

# ANTIMICROBIAL AND CLINICAL MICROBIOLOGY GUIDEBOOK

**Ninth Edition** 

January 2023



#### **TABLE OF CONTENTS**

Introduction	Page #
Contacts	3
Clinical Microbiology	4
Organism Identification Flowcharts	
Gram Positive Organisms	4
Gram Negative Organisms	5
Key phrases in microbiology	6
Bacteriology	
Susceptibility Testing	6
Intrinsic Resistance Tables for Organisms	16
Anaerobe Antibiogram data, CLSI	19
Interpretation of Microbiology Reports	21
Specimen Requirements	24
<u>Timing of Reports</u>	24
<u>Urinalysis and Urine Culture</u>	24
Stool Testing/C. difficile	25
Blood Cultures	26
Respiratory Cultures	29
Genital tract Cultures	30
Mycobacteriology	31
Virology	33
Mycology	34
Parasitology	34
Antimicrobial Formulary/Cost Table	36
Renal Dosage Adjustment Guidelines for Antimicrobials	45
Reportable diseases	58-60

#### ANTIMICROBIAL AND CLINICAL MICROBIOLOGY GUIDEBOOK

#### 2023

#### **INTRODUCTION**

This is the Ninth Edition of the Antimicrobial and Clinical Microbiology Guidebook at LMH Health. The development of this guidebook has been a joint effort of the Antimicrobial Stewardship Program, Infectious Disease and Prevention, Pulmonology Specialist Group, Pharmacy, and the Microbiology Department. The purpose of the booklet is to optimize antimicrobial usage and patient outcomes for infectious disease-related issues.

Every effort has been made to ensure that the information is complete, accurate, and up to date; however, this booklet does not serve as a substitute for clinical judgment or consultation with experts in Infectious Diseases. Application of this information to each clinical situation is the responsibility of the practitioner.

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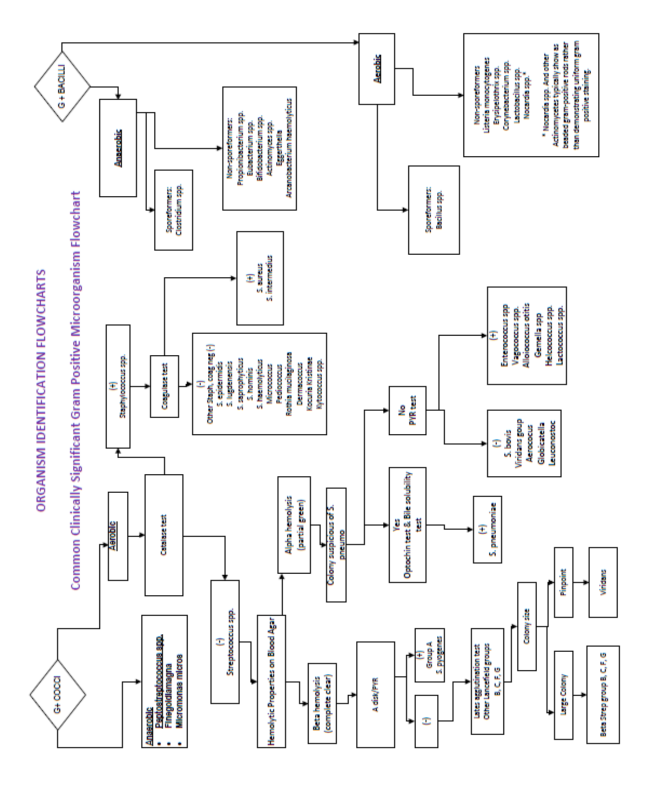
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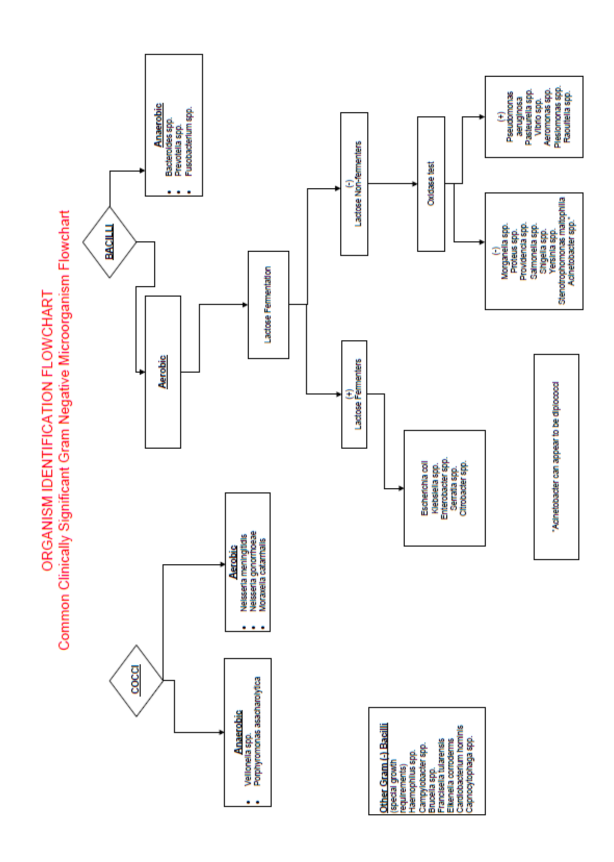
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#### **Key phrases from the Microbiology Laboratory**

Phrase	Suggested findings
Gram positive cocci in clusters	Staphylococcus species
Gram positive cocci in pairs and chains	Streptococcus species or Enteroccoccus species
Gram positive cocci in pairs	Streptococcus pneumoniae
Small pleomorphic Gram negative coccobacilli	Haemophilus species
Pleomorphic Gram positive bacilli	Corynebacterium species or Cutibacterium (prev.
	Propionobacterium) species
Branching Gram positive bacilli	Actinomyces or Nocardia species
Budding yeast or pseudohyphae	yeast
Fungal elements or hyphal elements	mold
Gram negative fusiform bacilli	Fusobacterium species or Capnocytophaga species

#### **BACTERIOLOGY**

#### **Susceptibility Testing**

Susceptibility testing is an *in vitro* assay that allows us to detect resistance to antimicrobial agents that may be used to treat an infection. It is important to note however, that clinical outcome may be dependent on various patient specific factors such as immune status or surgical treatment that are not reflected in laboratory tests.

All methods of susceptibility testing are based on diffusion or dilution.

#### A. Semi-Automated Susceptibility Testing

Semi-automated antimicrobial susceptibility testing is performed using the Vitek 2 system which is based on broth microdilution. This system allows the laboratory to rapidly perform identification and susceptibility testing on most common pathogens (e.g. Staphylococci, Enterobacteriaceae, Enterococci, *Pseudomonas* species, etc...). The antibiotics tested vary based upon the Vitek panel used and the antibiotics that are currently on the LMH formulary. The Vitek panels are chosen and agreed upon by the Antimicrobial Stewardship Committee, Pharmacy, Infectious Disease Physicians, and the microbiology laboratory.

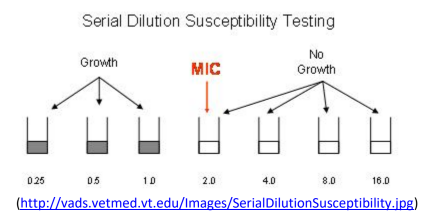
The microbiology laboratory reports antibiotics from most antibiotic classes that are appropriate for the specific organism tested. For example, if the laboratory recovers an isolate from urine, the results from the following antibiotic classes are reported: penicillin, Beta lactam/Beta lactamase combination, cephems, carbapenems, aminoglycosides, fluoroquinolones, nitrofurantoin, and trimethoprim-sulfamethoxazole.

