathogenesis and methods of diagnosis of chlamydia.
- 43. General characteristics of mycoplasma, pathogenicity factors, role in human pathology. Methods of mycoplasmosis 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.
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<u>Class № 10.</u> Clinical microbiology. Microbiological diagnostics of sepsis and purulent infections of the skin

Date

List of questions to study:

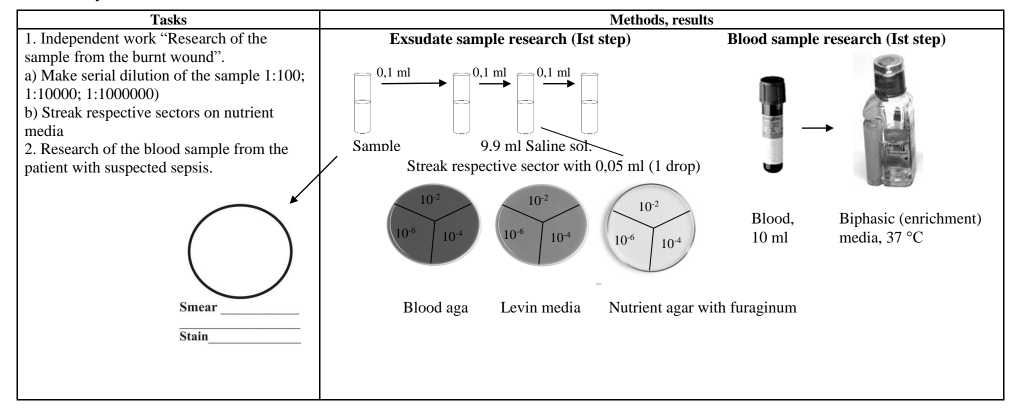
Clinical Microbiology: definition, objectives. Opportunistic microbes (OPM). Epidemiology, pathogenesis, diagnosis of diseases caused by UPM. Criteria of etiological significance.

Clinical forms and the etiology of septic infections of the skin and subcutaneous tissue. Methods of microbiological diagnostics.

Bacteriological method. Material for the research (pus, exudate), rules and methods of sampling. Criteria for assessment of the etiological significance of isolated microorganisms. Susceptibility to antibiotics.

Bacteremia. Sepsis. Pyosepticemia. Etiology, definitions. Methods of microbiological diagnosis of sepsis. Bacteriological method. Rules and methods of blood collection for the research, peculiarities of pathogen isolation and results interpretation Susceptibility to antibiotics testing.

Laboratory work



3. Research of the sample from the bronchi washings:

a) Prepare the slide from the material, Gram staining;
b) perform quantitative seeding of the

material on selective media.

Sample research (Ist step)

Sample 9.9 ml Saline sol.
Streak respective sector with 0,05 ml (1 drop)

