Medical Oral Microbiology	Topic A – 04
Name:	Date:

Topic P04: Diagnostics of enterobacteria and bacterial agents of gastrointestinal infections

Table for major results of Task 1 to Task 5 (to be filled step by step):

ı abie	tor maj	jor rest	lits of i	ask 1 t	o rask	5 (to b	e fillea	step by	/ step):
Strain		K	L	M	N	P	Q	R	S
Gram stai	n – Task 1								
Culture	Size on								
(blood	BA								
agar and	Colour								
Endo	on BA								
agar)	Other								
Task 2	on BA								
	Size on								
	Endo								
	Colour								
	on Endo								
	Other								
	on Endo								
Hajna me	dium								
Task 3a									
Oxidase to	est								
Task 3b									
PARTIA									
CONCLU									
More	XLD								
media	agar								
Task 4a	MAL								
	agar								
	CIN								
	agar								
EnteroTes									
(Task 4b)									
	Antigen analysis								
(Tasks 5a	and 5b)								
FINAL									
CONCLU	JSION								

Task 1: Microscopy of suspicious strains

There are letter-labelled strains on the table. Gram-stain them and write your results to the table. Strain that is NOT G- rod should not be used in tasks 3 to 5. Draw the results of your observation.

Task 2: Cultivation on blood agar and Endo agar

Using standard procedure, describe colonies of all strains on blood agar and Endo agar. If the strain on the medium does not grow, cross the corresponding cell in the table. Bacteria, that do not grow on any of the media and morphologically look like curved gram-negative rods, might be *Campylobacter*. A G- rod, that does not grow on any of the media, but is not curved, will be studied in the next lesson. For comparison describe also the

Name: Date:

strain, that appeared morphologically as a gram-positive coccus. Stain just one strain and share the reset with other student around the table. Using the dermograph, mark the slides with corresponding letter.

Task 3: Group diagnostics of the most important gram-negative rods growing on Endo agar

a) Reading of an examination on oblique agar according to Hajna

Agar according to Hajna is a combined diagnostic medium. Nevertheless, in this task we will mostly search for biochemically non-active, neither glucose nor lactose splitting and sulphan non forming rods – the gram-negative non-fermenting bacteria ("non-fermenters"). All strains, growing on Endo, were inoculated on Hajna medium. Have a look to the result. Where the medium remained fully red, it is a biochemically non-active strain – very likely, a gram-negative non-fermenter. This strain will not be used in Task 4 and Task 5.

b) Oxidase test

The teacher will do as a demonstration oxidase test for all Gram-negative, on Endo agar growing bacteria. Oxidase-positive are members of family *Vibrionaceae* and some gram-negative non-fermenters; the *Enteobacteriaceae* are (with exception of *Plesiomonas*) oxidase negative.

Make partial conclusion after tasks 1 to 3. What bacteria are enterobacteria? Tasks 4 and 5 will be only performed with strains proven to be enterobacteria.

Task 4: Genus and species determination of enterobacteria using cultivation and biochemical tests

a) Cultivation of enterobacteria on other media

You have already seen, how the colonies look like on BA and Endo agar. Exam and describe the appearance of the enterobacterial colonies on CIN, XLD and MAL (fill the table).

b) Biochemical behaviour of enterobacteria

Evaluate given results of ENTEROtest 16, incubated a day before. Check, whether the results comply with others, already done tests; e.g. strains with sulphan formation lead to black colour of Hajna medium, *Yersinia* has tiny pink colonies, *Salmonella* pale transparent colonies with black centre on XLD and MAL medium... For the strain found to be *Salmonella*, write *Salmonella* sp. only. Count % of probability as a total of all % of probability of individual Salmonellas at the code: T index should be taken from the first *Salmonella*.

prob	ability of	marv	iduai	Samo	mena	s at u	e cou	e; 1 11	naex s	siloula	be ta	ken n	rom u	ie ms	t Saime	эпена.	
	ONPG	1H	1G	1F	1E	1D	1C	1B	1A	2H	2G	2F	2E	2D	2C	2B	2A
Strain:										·							
	1	2	4	1	2	4	1	2	4	1	2	4	1	2	4	1	2
	Code:						Iden	tificat	tion				% o	f prob).	T index	
	ONPG	1H	1G	1F	1E	1D	1C	1B	1A	2H	2G	2F	2E	2D	2C	2B	2A
Strain:																	
	1	2	4	1	2	4	1	2	4	1	2	4	1	2	4	1	2
	Code:						Iden	tificat	tion				% o	f prob).	T index	
	ONPG	1H	1G	1F	1E	1D	1C	1B	1A	2H	2G	2F	2E	2D	2C	2B	2A
Strain:	•																
	1	2	4	1	2	4	1	2	4	1	2	4	1	2	4	1	2