Selective reporting of antibiotics for each organism group is reviewed and approved by the Antimicrobial Stewardship Committee and the medical director of the laboratory annually.

The results obtained from the Vitek system are based on the minimum inhibitory concentration (MIC). The MIC is defined as the lowest concentration of antibiotic that completely inhibits growth of the specific organism being tested.

For example, in figure 1 below, the organism being tested grew in wells containing 0.25, 0.5, and 1.0 ug/ml of antibiotic. The lowest concentration of antibiotic (MIC) that completely inhibits growth was 2.0 ug/ml.

Figure 1:



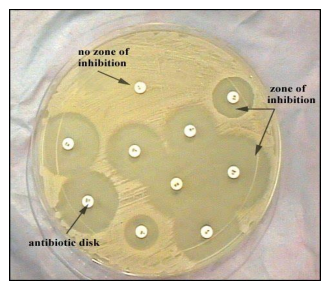
The MIC is then interpreted (S=susceptibile, I=Intermediate, or R=resistant) using FDA approved guidelines. These guidelines are based on many studies, including clinical, pharmacokinetic/pharmacodynamic, and microbiological studies.

If susceptibility testing for a specific organism cannot be reported using the automated Vitek method, disc diffusion testing may be performed and results will be reported based on CLSI (Clinical and Laboratory Standards Institute) standards if available.

If the laboratory cannot perform testing in-house, the physician may notify the microbiology laboratory (505-6177) and request the isolate be submitted to a reference laboratory for further testing.

#### B. Disk Diffusion

The LMH microbiology laboratory does routinely perform disk diffusion (Kirby Bauer) antimicrobial susceptibility testing for *Pseudomonas aeruginosa* isolates from cystic fibrosis patients, beta-lactamase positive *Haemophilus* species, and other fastidious organisms. Disk diffusion allows for measurement of the zone of growth inhibition. See figure 2. Figure 2:



(http://mrsa30day.com/wp-content/uploads/2012/08/Antibiotic-sensitivity.jpg)

The CLSI provides interpretive standards for reporting an organism as S, I, or R based on the zone of inhibition. The main difference between disk diffusion testing and MIC testing is that disk diffusion provides clinicians with qualitative results, whereas MIC testing provides the clinicians quantitative results.

Knowing the MIC can help clinicians incorporate pharmacodynamic/pharmacokinetic principles into the design of the treatment regimen.

The size of the zone of inhibition does not give us the MIC, and so in this specific case disk diffusion is not helpful. Also, zone sizes cannot be compared between drugs. Just because drug A has a larger zone size than drug B, it does not mean that drug A will work better. Zone sizes must be correlated back to the CLSI interpretive standards in order to determine susceptibility or resistance.

#### C. E-test

The LMH laboratory does perform E-test susceptibility testing for daptomycin on all MRSA and VRE isolates from non-respiratory tract sites per the Infectious Disease practitioner. Daptomycin susceptibility testing is available by E-test upon special request by ID or Pharmacy. Penicillin susceptibility testing is also available by E-test for unusual Strep species.

The E-test is an agar based method that uses a plastic strip with antibiotic concentrations in variable size plastic disks on its underside. When placed on an agar surface pre-inoculated with the bacterial isolate, the diffusing antibiotic creates a concentration gradient in the agar. Decreasing concentrations of antibiotic from the top of the strip to the bottom create

an ellipsoid diffusion pattern around the strip. The resultant elliptical zone of inhibition allows the MIC to be read at the point where the zone crosses the E-test strip (figure 3).

Figure 3:



### D. Special Susceptibility Testing Issues:

#### **Extended-spectrum Beta-lactamases (ESBL's)**

ESBL's are Beta-lactamases that are capable of hydrolyzing expanded-spectrum cephalosporins (ceftriaxone, cefotaxime, and ceftazidime) as well as cefepime and aztreonam. ESBL's can be isolated from many different Enterobacteriaceae species but are most commonly isolated from *Klebsiella pneumoniae*, *Klebsiella oxytoca*, *E. coli*, and *Proteus mirabilis*.

Using the Vitek, isolates that carry ESBL's can initially be intermediate or resistant to one or all of the extended spectrum cephalosporins, cefepime or aztreonam. This is due to the fact that there are many different ESBL's with different substrate specificities. The Vitek gram negative panel includes an ESBL confirmatory test and all ESBL positive isolates are verified. If a particular isolate is confirmed as resistant or intermediate to any of the extended-spectrum cephalosporins, cefepime or aztreonam, the following statement will be included in the report: "Isolate is an ESBL producing strain resistant to all penicillins, cephalosporins, and aztreonam."

ESBL positive *E coli* isolates from urinary tract specimens will be tested for fosfomycin and amoxicillin/clavulanate susceptibility per the Antimicrobial Stewardship Committee.

The GenMark Gram Negative Blood culture assay can detect the CTX-M beta-lactamase resistance mechanism. CTX-M is a beta-lactamase that has potent activity against cefotaxime. The mechanism of resistance is plasmid-mediated.

Because of the significant public health implications, the spread of CTX-M beta-lactamase producers' merits close monitoring.

#### **Carbapenemases:**

Carbapenemases are beta-lactamases that are capable of hydrolyzing all beta-lactams, including the carbapenems. Carbapenemases can be isolated from many different Enterobacteriaceae species.

Carbapenemases are particularly dangerous resistance mechanisms, since they can inactivate a wide range of different antibiotics. Enterobacteriaceae, Pseudomonas, and Acinetobacter isolates that produce carbapenemase have been referred to as "Infection Control Emergencies." Bacteria that produce carbapenemases (carbapenemase-producing-Carbapenem resistant organisms, ie. CP-CRO) are often referred to in the news as "superbugs" because infections caused by them are extremely difficult to treat.

For all acute care facilities, CDC and HICPAC recommend an aggressive infection control strategy, including managing all patients with carbapenem resistant organisms (CRO) and carbapenemase producing-carbapenem resistant organisms (CP-CRO) using contact precautions.

There are several important mechanisms of carbapenemase producing carbapenem resistance that increases the risk for dissemination:

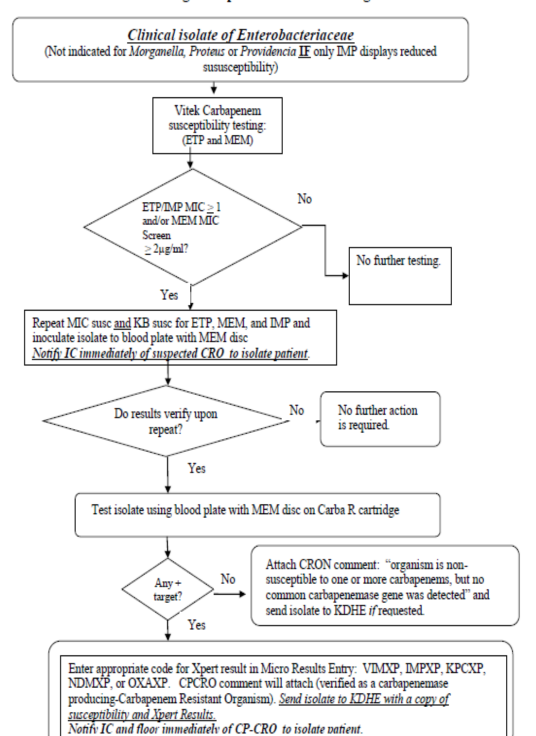
- KPC: Klebsiella pneumoniae carbapenemase (KPC) refers to the production of a
  carbapenemase enzyme, bla<sub>kpc</sub>. The gene that encodes the bla<sub>kpc</sub> enzyme is carried on a
  mobile piece of genetic material (transposon), which can easily be passed from one
  organism to the other.
- 2. <u>NDM</u>: New Dehli Metallo-beta-lactamase-1 refers to the production of a carbapenemase referred to as NDM-1 enzyme. The most common bacteria that produce this enzyme are gram negatives such as *E coli* and *Klebsiella pneumoniae*, but the gene for NDM *bla*<sub>NDM-1</sub> can spread from one strain of bacteria to another by horizontal gene transfer.
- 3. <u>IMP</u>: IMP-type carbapenemases are plasmid mediated Class B metallo-beta-lactamases that can be found in both enteric Gram-negative organisms and in *Pseudomonas* and *Acinetobacter* species.
- 4. <u>VIM</u> (Verona integron-encoded metallo-beta-lactamase): The VIM family of carbapenemases occur mostly in *Pseudomonas aeruginosa* and *P. putica* and vary rarely in Enterobacteriaceae. The VIM enzymes are integron-associated and hydrolyse all beta-lactams and can evade all beta-lactam inhibitors.
- 5. <u>OXA</u> (Oxacillinase): The OXA group of beta-lactamases occur mainly in Acinetobacter species.

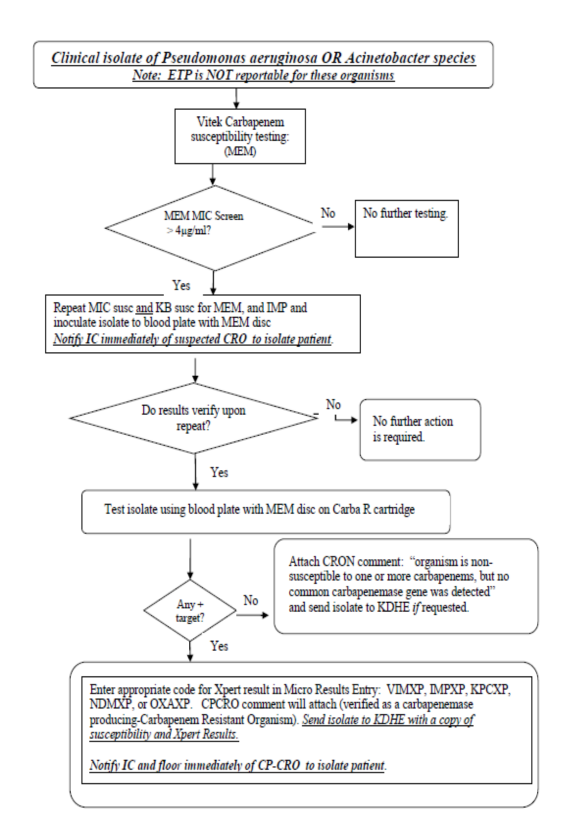
The GenMark Gram Negative Blood Culture assay can detect the following carbapenem resistance mechanisms in positive blood cultures: KPC, NDM, VIM, OXA, and IMP. Infectious disease consultation is suggested when one of these resistance mechanisms is detected.

If the Vitek susceptibility profile suggests the presence of a carbapenemase producing organism, the microbiology department will repeat testing and will also perform Kirby bauer susceptibility testing to verify. In addition, the lab will perform a Carba-R test on the Gene Xpert on the isolate. The Carba-R cartridge can detect KPC, NDM, VIM, OXA, and IMP. The following flowcharts were developed for testing based on CDC recommendations.

It is important to note that if an isolate is suspected to be a carbapenemase producing carbapenem resistant organisms (CP-CRO), the Infectious Disease specialist and the Infection Preventionist will be notified immediately to ensure the patient is placed in isolation. Carbapenemase screening can be performed if ordered by Infection Prevention. A rectal swab (dual swab culturette with Liquid Stuart's media) should be collected for screening.

#### Practical Testing Schemes to Carbapenemase Producing-Carbapenem Resistant Organism (CP-CRO) vs. Carbapenem Resistant Organism (CRO) Using the Xpert Carba-R Cartridge





#### Inducible clindamycin-resistance in *Staphylococcus* and *Streptococcus* species:

Erythromycin resistance within staphylococci is typically mediated through two distinct mechanisms. The first mechanism entails protection of the ribosome from erythromycin and clindamycin through methylation (referred to as MLS<sub>B</sub> resistance). This mechanism may be constitutive (conferring resistance to both erythromycin and clindamycin) or inducible (conferring resistance only to erythromycin). Published clinical reports have demonstrated that *S. aureus* isolates carrying an inducible MLS<sub>B</sub> resistance gene should be considered resistant to clindamycin even if the *in vitro* result considers the isolate susceptible to clindamycin. The second resistance mechanism is conferred through efflux of erythromycin out of the cell through specific pumps (encoded by the msrA gene). Staphylococcal isolates carrying the MsrA efflux pump are resistant only to erythromycin and not clindamycin.

The Vitek performs an inducible clindamycin resistance (ICR) test (otherwise known as D-test) automatically on *Staph* species, group A Beta *Strep* and group B Beta *Strep*. If the test is positive for ICR, the clindamycin result is not reported and a comment is attached to the report "isolate presumed resistant to clindamycin by ICR testing."

Isolates of *Streptococcus pneumoniae*, and other Beta *Streptococcus* species (as specified by CLSI) may require a manual D-test performed (Figure 4). If the D-test is positive the clindamycin result is not reported and a comment is attached to the report "isolate presumed resistant to clindamycin by ICR testing."

Figure 4 (positive D-test by manual method):



#### **Inducible Methicillin Resistance in Staph aureus:**

Methicillin resistance in S. aureus is generally due to the presence of the mecA gene located on the staphylococcal cassette chromosome mec(SCCmec). The mecA gene codes for production of an altered penicillin binding protein PBP2', also referred to as PBP2a. PBP2' has a low binding affinity for beta-lactam antibiotics. Because oxacillin and other betalactams cannot bind to the altered PBP2' site, these antibiotics are ineffective. The 'gold standard' for the detection of MRSA is molecular methodology using either polymerase chain reactions (PCR) or nucleic acid amplification. Molecular methods for detection of genes such as mecA that code for resistance are being used in laboratories with increasing frequency. However, routine susceptibility testing by microdilution or disk diffusion is still the method of choice for resistance determination in most laboratories. Until recently, our laboratory used microdilution and PBP2' latex methods to detect MRSA. In January 2012, the Cepheid SSTI MRSA/MSSA cartridge for use on the GeneXpert platform was added to the laboratory's test menu. In addition to a routine wound culture, we began to offer a special site culture with PCR for MRSA/MSSA. PCR results are available in just over one hour from the time of specimen receipt in the laboratory. PCR testing performed on these specimens is followed up with routine culture and susceptibility testing.

Our laboratory has previously identified *S aureus* isolates from outpatient wound specimens that were positive for the *mec*A gene by PCR that were found to be sensitive to oxacillin by follow-up routine susceptibility testing. Upon further investigation, these isolates were found to demonstrate resistance in vitro, after exposure to a beta-lactam (cefoxitin) antibiotic. In other words, the *S aureus* isolates demonstrated "inducible methicillin resistance."