Smear

Stain

Blood agar

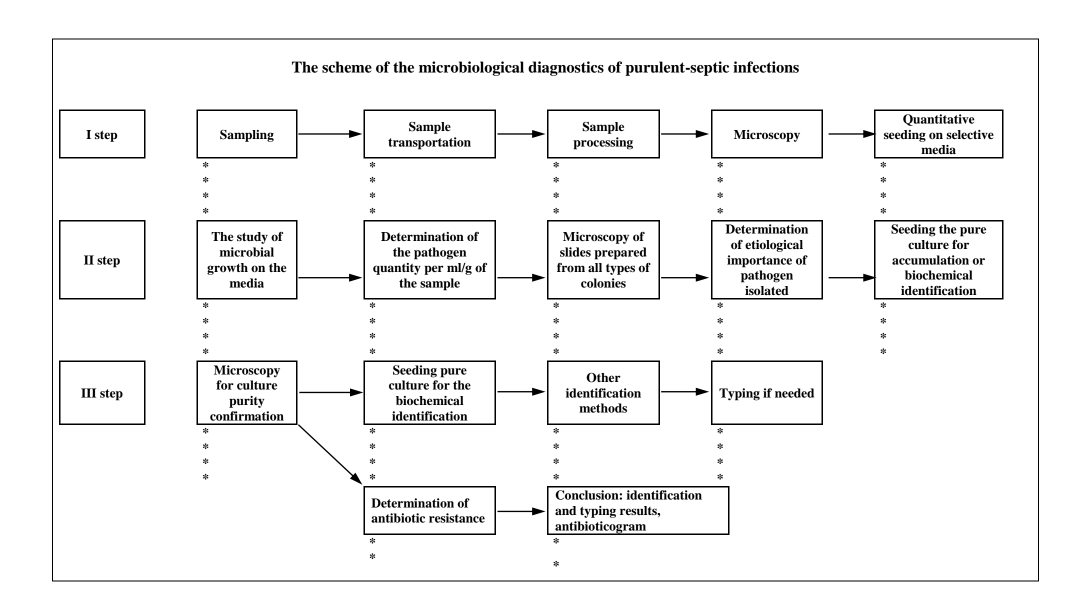
Levin media

Nutrient agar with lactose and bromothymol blue

Signature of the tutor_____

Additional materials for independent work for class N_2 10.

Criteria of etiological importance off opportunistic pathogens	Etiology (main pathogens) of purulent infection of the skin
1.	1.
2.	2.
4.	3.
5.	4.
6.	5.
$\frac{7}{8}$.	
9.	1
10.]



<u>Class No 11.</u> Clinical microbiology. Microbiological diagnostics of purulent infections of urinary tract. Hospital-acquired infection.

The list of questions to study:

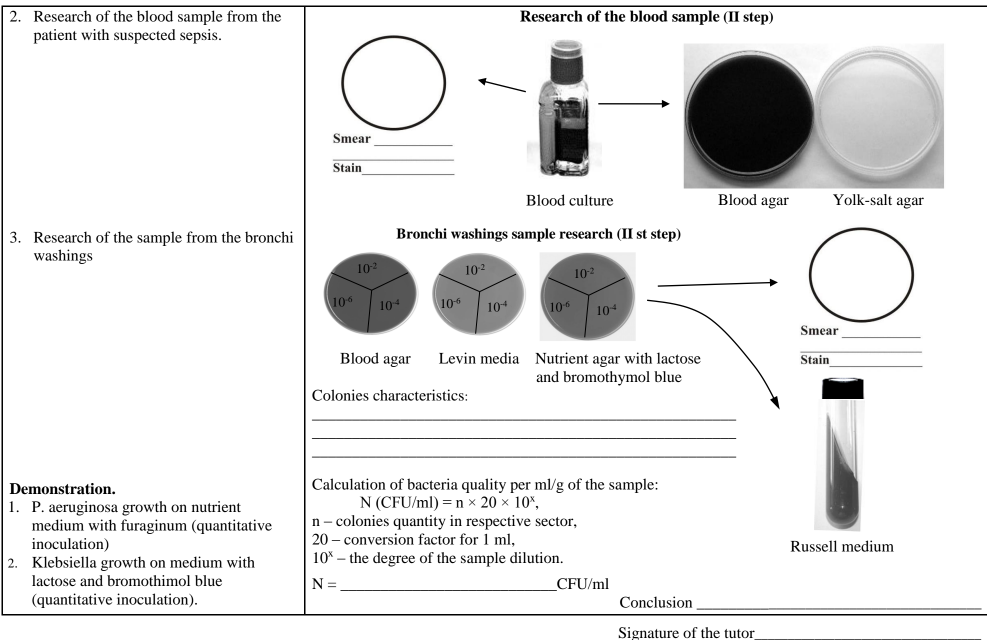
Clinical forms and etiology of septic-purulent (opportunistic) infections of the bronchi and lungs. Methods of microbiological diagnostics. Material for the research, rules and methods of sampling. Bacteriological method. Criteria for assessing the etiological role of isolated bacteria. Susceptibility to antibiotics.

Etiology and clinical forms of septic-purulent (opportunistic) infections of the urogenital tract. Methods of microbiological diagnostics. Material for the study, rules and methods of sampling. Urine culture. Criteria for assessing the etiological role of isolated microbes. Susceptibility to antibiotics. Antibioticogramm.

Nosocomial infections. Pathogens. Principles of microbiological diagnosis. Prevention.