Name: Date:

Code:						Identification					% of prob.			T index		
ONPG	1H	1G	1F	1E	1D	1C	1B	1A	2H	2G	2F	2E	2D	2C	2B	2A
1	2	4	1	2	4	1	2	4	1	2	4	1	2	4	1	2
														•		
Code:					Identification					% of prob.			T index			
	ONPG 1	ONPG IH 1 2	ONPG 1H 1G 1 2 4	ONPG 1H 1G 1F 1 2 4 1	ONPG 1H 1G 1F 1E 1 2 4 1 2	ONPG	ONPG 1H 1G 1F 1E 1D 1C 1 2 4 1 2 4 1	ONPG 1H 1G 1F 1E 1D 1C 1B 1 2 4 1 2 4 1 2	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 1	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2H 2H 2H 4H 1H 2H 2H 4H 1H 2H 4H 4H 4H 4H 4H 4H 4	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2G 1 2 4 1 2 4 1 2 4 1 2	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2G 2F 1 2 4 1 2 4 1 2 4 1 2 4 1 2 4	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2G 2F 2E 1	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2G 2F 2E 2D 2D 2D 2D 2D 2D 2D	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2G 2F 2E 2D 2C 1 2 4 1 2 4 1 2 4 1 2 4 1 2 4 1 2 4	ONPG 1H 1G 1F 1E 1D 1C 1B 1A 2H 2G 2F 2E 2D 2C 2B 2D 1 2 4 4 1 2 4 4 1 2 4 4 1 2 4 4 1 2 4 4 4 4 4 4 4 4 4

Task No. 5 Antigen analysis in diagnostics of enterobacteriae – excluding of EPEC

In strain identified as *Escherichia coli*, perform antigen analysis using slide agglutination with two polyvalent sera. If both results will be negative, the strain does not belong into EPEC group.

Task No. 6: Susceptibility tests of enterobacteria to antibiotics

On your table, you will find diffusion disc tests for strains found to be *Enterobacteriaceae*. Write abbreviations of antibiotics according to the card and measure susceptibility zones for all tested strains. Borderline zones are written on the cards; using them, interpret the strains as susceptible (S) resistant (R) and dubious (D).

Strain →		•		•		· ,	, ,	
Antibiotic (full name)	Zone Ø (mm)	Interpr.						

Task No. 7 Diagnostics of Campylobacter

Observe the cultivation appearance that did not grow neither on BA nor on Endo agar and which, according to the morphology, is supposed to be a *Campylobacter* (because of being curved), on a special medium. Remember four main conditions for cultivation of *Campylobacter*: (a) special medium with charcoal and addition of antibiotics and antimycotics to prevent growth of other microbes, (b) microaerofilic conditions, (c) temperature elevated to 42 °C, what correspons to body temperature of birds – natural hosts, and (d) prolongation of the cultivation to 48 hours.

Describe the colonies write down the result of oxidase test (teacher will perform it as a demonstration). For *Campylobacter* a retarded positivity is typical, e. g. the strip becomes blue, but not immediately, but after a while.

Medical Oral Microbiology		Topic A – 04						
Name:		Date:						
Task No. 8: Urease te In diagnostics of helicobacters we u mucosa. Excised tissue is placed on negative yellow. Among two specim Result: Positive urease test was fou	the surface of media contains nens (X and Y) find the positive	directly with a bioptic spec urea and pH indicator. A we one.						
Task No. 9 Diagnostic <i>Vibrionaceae</i> is a bacterial family s culture <i>Vibrionaceae</i> . Mutual differ Enterotest 16 could be used, but a s how a <i>Vibrio</i> looks like microscopia	imilar to Enterobacteriaceae, rentiation is possible through b pecial codebook is required. A cally, and add some more prop	but oxidase-positive. We use in the properties of the state of the state of the state of the slide of the sli	nterobacteria. Even sed, too. Draw here,					
Microscopy:	Most important solid medium							
	Most important liquid medium	n for <i>Vibrio</i> :						
	The two most important sero	vars of <i>V. cholerae</i>						
	The two most important biov	ars of <i>V. cholerae</i> O1						
Check-up questions: 1. Do you know, what is the result of 2. For practical reasons, one medium	m used in diagnostics of entero		nite broth. What type					
of medium is it and what is its use?	(See practical J03)							
3. Do you know at least some antigonal antigonal antigonal and a some antigonal antigonal antigonal and a some antigonal an	enic types of EPEC?							
4. What pathogen is diagnosed by V	Vidal reaction? Is it a direct, o	r indirect method? Which	type of reaction is it?					
5. Is it recommended to use antibio	tic treatment for intestinal infe	ction? Why?						
6. In which clinical material it is mo	ore likelly to find Salmonella T	Typhi in typhoid fever rath	er than in the stool?					