The prevalence of this inducible methicillin resistant *S aureus* strain (Ridom spa type T175) has been found to be low however susceptibility testing alone can miss these strains. It is important to be aware that this strain has historically been found in our community. Also, it is important to note that if an inducible MRSA isolate were isolated from a serious infection, such as sepsis, and the infection were treated with first-generation cephalosporins or semi-synthetic penicillin, the patient may fail therapy.

#### **INTRINSIC RESISTANCE TABLES**

Intrinsic resistance is the innate ability of a bacterial species to resist activity of a particular antimicrobial agent through its inherent structural or functional characteristics. Such natural resistance can be due to: lack of affinity of the drug for the bacterial target, inaccessibility of the drug into the bacterial cell, extrusion of the drug by chromosomally encoded active exporters, or innate production of enzymes that inactivate the drug.

Below are lists of organisms with their respective intrinsic antimicrobial resistance.

Enterococcus species				Antimi	crobial	agents			
Organisms	Cephalosporins	Vancomycin	Teicoplanin	Aminoglycosides	Clindamycin	Quinupristin-dalfopristin	Trimethoprim	Trimethoprim/Sulfameth oxazole	Fusidic Acid
Enterococcus faecalis	R*			R*	R*	R	R	R*	R
Enterococcus faecium	R*			R*	R*		R	R*	R
Enterococcus gallinarum/ Enterococcus casseliflavus	R*	R		R*	R*	R	R	R*	R

<sup>\*</sup>Warning: For *Enterococcus* spp., cephalosporins, aminoglycosides (except for high level resistance screening), clindamycin, and trimethoprim-sulfamethoxazole may appear active in vitro, but are not effective clinically.

Note: Gram-positive bacteria are also intrinsically resistant to aztreonam, polymyxin B/colistin and naladixic acid.

Staphylococci		Antimicrobial Agents									
Organisms	Novobiocin	Fosfomycin Fusidic Acid									
S. aureus	There is no	There is no intrinsic resistance in these species.									
S. lugdenensis											
S. epidermidis											
S. haemolyticus											
S. saprophyticus	R	R	R								
S. capitis		R									
S. cohnii	R										
S. xylosus	R										

#### Notes:

- 1. Gram positive bacteria are also intrinsically resistant to aztreonam, polymyxin B/colistin and naladixic acid.
- 2. MRSA and oxacillin resistant coagulase negative staphylococci are considered resistant to other beta-lactam agents, (penicillins, beta-lactam/beta-lactamase inhibitor combinations, cephems with the exception of ceftaroline), and carbapenems.

Enterobacterales					A	ntimic	robial /	Agent				
Organisms	Ampicillin	Amoxicillin-clavulanic acid	Ampicillin-sulbactam		Ticarcillin	Cephalosporin I: Cefazolin, Cephalothin	Cephamycins: Cefoxitin, Cefotetan	Cephalosporin II: Cefuroxime	Imipenem	Tetracyclines and Tigecycline	Nitrofurantoin	Polymyxin B/Colistin
Citrobacter freundii	R	R	R			R	R	R				
Citrobacter koseri	R				R							
Klebsiella aerogenes	R	R	R			R	R	R				
(prev.Enterobacter)												
Enterobacter cloacae	R	R	R			R	R	R				
complex												
Escherichia coli		TI	nere is r	no i		resistan	ce to beta	-lactams	in th	is orgai	nism	
Escherichia hermannii	R				R							
Hafnia alvei	R	R	R			R	R					
Klebsiella pneumoniae	R				R							
Morganella morganii	R	R				R		R	*	R	R	R
Proteus mirabilis	No	intrinsi	c resista	nce	to penio	illins and	cephalospo	rins	*	R	R	R
Proteus penneri	R					R		R	*	R	R	R
Proteus vulgaris	R					R		R	*	R	R	R
Providencia rettgeri	R	R				R			*	R	R	R
Providencia stuartii	R	R				R				R	R	R
Salmonella and Shigella spp	1 <sup>ST</sup> AN	D 2 <sup>ND</sup> ge					beta-lactar appear acti				effective	clinically.
Serratia marcescens	R	R	R			R	R	R			R	R
Yersinia enterocolitica	R	R			R	R						

Note: Cephalosporins III, cefepime, aztreonam, ticarcillin-clavulanate, piperacillin-tazobactam, and the carbapenems are not listed, because there is no intrinsic resistance in Enterobacteiaceae.

Enterobacteriaceae are also intrinsically resistant to clindamycin, daptomycin, glycopeptides, (vancomycin), linezolid, macrolides (erythromycin, clarithromycin, azithromycin), quinupristin-dalfopristin, and rifampin.

Non- Enteroba cteriacea e		Antimicrobial Agents																				
Organism s	Piperacillin	Ticarcillin	Ampicillin-Sublactam	Amoxicillin-clavulanic	Piperacillin-tazobactam	Ampicillin, Amoxicillin	Cefotaxime	Ceftriaxone	Ceftazidime	Cefepime	Aztreonam	Imipenem	Meropenem	Ertapemem	Polymyxin B/Colistin	Aminoglycosides	Tetracycline	Tigecycline	Trimethoprim	Trimethoprim/Sulfa	Chloramphenicol	Fosfomycin
Acinetobac ter			*	R		R					R			R					R		R	R
baumannii																						
/A.																						
calcoacetic us complex																						
Burkholderi	R	R	R	R	R	R	R	R		R	R	R		R	R	R			R			R
a cepacia complex																						
Pseudomon			R	R		R	R	R						R			R	R	R	R	R	R
as aeruginosa																						
Steno.	R	R	R	R	R	R	R	R			R	R	R	R		R	#	#	R			R
maltophilia	' '	.,		.,	'`	.,	,	.,			•		٠,	٠,		٠,	"	,	.,			.,

- 1. \*Acinetobacter baumannii/calcoaceticus may appear to be susceptible to ampicillin-sulbactam due to the activity of sulbactam with this species.
- 2. # Stenotrophomonas maltophilia is intrinsically resistant to tetracycline but not to doxycycline or minocycline.
- 3. Note: Nonfermentative gram-negtive bacteria are also intrinsically resistant to cephalosporin I (cephalothin, cefazolin), cephalosporin II (cefuroxime), cephamycins (cefoxitin, cefotetan), clindamycin, daptomycin, fusidic acid, glycopeptides (vancomycin, teicoplanin), linezolid, macrolides (erythromycin, azithromycin, clarithromycin), penicillin, quinupristin-dalfopristin, and rifampin.