Laboratory work

Tasks	Methods, results	
1. Independent work: "Research of the	Exsudate sample research (Hst step)	
sample from the burnt wound"	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	
	Blood agar Levin media Nutrient agar with p-Phenylenediamine	
	Colonies characteristics: (PPD)	
	Calculation of bacteria quality per ml/g of the sample: $N (CFU/ml) = n \times 20 \times 10^{x}$,	
	n – colonies quantity in respective sector, 20 – conversion factor for 1 ml, $\frac{\text{Smear}}{10^{\text{x}} - \text{the degree of the sample dilution.}}$	
	N =CFU/ml	
	Conclusion	



Additional materials for independent work for the class № 10 Etiology (main pathogens) of respiratory septic-purulent diseases

1.			
2.			
3.			
4.			
5.			

Etiology (main pathogens) of urogenital septic-purulent diseases

1.	٦
2.	
3.	
4.	
5.	

Hospital acquired infections (HAI, nosocomial infections) - any clinically recognizable infection contracted by patient due to residence or receiving various types of inpatient and outpatient medical care, the delivery of emergency medical services both in health care organizations and at home, as well as infectious disease contracted by medical staff as a result of professional activity, regardless of time of symptoms onset.

Nosocomial infections should be distinguished (introduced) from cases of infectious diseases registered in the delivery of health care in inpatient, outpatient medical institutions, or at home. Their main features are: the absence of a causal connection with the performance of therapeutic and diagnostic procedures and manipulations; acquisition of infection within the minimum incubation period before seeking medical help.

Etiology (main pathogens) of nosocomial infections

1.	
2.	
3.	
4.	
5	

CLASSIFICATION of HAI

HAI etiology includes bacteria; viruses; fungi; protozoa and metazoa.

By source of infection HAI can be exogenous; endogenous and auto-infection.

Depending on the profile of medical care nosocomial infections are divided into: surgical infection, obstetric infections; neonatal infections; other infections.

Depending on the entrance gate and localization of infection nosocomial infections are divided into: surgical wound infections; burn wound infection; infections of skin and soft tissue; primary bloodstream infections; sepsis; cardiovascular system infection; bone and joint infections; eye infection; ear infections; infection of the nose, throat, mouth and upper respiratory tract; lower respiratory tract infections; pneumonia; infections of the central nervous system; urinary tract infections; infections of the reproductive system; infections of the gastrointestinal tract.

Depending on the type of pathogen nosocomial infections are divided into: caused by obligate pathogens and opportunistic pathogens.

Depending on the spread in the organism HAI can be divided into: localized; generalized and systemic infections.

Depending on the course character nosocomial infections are divided into: acute; subacute and chronic.

By severity nosocomial infections are divided into: pathogen caring; mild; moderate and severe form.

Depending on the mechanisms, ways and factors of transmission of nosocomial infections are divided into: aerosol; contact (direct and indirect); parenteral; fecal-oral (food and water).

Class № 12. Microbiological diagnostics of fungal and protozoan imfections

Date	
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The list of questions to study:

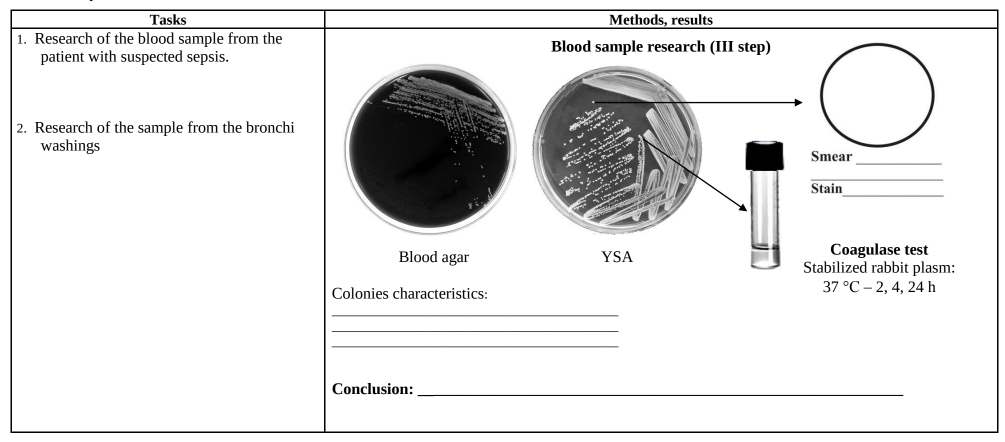
General characteristics and classification of protozoa. Pathogenic representatives. Laboratory diagnosis of malaria, toxoplasmosis, amebiasis, giardiasis, trichomoniasis.

