P14 Revision for the practical examination

This practical session is not compulsory but students are highly recommended to attend (even another than their own group session, though should a problem with the hall capacity occur, "native" students will receive precedence).

Task: Orientation at survey of knowledge for the practical examination

Follow the presented survey and add your own notes according to the teacher's explanation and practical demonstration.

Attention! It is only an orientation at survey; at the practical examination you cannot raise objections that something "was not in the survey". The practical examination assesses the knowledge obtained during two terms of education, **not** the knowledge of a survey.

of education, not the knowledge of a survey.	
The basic requirements for each topic	Student's notes
Microscopy	
Gram staining:	
be able to perform it	
be able to observe a preparation and to	
identify G+/G- cocci/bacilli (+arrangement),	
yeasts, epithelial cells, WBCs	
know the principle	
Wet mount, other staining methods perfored in	
practicals (survey)	
(Ziehl-Neelsen staining, see Acid fast bacteria)	
Interpretation of microscopic findings (importance of	
epithelial cells, leucocytes)	
Culture	
Most important culture media	
be able to recognize blood agar, Endo agar	
and Mueller Hinton agar	
be able to describe the function of all the	
fourteen media from J03	
Inoculation (be able to inoculate a strain/a swab)	
Description of colonies (practically)	
Biochemical identification	
Catalase test	
be able to perform it	
• understand its principle	
be able to give an example of its use in	
diagnostics	
Strip tests * know the most important ones (oxidase.	
know the most important ones (oxidase, PYR, INAC) and to give examples of their	
use	
be able to use them practically (incl. reading	
the results)	
Hajna, MIU and other similar tests	
* know their practical use and what they detect	
Enterotest-like tests	
be able to read an Entero- or Staphy-test and	
o describe its principle	
Further notes:	

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methods and the way they are used (chloramin, NaOCl, Ca(OCl) ₂ , iodine-povidone, hydrogen peroxide, peracetic acid, ajatin, UV-rays disinfection, hot air and steam sterilization, radiation sterilization) To understand the methodological difference between testing the growth limit and the survival limit To be able to read corresponding tests (Task 1, P06) To know how effect of disinfection and sterilization can be tested Antimicrobial drugs To know principles of microdilution test, diffusion disk test and E-test, to be able to read the results of all of them and to interpret them To understant the importance of MIC and its comparison with breakpoint level To know basic methods of testing the factors of resistance (beta-lactamases) Serological tests (J07 to J09) To be able to read the results any of these tests; students will get the necessary information (dilution in the first well, c. o. counting in ELISA etc.) To be able to describe the basic indication for the test and to interpret these results in combination with other parameters; including ASO! The principle of antigen/analysis reactions and its use for antigen detection in a specimen/antigen analysis of a strain/antibody detection To understand the major interpretation difference between direct and indirect diagnostic methods	
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To know the principles of agglutination, precipitation	
agglutination on carriers, CFT, neutralisation (ASO,	
HIT, VNT), reactions with labelled components,	
western blotting, incl. differences between them	
To understand titers, titer dynamics, seroconversion,	
importance of IgM/IgG (and knowing what reactions	
enable their detection – importance of conjugate),	
avidity (A-aspiring students)	
To be able to construct the scheme of HBsAg and anti- HBs testing	
To understand the terms "heterophilic antibodies" and	
"anticomplementarity test"	
Detection of nucleic acid	
To know the basic indication for these methods in	
microbiology	
To understand the difference between methods	
with/without amplification	
To know the basic principle of the reaction, including	
two major ways of product detection	
To understand the importance of internal control	
To be able to read practically a PCR result (in a	
picture), including IC result interpretation	
Further notes:	
Virology To know the ways of isolating a virus (including	

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individ	dual structures of a fertilized egg)	
To be	able to differentiate a cell culture with/without	
CPE (in simplex cases only) and to understand, what a	
CPE is		
	serology: HIT, VNT, see serology)	
Para		
	ow basic methods for parasites (Faust, Kato,	
	m; thick and thin smear; C. A. T. swab and	
	a stained smear for trichomonads; indirect	
	ostics of tissue parasites)	
	able to distinguish the most common helminth	
	tapeworm, pinworm, common roundworm,	
	vorm) and tapeworm proglottid	
	ow the basic principles of sampling for	
parasi		
Easil	y culturable bacteria and yeasts (P01–I	P06; P10)
To be	able to find out (and utilize practically) a	
diagno	ostic algorithm to identify common bacteria	
except	G+ rods (Staphylococcus aureus, coagulase-	
negati	ve staphylococci, Streptococcus pyogenes, S.	
	ctiae, S. non-A-non-B, S. pneumoniae, oral	
	ococci, Enterococcus faecalis, E. faecium,	
Eschei	richia coli, Klebsiella pneumoniae, Salmonella	
enterio	ca, Proteus sp., Pseudomonas aeruginosa, other	
G– no	n-fermenters, Haemophilus influenzae, H.	
	fluenzae, Pasteurella multocida, Neisseria	
	rhoeae, Neisseria meningitidis, oral neisseriae,	
	cella catarrhalis, Candida albicans, Candida	
sp.)		
	+ rods: to know their main characteristics; to be	
	identify practically coryneform rods according	
	r palisade arrangement	
	erobic bacteria	
	able to describe an anaerobic jar and an	
	blic box, their parts and their function	
	ostridia: to know their main characteristics; to be	
	o identify <i>C. tetani</i> according to its sphaerical	
	al endospore	
	-fast rods	
	ow the principle of Ziehl-Neelsen staining, to be	
	o distinguish between the pictures of positive and	
	ve findings and pictures stained using other	
Stallill To lea	g methods	
	by the principles of acid-fast rod culture, to	
	basic media, to be able to distinguish pictures of	
-	ve findings/negative findings/pictures describing	
	hing else	
	al bacteria	
	plain the use (and complications in use) of direct	
	ds in spirochete diagnostics	
	derstand screening/confirmatory reactions for	
	lia and Treponema	
To be	able to read and interprete the tests (see also	
Serolo	egy)	
	er notes:	
Fung	i	
	ow basic diagnostic methods used in mycology	
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Topic P14

To be able to read a microprecipitation test for lung	
aspergillosis and to explain its principle	
To know the basic principles of sampling for	
mycology	
See also "Easily culturable bacteria and yeasts (P01–	
P06; P10)"	
Biofilm	
To know the diagnostic methods of biofilm detection	
To know the difference between three most typical	
methods of venous catheter microbiologic diagnostic	
To be able to read the results of the biofilm growth:	
glucose/time experiment (see P12 Task 4)	
To be able to read MBEC values and to interpret the	
result (in comparison with MIC)	I
Clinical microbiology	
To be able to find a pathogen in phagyngeal flora (and	
to know the composition of normal pharyngeal flora,	
and common pharyngeal pathogens)	
To be able to read a result of urine culture	
semiquantitatively and qualitatively	
For a simple mini-casuistry, be able to find out the	
best sampling method, including finding the best swab	
or container (practically)	
To understand basic principles of sampling under	
various circumstances	

Further notes:

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