#### Other organisms:

Organisms	Natural Resistance Against
Anaerobic bacteria	Aminoglycosides
Aerobic bacteria	Metronidazole
Lactobacilli and Leuconostoc	Vancomycin
Aerococcus urinae	Sulfonamides and Netilmicin
Cardiobacterium hominis	Clindamycin

Anaerobic Organisms	Number of Strains	Amnicillin-	sulbactam	Number of Strains	Pineracillin.	tazobactam	Number of Strains	Cefoxitin		Number of Strains		Ertapenem	Number of Strains		Imipenem	Number of Strains		Meropenem
Percent susceptible (%S) and percent resistant (%R) <sup>b</sup>		%S	%R		%S	%R		%S	%R		%S	%R		%S	%R		%S	%R
Breakpoints, μg/mL		≤ 8/4	≥ 32/16		≤ 16/4	≥ 128/4		≤ 16	≥ 64		≤ 4	≥ 16		≤ 4	≥ 16		≤ 4	≥ 16
B. fragilis	129	84	2	1030	96	1	830	100	0	133	82	14	189	97	1	1505	93	5
B. thetaiotaomicron	76	82	5	252	87	0	258	13	54	-	-	-	70	100	0	328	99	0
B. ovatus	30	80	3	206	94	0	177	20	34	19 <sup>c</sup>	84 <sup>c</sup>	16 <sup>c</sup>	49	100	0	236	95	1
B. vulgatus	20 <sup>c</sup>	45 <sup>c</sup>	15 <sup>c</sup>	168	92	0	153	73	14	-	-	-	35	97	0	171	96	4
B. uniformis	19 <sup>c</sup>	84 <sup>c</sup>	O <sub>c</sub>	78	96	0	72	85	10	-	-	-	19 <sup>c</sup>	100 <sup>c</sup>	Oc	93	100	0
Parabacteroides distasonis	27 <sup>c</sup>	59 <sup>c</sup>	19 <sup>c</sup>	92	95	1	82	29	43	-	-	-	26 <sup>c</sup>	100°	0	119	97	2
Anaerobic Organisms			Number of Strains		Clindamycin			Number of Strains			Moxifloxacin			Number of Strains		Metronidazole		
Percent susceptibl (%S) and percent resistant (%R) <sup>b</sup>	le			%5	;	%R				%S	%	SR.				%S	9	6R
Breakpoints, μg/m	L			≤ 2	2	≥ 8				≤ 2	≥	8				≤ 8	≥	32
B. fragilis		1	013	26	-	22	25	6		61	3	32	1	140		100		0
B. thetaiotaomicro	n	3	328	28		49	70	)		54	3	6	,	322		100		0
B. ovatus		2	207	46		51	59	)		41	2	25	:	236		100		0
B. vulgatus		1	171	53	53 46		29	с		31 <sup>c</sup>	4	5 <sup>c</sup>		186		100		0
B. uniformis			87	45	45 48		25	с		48 <sup>c</sup>	c 40°		89			100		0
Parabacteroides distasonis		1	108	43		44	37	,		62	3	5		118		100		0

#### Footnotes

a. Data were generated from unique isolates from patient specimens submitted to Tufts Medical Center, Boston, Massachusetts; International Health Management Associates, Inc., Schaumburg, Illinois; R.M. Alden Research Laboratory, Culver City, California; Creighton University School of Medicine, Omaha, Nebraska; Mayo Clinic College of Medicine and Science, Rochester, Minnesota; and the Centers for Disease Control and Prevention, Atlanta, Georgia. All testing was performed by the agar dilution method. Information and analysis of previous versions of this table have been published.

- b. Intermediate category is not shown but can be derived by subtraction of %S and %R for each antimicrobial agent from %100.
- c. Calculated from fewer than the CLSI document  ${
  m M39}^{1}$  recommendation of 30 isolates.

Anaerobic Organisms	Number of Strains	Ampicillin-	sulbactam	Number of Strains	Strains Piperacillin- tazobactam		Number of Strains	Imipenem		Number of Strains	Strains		Number of Strains	o illinia	remomina
Percent susceptible (%S) and percent resistant (%R) <sup>b</sup>		%S	%R		%S	%R		%S	%R		%S	%R		%S	%R
Breakpoints, μg/mL		≤ 8/4	≥ 32/16		≤ 32/4	≥ 128/4		≤ 4	≥ 16		≤ 4	≥ 16		≤ 0.5	≥ 2
Prevotella spp.	29 <sup>c</sup>	97 <sup>c</sup>	3c	63	100	0	29 <sup>c</sup>	100	0	92	98	0	63	100	0
Fusobacterium spp.	20 <sup>c</sup>	100 <sup>c</sup>	O <sub>c</sub>	55	96	2	75	95	4	20 <sup>c</sup>	100 <sup>c</sup>	0°	_d	_d	_d
Anaerobic gram- positive cocci <sup>e</sup>	_d	_d	_d	1853	99	1	134	99	0	1647	100	0	1647	100	0
Cutibacterium (formerly Propionibacterium) acnes <sup>f</sup>	_d	_d	_d	18 <sup>c</sup>	100 <sup>c</sup>	0°	17 <sup>c</sup>	94 <sup>c</sup>	O <sup>d</sup>	_d	_d	_d	_d	_d	_d
Clostridium perfringens	15 <sup>c</sup>	100 <sup>c</sup>	0	410	100	0	23 <sup>c</sup>	100 <sup>c</sup>	0°	417	100	0	402	90	4
Clostridioides (formerly Clostridium) difficile <sup>8</sup>	76	99	0	542	93	0	480	69	4	609	99	0	533	6	37
Other <i>Clostridium</i> spp.	_d	_d	_d	439	94	1	71	99	0	390	100	0	390	69	13

Anaerobic Organisms	Number of Strains		Clindamycin	Number of Strains	Moxifloxacin		Number of Strains	Metronidazole		
Percent susceptible (%S) and percent resistant (%R) <sup>b</sup>		%S	%R		%S	%R		%S	%R	
Breakpoints in µg/mL		≤ 2	≥ 8		≤ 2	≥ 8		≤ 8	≥ 32	
Prevotella spp.	29 <sup>c</sup>	69 <sup>c</sup>	28 <sup>c</sup>	92	66	25	92	99	0	
Fusobacterium spp.	75	77	21	75	68	23	75	95	5	
Anaerobic gram- positive cocci <sup>e</sup>	1826	97	3	300	72	21	1692	100	0	
C. (formerly <i>P.)</i> acnes <sup>f</sup>	17 <sup>c</sup>	53 <sup>c</sup>	35 <sup>c</sup>	114	95	4	18 <sup>c</sup>	O <sub>c</sub>	100 <sup>c</sup>	
C. perfringens	425	83	12	23 <sup>c</sup>	83°	9°	425	100	0	
Clostridioides (formerly Clostridium) difficile <sup>3</sup>	1013	32	38	480	74	25	1343	100	0	
Other <i>Clostridium</i> spp.	461	67	25	71	62	35	461	100	0	

#### **Footnotes**

- a. Data were generated from unique isolates from patient specimens submitted to Tufts Medical Center, Boston, Massachusetts; International Health Management Associates, Inc., Schaumburg, Illinois; R.M. Alden Research Laboratory, Culver City, California; Creighton University School of Medicine, Omaha, Nebraska; Mayo Clinic College of Medicine and Science, Rochester, Minnesota; and the Centers for Disease Control and Prevention, Atlanta, Georgia. All testing was performed by the agar dilution method. Information and analysis of previous versions of this table have been published.
- b. Intermediate category is not shown but can be derived by subtraction of %S and %R for each antimicrobial agent from %100.
- c. Calculated from fewer than the CLSI document M39<sup>1</sup> recommendation of 30 isolates.
- d. A dash (-) indicates that data were not available.
- e. Anaerobic gram-positive cocci include Peptococcus, Peptostreptococcus, Finegoldia, Peptoniphilus, and Anaerococcus species.
- f. 80 isolates of *Cutibacterium* (formerly *Propionibacterium*) *acnes* from two of the sites generated MIC values for rifampin ≤ 0.03 µg/mL using the agar dilution method. There are no interpretive breakpoints for this organism/antimicrobial agent combination.
- g. Clostridioides (formerly Clostridium) difficile isolates are from an intestinal source; these results do not imply efficacy for intraluminal infections. Vancomycin minimal inhibitory concentrations for isolates were < 4 µg/mL.