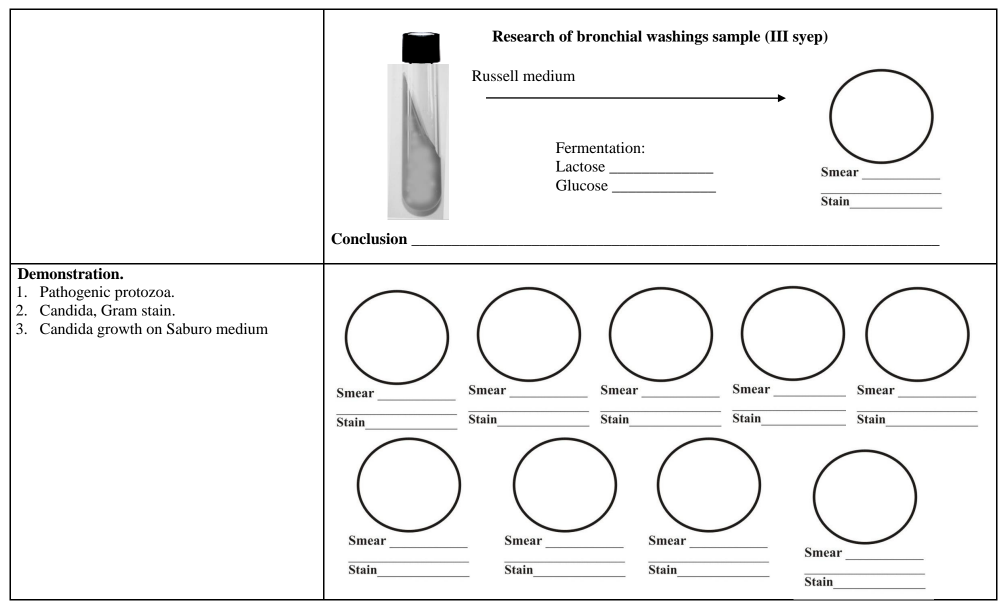
The causative agent of cryptosporidiosis.

Classification and general characteristics of fungi. Pathogens of ringworm, keratomycosis, deep mycoses. Candidiasis and conditions which promote its development. General principles of fungal infections diagnostics.

Pathogen of pneumocystosis.

Laboratory work





Signature of the tutor_____

DIAGNOSTICS OF MYCOSIS

Microscopic method. High diagnostic value of the method caused by significant differences in fungal morphology, simplicity and speed of the research. The result can be obtained within 1-2 hours. Microscopy can be conducted in native preparations without staining. For visualization of the pathogen in the biological material which is poorly transparent (hair, skin, nails, etc.) it should be processed with 10-20% alkaline (KOH), which dissolves keratin and has no effect on the morphology of the fungal cells. Fixed smears may be stained by Gram (fungi are Gram-positive), Romanovsky-Giemsa, special techniques. Dimorphic fungi in biological material are in the form of yeast. Microscopy of histological preparations is also possible.

Serological method:

Immunofluorescence is sensitive, specific and rapid method based on the identification of fungal Ag in biological materials.

PHAT, latex agglutination, PT, CFT, ELISA are used to detect fungal antigens and antibodies in blood, CSF, urine. Serological reactions not always highly specific, but produce results earlier than culture method.

Culture (mycological) method. Most pathogenic fungi are mesophiles (20-45 °C) and not demanding for the nutrient medium. Optimal pH ranges from 4.0 to 6.5. Growing time depends on the kind of fungus and can be from several weeks to 2-3 days. The most frequently used medium is Saburo agar (peptone agar with glucose or maltose). The acidity of the medium and high carbohydrate content inhibits the growth of bacteria. Dimorphic fungi (pathogens caused subcutaneous and deep mycoses) grow in the mycelial form at 20-25 °C. The identification of a pure culture is carried out by morphological and biochemical characteristics.