#### Reference for D2

<sup>1</sup> CLSI. Analysis and Presentation of Cumulative Antimicrobial Susceptibility Test Data; Approved Guideline—Fourth Edition. CLSI document M39-A4. Wayne, PA: Clinical and Laboratory Standards Institute; 2014.

**Development of Resistance and Testing of Repeat Isolates:** Isolates that are initially susceptible may become intermediate or resistant after initiation of therapy. Therefore, subsequent isolates of the same species from a similar body site should be tested in order to detect resistance that may have developed. This can occur within as little as three to four days and has been noted most frequently in *Enterobacter, Citrobacter*, and *Serratia* spp. with third-generation cephalosporins; in *P. aeruginosa* with all antimicrobial agents, and staphylococci with quinolones. For *S. aureus*, vancomycin-susceptible isolates may become vancomycin intermediate during the course of prolonged therapy.

In certain circumstances, testing of subsequent isolates to detect resistance that may have developed might be warranted earlier that within three to four days. The decision to do so requires knowledge of the specific situation and the severity of the patient's condition.

#### **Interpretation of Microbiology Reports**

#### Semi-quantitative Terminology (rare/few/moderate/many):

The amount of each type of organism seen is quantified when reading gram stains using the following interpretive criteria:

Description	No. per oil ir	mmersion field (×1000)
	Cells	Bacteria
Rare	<1	<1
Few	1–5	2–10
Moderate	6–10	11–50
Many	>10	>50

Of note, when a report says, "rare gram negative bacilli" it does not mean rare as in unusual, it means rare as in very few.

#### **Contaminant vs. Pathogen:**

#### **BLOOD**

Normally Sterile

#### **PATHOGENS**

• Any organism isolated

#### **LIKELY CONTAMINANTS**

- Coagulase negative staphylococci
- Alpha-hemolytic streptococci
- Bacillus spp.
- Corynebacterium spp. (except C. jeikeium)
- Cutibacterium acnes (prev. Propionibacterium acnes)

#### **TISSUE AND BODY FLUIDS**

Normally Sterile

**PATHOGENS** - Any organism isolated; use judgment to evaluate the possibility of normal flora being present in relation to the source of specimen.

#### EYE/EAR

#### **NORMAL FLORA:**

- coagulase negative Staphylococci
- Non-hemolytic streptococci
- Alpha-hemolytic streptococci
- Diphtheroids

#### SKIN

#### **NORMAL FLORA:**

- coagulase negative staphylococci
- Cutibacterium acnes (prev. Propionibacterium acnes)
- Diphtheroids
- Alpha-hemolytic streptococci
- Bacillus spp.

#### **GENITAL**

#### **PATHOGENS**

- Neisseria gonorrhoeae
- Beta-hemolytic streptococci (GBS will be held if present for susceptibility testing if needed)
- Predominant growth of Yeast or S aureus

#### URINE

Should be sterile. Cultures with mixed flora will be reported as such if contamination is suspected.

#### **PATHOGENS**

- Enterobacterales
- Enterococcus spp.
- Pseudomonas spp. and other non-fermenters
- Group B Streptococcus (Streptococcus agalactiae)
- S. aureus and S. saprophyticus
- Yeast
- Aerococcus urinae

#### STOOL (CULTURES NO LONGER PERFORMED IN-HOUSE):

#### **ENTERIC PATHOGEN TESTING BY PCR/HYBRIDIZATION:**

- Shigella spp.
- Salmonella spp.
- Campylobacter group (C. coli, C. jejuni, and C. lari)
- Yersinia enterocolitica
- Vibrio group (V. cholera and V. parahaemolyticus)
- Shiga-toxin producing strains of E coli (both shiga toxin 1 and 2)
- Norovirus (GI and GII)
- Rotavirus A

Note: If cultures for *Aeromonas or Plesiomonas* are needed, specific orders for these organisms will need to be requested and these specimens will be referred to the reference laboratory.

#### RESPIRATORY TRACT

#### **PATHOGENS:**

- Group A Streptococcus (Streptococcus pyogenes)
- Group C Streptococcus (large colony)
- Group G Streptococcus (large colony)
- Arcanobacterium haemolyticum
- Streptococcus constellatus subsp. pharyngis
- Streptococcus pneumoniae
- Haemophilus influenzae
- Moraxella catarrhalis (predominant)
- Enterobacterales (lower respiratory tract)
- Pseudomonas spp. and other non-fermenters
- Burkholderia cepacia
- Yeast (predominant)
- S. aureus (predominant)

#### **Specimen Requirements**

Specimen requirements can be found by going to the on-line LMH Lab Test Directory (link as follows): <a href="http://www.testmenu.com/lmh">http://www.testmenu.com/lmh</a>

#### **Timing of Reports**

#### **Preliminary/Final Reports:**

Cultures are examined each day and preliminary reports are generated on a daily basis that include any information or presumptive identification of organisms isolated that would be helpful to the physician. When the culture is completed, a final report is released.

#### **Gram Stain:**

Surgical specimens, sterile body fluids and bronch washings and brushings-within 1 hour of receipt in lab

#### Organism identification:

The organism will be identified within 24 hours of isolation of an organism unless it is an unusual or fastidious organism and/or requires further work-up to confirm.

#### Susceptibility results:

The susceptibility results will be reported within 24-48 hours of isolation of an organism unless it is an unusual or fastidious organism and/or requires further work-up to confirm identification.

#### **Antibiogram:**

The hospital-wide antibiogram can be found on the intranet by selecting resources, clinical, medical staff, resources, antibiogram. It is updated every 12 months and approved by the antibiotic stewardship committee.

#### **Urinalysis and Urine Culture**

#### **Indicators of Infection from a Urinalysis:**

- Positive nitrite
- >10 WBC's/hpf on patients >60 years of age
- >10 WBC's/hpf and epithelial cells <16/hpf on patients <60 years of age (if it is a cath specimen, epi's don't count)

Note: A urine culture should be ordered separately for neutropenic patients with possible urinary tract infection. Urinalysis with culture if indicated is not appropriate for those patients.

#### **Urine Culture:**

A urine culture must ALWAYS be interpreted in the context of a urinalysis and patient symptoms. Ideally, a urine culture would not be performed unless a urinalysis indicated a possible infection. Typically, catheterized patients can become colonized within 48 hours of catheterization. The only

patient populations for which it is recommended to screen for and treat asymptomatic bacteriuria are pregnant women and patients scheduled for genitourinary surgical procedure.

Urine cultures are held for 2 days before finalizing as "No Growth."

In February 2020 The American College of Obstetricians and Gynecologists (ACOG) published Committee Opinion number 797 to update the Guidelines for the Prevention of Perinatal Group B Streptococcal (GBS) Disease published in Nov. 2010. It notes that the presence of group B Streptococcus at any colony count during pregnancy is considered indicative of heavy maternal colonization of GBS in the vaginal flora. (i.e. Maternal GBS bacteriuria at any point during pregnancy is a recognized risk factor for early-onset GBS disease and therefore has been included as an indication for intrapartum antibiotic prophylaxis.)

If GBS is present as the predominant organism, susceptibility will be performed and reported. If susceptibility testing is not performed and the patient is female, the organism will be held for 2 weeks and the following comment is attached to the report: "Presence of group B Strep in urine samples of pregnant individuals during any trimester is indicative of heavy maternal GBS colonization if the patient is pregnant and penicillin allergic, notify the laboratory if susceptibility testing is required. Disregard if susceptibility testing was performed."

#### **Stool Testing**

**Stool for Lactoferrin:** The presence of fecal lactoferrin suggests inflammation of the bowel. This should lead the physician to evaluate for the cause of inflammation and consider selective cultures for the most common invasive pathogens such as: *Campylobacter* spp., *Salmonella* spp., *Shigella* spp., and enterohemorrhagic *E coli* 0157:H7.

**Enteric pathogen (EP) testing:** Stool cultures (in-house) were discontinued in July 2015. Testing of liquid and soft stool specimens is now performed on the Verigene Nanosphere system. This assay utilizes amplification (PCR) and hybridization to qualitatively detect and identify common gastrointestinal pathogens.

The EP assay detects and identifies the following enteric bacteria/toxins and viruses: *Campylobacter group (C. coli, C. jejuni, and C. lari), Salmonella* species, *Shigella* species, *Vibrio* group (*V. cholera* and *V. parahaemolyticus), Yersinia enterocolitica,* Shiga toxins 1 and 2 (STEC), Norovirus and Rotavirus. If the sample is positive for *Vibrio, Salmonella, Shigella* or STEC, the specimen will be sent to KDHE for further identification.

Note: This assay is FDA-approved for testing of liquid or soft stools. Testing of formed stools is not indicated.

If *Plesiomonas shigelloides or Aeromonas* spp. are suspected, separate orders are required for referral to the reference laboratory.

**Note:** It is inappropriate to order enteric pathogens testing on patients who have been hospitalized for more than 3 days and then develop diarrhea. In these situations, studies have shown that the most common pathogen is *C.difficile* and PCR testing should be ordered.

Stool should be tested for *Clostridioides* (prev. Clostridium) difficile on patients over 6 months of age with clinically significant diarrhea and a history of antibiotic exposure. Current College of American

Pathologists (CAP) guidelines indicate that a provider should consider *C. difficile* testing as an alternative to routine microbiologic studies for inpatients over 6 months of age who have test requests for routine enteric pathogens.

Additionally, CAP clinical guidelines indicate that no more than 2 stool specimens/patient should be accepted without prior consultation with the provider who can explain the limited yield provided by additional specimens.

#### Clostridioides (prev. Clostridium) difficile testing by PCR:

The Cepheid Gene XPert C diff/epi assay is the test currently used for diagnosing *C. difficile* infection (CDI) at LMH. This assay is a qualitative *in vitro* diagnostic test for rapid detection of toxin B gene sequences and for the presumptive identification of 027/NAP1/BI strains (epidemic, "epi"), of toxigenic *C. difficile* from unformed (liquid or soft) stool specimens collected from patients suspected of having CDI.

The test utilizes automated real-time polymerase chain reaction (PCR) to detect toxin gene sequences associated with toxin producing *C. difficile*. The assay is intended as an aid in the diagnosis of CDI.

Detection of 027/NAP1/B1 strains of *C. difficile* by the assay is presumptive and is solely for epidemiological purposes and is not intended to guide or monitor treatment for CDI. 027/NAP1/B1 strains of *C. difficile* have been referred to as "hypervirulent." These strains exhibit increased toxin production and are thought to produce more spores leading to enhanced persistence in the environment.

<u>Testing will not be performed on formed stool (stool that does not take the shape of the container)</u> unless a rare case of ileus is suspected. In cases of suspected ileus, a formed stool may be tested by special request of the physician.

The *C. difficile* assay should not be used to assess response to therapy. Patients can continue to test positive after treatment.

#### **Blood Cultures**

A minimum of two sets (four bottles; one set = one aerobic bottle + one anaerobic bottle) should usually be obtained. The suggested volume of blood per bottle for adults is 10 ml.

Ordering one set may lead to confusion if the culture is positive for an organism that is commonly a contaminant. For example, if one set is ordered and is positive for coagulase-negative staphylococci (CoNS), a common contaminant, it is difficult to determine if this represents contamination or infection. However, if two sets are ordered, and only one is positive for CoNS, this most likely represents contamination.

Please specify the desired sites of the blood draw (e.g., one from line, one peripherally). Ideally, blood cultures should be drawn before the first dose of antibiotics, but antibiotics should not be withheld because of a delay in getting cultures drawn. Although it is common practice to wait 30-60 minutes between blood cultures, there is little data to support this practice, and we do not recommend it.

If a vascular catheter is thought to be a potential site of infection, blood should be drawn from the catheter and the periphery. Site and time of phlebotomy should be noted. The differential time to positivity can help in assessing whether the catheter is the likely source.

Differential Time to Positivity (DTP)

- a. A positive line culture result is obtained at least 2 hours earlier than a positive peripheral blood culture result.
- b. Typically, the diagnosis of a line-associated infection can be made according to the following criteria: presence of an intravascular device, at least one positive blood culture obtained from a peripheral site, clinical manifestations of infection, and no other apparent source for bloodstream infection

Blood Cultures are held for 5 days prior to reporting the culture as final for no growth.

Blood cultures are continuously monitored by the BacTAlert 3D analyzer. If a positive bottle is detected, a gram stain is prepared and the bottle is sub-cultured at that time. The result of the gram stain is called to the physician immediately.

If the Gram stain indicates the presence of Gram positive cocci or bacilli, PCR testing will be performed on the GenMark analyzer for the presence or absence of targets specific for:

Streptococccus pyogenes (group A Beta Strep)

Streptococcus agalactiae (group B Beta Strep)

Streptococcus pneumoniae (Streptococcus mitis can cross react with S pneumo target)

Streptococcus anginosus group

Streptococcus species

Enterococcus faecalis

Enterococcus faecium

*Listeria* species

Listeria monocytogenes

Bacillus cereus group

Bacillus subtilis group

Cutibacterium acnes (prev. Propionibacterium acnes)

Lactobacillus species

Staphylococcus species

Staphylococcus aureus

Staphylococcus epidermidis

Staphylococcus lugdunensis

mecA and mecC for methicillin resistance in Staph species

vanA and vanB for vancomycin resistance in Enterococcus

The assay includes a pan Candida and pan Gram negative target to detect mixed infections, further testing would be performed if positive.

If the Gram stain indicates the presence of Gram negative bacilli (rods), micro-array testing will be performed on the GenMark analyzer for the presence or absence of targets specific for:

Acinetobacter baumannii

Bacteroides fragilis

Citrobacter species

Enterobacter cloacae complex and Enterobacter non-cloacae complex

E coli

Fusobacterium necrophorum and Fusobacterium nucleatum

Haemophilus influenzae

Klebsiella oxytoca

Klebsiella pneumoniae group

Morganella morganii

Neisseria meningitidis

Proteus species and Proteus mirabilis

Pseudomonas aeruginosa

Salmonella

Serratia and Serratia marcescens

Stenotrophomonas maltophilia

The assay includes a pan Candida and pan Gram positive target to detect mixed infections, further testing would be performed if positive.

The assay can also detect the CTX-M, KPC, NDM, OXA, IMP and VIM resistance mechanisms if an organism is detected.

If yeast are present on Gram Stain a fungal panel can be performed on the GenMark analyzer that can detect the following fungal species: *Candida albicans, Candida auris, Candida dubliniensis, Candida* 

famata, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Cryptococcus gattii, Cryptococcus neoformans, Fusarium and Rhodotorula.