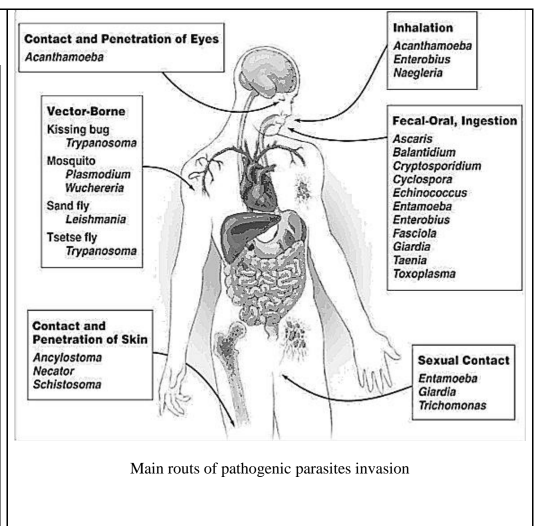
Allergic method. Skin tests are performed with fungal allergens (eg. Candide). Method is not very specific because of the group antigens presence.

Biological method. Bioassays in laboratory animals allow us to estimate the virulence of the pathogen, get in tissue culture of the fungus (usually in a form of yeast).

Molecular genetic methods. PCR and molecular hybridization are used. Among advantages - very high sensitivity and specificity, relative safety and short time needed for results.

The comparison of eu- and prokaryotic cell

Main characteristics	Prokaryotic cell	Eukaryotic cell
Cell size	Average 0,2-2,0 mkm	
Nucleus	Does not have a true nucleus. Nucleoid, is not separated from the cytoplasm by a membrane	
Chromosomes	Ring-like	
Number of chromosomes per cell	Usually one	
Mitochondria	No	
Endoplasmatic reticulum	No	
Ribosomes location	Dispersed in cytoplasm	
Sedimentation constant	70S	
Teichoic asides in cell wall	Gram positive bacteria	
Peptidoglycane in cell wall	All bacteria with exception of mycoplasm	
Endospores	Some has	
Division of cell	Binary (mitosis)	
Gametes, zygotes	No	

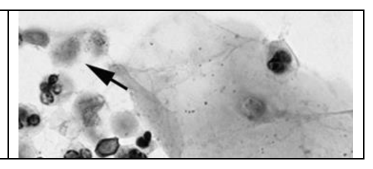


Protozoa belong to the domen – EUKARYA, kingdom – ANIMALIA, subkingdom – PROTOZOA, which includes 7 types, Four types of medical importance are showed in the tableu

Taxons	Representatives	Disease	Morphology
TYPE SARCOMASTIGOPHORA subtype Sarcodina	AMOEBAE Entamoeba histolytica	Amebiasis	
	Naegleria, acanthamoeba, hartmanella	Amoebic meningoencephalitis, keratitis	
subtype Mastigophora	LEISHMANIA Leishmania species	Leishmaniasis	
	TRYPANOSOMES Tripanosoma gambiense, Tripanosoma rodesiense Tripanosoma cruzi	African trypanosomiasis (sleeping disease) Chagas disease (American trypanosomiasis)	
	GIARDIA: Lamblia intestinalis (Giardia lamblia)	Diarrhea, malabsorption syndrome	

<u>Trichomonas</u> Trichomonas vaginalis

Trichomonas vaginalis vaginitis, urethritis, prostatitis



Medical-Enc.ru **PLASMODIUM MALARIA:** TYPE -Plasmodium vivax **APICOMPLEXA** Plasmodium ovale class – Sporozoa Plasmodium malariae Plasmodium falciparum

TOXOPLASMA: Toxoplasma gondii	Toxoplasmosis	
SARCOCYST: Sarcocystis species	Sarcocystosis	Sarcocystis 20um
ISOSPORA: Isospora species	Diarrhea	
CRYPTOSPORIDIUM: Cryptospodium species	Diarrhea	
CYCLOSPORA: Cyclospora cauetanensis	Diarrhea	o la
BABESIA: Babesia species	Babesiosis	

TYPE – CILIOPHORA class Kinetofragminophorea	BALANTIDIUM: Balantidium coli	Balantidiasis	
TYPE – MICROSPORA class Microsporea	MICROSPORIDIA: Encephalitozoon species Enterocytozoon species	Microsporidiasis	
	BLASTOCYST: Blastocystis hominis	Vacuolar	Cyst G Granular Amoeboid

MICROBIOLOGICAL DIAGNOSTICS OF PROTOZOAN INVASIONS

<u>AMEBIASIS</u> Microscopic method. Materials: samples of faeces or exudates from abscesses. Smears are stained with iodine solution or hematoxylin. Tissue forms with phagocytized erythrocytes or quad cysts. can be identified. In native specimens characteristic motile vegetative forms can be noted. IF may be used for the identification of pathogen

Serological method: PHA test, ELISA, CFT, and other tests may be used. The highest antibody titer can be detected in extraintestinal amebiasis.

Some non-pathogenic amoeba are morphologically identical to Entamoeba histolytica. The differentiation is based on the enzymatic, immunological and molecular genetic analysis.

TRYPANOSOMES

Microscopic method. Materials: samples of blood, punctate from cervical lymphatic nodes, cerebrospinal fluid. Smears are stained by Romanovsky-Giemsa method.

Cultural method. Trypanosomes can be cultured on a nutrient medium with blood as well as in white mice or rats.

Serological method. The determination of specific IgM by ELISA is used.

TRICHOMONIASIS

Microscopic method. Materials: samples from urethral discharge, prostatic secretions or urine sediment are studied. Smears are stained by Romanovsky-Giemsa (trophozoite nucleus is violet-ruby, cytoplasm - blue and blefaroplast, flagella and aksostil - pink-red), methylene blue. IF is also used.

Cultural method. In chronic trichomoniasis pathogen can be cultured on nutrient media with protein. The method gives good results when confirmation of convalescence is needed.

<u>TOXOPLASMOSIS</u> Microscopic method. Materials: biopsy, samples of body fluids (blood, cerebrospinal fluid, lymph node puncture, etc.). Smears are stained by Romanovsky-Giemsa method. Toxoplasma Ags may be detected by IF test.

Cultural method. Cultivation of Toxoplasma is possible in cell cultures and chicken embryo.

Serological method. Detection of specific IgM indicates the early stages of the disease. IgG peaks at 4-8 week of disease. ELISA is widely used.

Biological method. Mice are infected in the abdominal cavity or intracranially. They usually succumb 7-10 days after infection. The pathogen is identified microscopically or by serological method.

LEISHMANIASIS

Microscopic method. Materials: skin lesions (bumps, ulcers), bone marrow. Smears are stained by Romanovsky-Giemsa method. The detection of amastigote (nucleus and kinetoplasts are of red-purple color and cytoplasm is bluish) is of importance. IFT is also used.

Cultural method. Leishmania can be cultured on blood agar.

Biological method. Infection of mice or hamsters is possible.

Serological method. Specific antibodies may be detected by CFT, passive hemagglutination or ELISA.

Allergic method. Skin test with leishmania Ags may be used.

GIARDIASIS

Microscopic method. Materials: feces, duodenal secretion. In smears cysts or vegetative forms, can be detected. Iodine staining is usually used. IFT is also applicable.

Cultural method. Giardia can be cultured nutrient media.

Serological method. Specific antibody titers are higher in symptomatic giardiasis.

MALARIA

Microscopic method. Smears of blood are stained by Romanovsky-Giemsa method. Various forms of pathogen can be identified (red nucleus, blue cytoplasm).

Differentiation of species is carried out by morphological features of parasites and parasitized erythrocytes.

Serological method. Specific antibodies are detected by ELISA. IFT is applicable for diagnostics. **Molecular genetic method**. PCR.

BALANTIDIASIS

Microscopic method. Microscopy of smears from feces under low magnification allows to reveal large motile balantidiums.

Cultural method. Possible, but rarely used

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