#### **Respiratory Cultures/Testing**

**Lower respiratory tract:** Appropriate specimens to identify pathogens causing disease of the lower respiratory tract (tracheitis, bronchitis, pneumonia, lung abscess, and empyema) include expectorated and induced sputum, endotracheal tube aspirations, bronchial brushings, washes, or alveolar lavages collected during bronchoscopy and pleural fluid.

**Sputum evaluations** are performed on all sputum specimens using the gram stain to assess for quality (lack of contaminating oral respiratory tract flora and epithelial cells). An acceptable specimen generally yields less than 10 squamous epithelial cells per low power field. The presence of 25 or more polymorphonuclear leukocytes per low power field together with few squamous epithelial cells, implies an excellent specimen.

If the specimen is of poor quality, a comment is attached to the gram stain report, "Suggests orophapharyngeal contamination, interpret accordingly." The number of white blood cells may not always be relevant, because many patients are severely neutropenic and specimens from these patients will not show white blood cells on gram stain examination. If the specimen is from a patient (from either the ED or an inpatient floor) with a diagnosis of pneumonia and the specimen is poor quality, a new specimen may be requested.

**Upper respiratory tract:** Appropriate specimens to identify pathogens causing upper respiratory tract infections include samples from the nasopharynx, throat, oral ulcerations, and inflammatory material from the nasal sinuses.

Specific pathogens or normal respiratory flora are quantified in the culture report using the terms –many, -moderate, or –few.

All negative rapid strep tests for group A Strep are followed up with culture for group A, C, F, and G Beta hemolytic *Streptococcus*, *Arcanobacterium haemolyticum* and *Streptococcus constellatus ssp. pharyngis*.

**Legionella** and **S pneumo** antigen testing: It is important to note that as an alternative to culture for *Legionella*, the laboratory offers a highly sensitive rapid EIA for the detection of *L. pneumophilia* antigen in the urine of patients suspected of having legionellosis. The laboratory offers a similar rapid EIA test for the detection of *S. pneumoniae* antigen in the urine of patients suspected of having pneumococcal pneumonia or sepsis. Note: A negative result does not rule out the possibility of infection but indicates that the antigen may be below the limits of detection for this assay.

**Mycoplasma pneumoniae and Chlamydia pneumoniae:** See respiratory panel in Virology section to follow. Testing is performed as part of the Respiratory panel on the GenMark by pcr.

#### **Genital Tract Cultures/Testing**

Genital tract cultures are performed in the microbiology laboratory to determine the etiology of various clinical syndromes, including vulvovaginitis, genital ulcers, urethritis, cervicitis, endometritis, salpingitis, and ovarian abscess in females and urethritis, epididymitis, prostatitis, and genital ulcers in males.

Genital tract cultures are held for 3 days before finalizing the report.

**Bacterial Vaginosis:** The gram stain is the preferred method for detecting bacterial vaginosis (BV). While it is true that *Gardnerella vaginalis* (one of the prevalent organisms in BV) is easily cultured, it can be present in greater than 60% of normal patients. The presence of "clue cells" an indication of BV, can only be determined from a gram stain and culturing the anaerobic bacteria associated with BV is too costly and time-consuming, therefore the culture should not be used for the diagnosis of bacterial vaginosis.

In our laboratory graded criteria for evaluation of the gram stain is used for determining the presence or absence of suspected bacterial vaginosis. One of the following interpretations is attached to the gram stain report: "No Evidence of Bacterial Vaginosis" (Grade 1); "Intermediate for Bacterial Vaginosis" (Grade 2); or "Bacterial Vaginosis Indicated upon Smear Review." If the specimen quality was poor a comment will be attached to the report indicating "Submitted specimen does not have sufficient vaginal material to evaluate for bacterial vaginosis."

**GBS screens:** Our laboratory offers two separate orders for GBS screens on pregnant females at 36 to 37 weeks of gestation. STREPB is the pneumonic developed at LMH for GBS screening on non-allergic patients. STRPBS is the pneumonic developed at LMH for GBS screening on penicillin-allergic patients to ensure that if the test is positive susceptibility testing will be performed.

**Chlamydia/GC testing:** The laboratory offers in-house testing using real-time PCR for the detection of *Chlamydia trachomatis* (CT) and *Neisseria gonorrhoeae* (NG) from endocervical swabs, rectal swabs, pharyngeal swabs, and urine specimens. A specimen adequacy control has been added to this test to detect human cells and DNA which ensures the specimen quality is acceptable.

Results are reported as Detected or Not Detected for each specific organism. An Invalid result can be due to the presence of interfering substances or it can be due to the absence of human cells or DNA.

This test should not be used for the evaluation of suspected sexual abuse or for other medico-legal indications. This assay has not been evaluated in patients less than 14 years of age. NOTE: A negative test result does not exclude the possibility of infection because test results may be affected by improper specimen collection, concurrent antibiotic therapy or the number of organisms in the specimen which may be below the sensitivity of the test.

The test takes 90 minutes to complete. Specimens collected from patients in the Emergency Department will be performed stat. Specimens received from clinics will be performed on a routine basis ensuring results in less than 24 hours Monday through Friday.

The sensitivities and specificities of the various specimen types are as follows:

	Chlamydia		Gonorrhea	
Sample type	Sensitivity	Specificity	Sensitivity	Specificity
Endocervical swabs	96.0%	99.6%	100%	>99.9%
Female Urine	98.1%	99.8%	94.4%	>99.9%
Male Urine	98.5%	99.8%	98.3%	99.9%
Pharyngeal swabs	95.9%	99.7%	94.7%	98.8%
Rectal swabs	86.0%	99.4%	91.2%	99.6%

#### **Trichomonas vaginalis (TVPCR):**

The laboratory offers in-house testing using real-time PCR for the detection of *Trichomonas vaginalis* (TV) from both endocervical swabs and urine specimens. A specimen adequacy control has been added to this test to detect human cells and DNA which ensures the specimen quality is acceptable.

Results are reported as Detected or Not Detected. An Invalid result can be due to the presence of interfering substances (such as blood or mucous) or it can be due to the absence of human cells or DNA.

This test should not be used for the evaluation of suspected sexual abuse or for other medico-legal indications.

A negative test result does not exclude the possibility of infection because test results may be affected by improper specimen collection, concurrent antibiotic therapy or the number of organisms in the specimen which may be below the sensitivity of the test.

The test takes 70 minutes to complete, however positives may come off as soon as 40 minutes. Specimens collected from patients in the Emergency Department will be performed stat. Specimens received from clinics will be performed on a routine basis ensuring results in less than 24 hours Monday through Friday.

The sensitivities and specificities of the various specimen types are as follows:

	<b>Endocervical Swabs</b>	Urine
	TV	TV
Sensitivity	98.9%	98.4%
Specificity	98.9%	99.7%

#### Mycobacteriology

Specimens submitted to the laboratory for acid fast culture will be sent to reference laboratory for testing. The reference laboratory will perform a culture and fluorescent stain for *Mycobacterium* spp. and will concentrate specimens as necessary.

#### **Specimen requirements:**

**Sputum:** Spontaneously produced sputum is the specimen of choice. A good sputum is 5-10 mls with a minimal amount of oral or nasal secretion. Three specimens (at least one must be a first morning specimen) should be submitted, refrigerated until processed, are desirable. These specimens should be collected at least 8 hours apart. Respiratory Therapy may collect an induced specimen using inhalation treatment or nebulization. See notation regarding Acid Fast Bacilli Culture and Smear with PCR for M-TB complex below.

<u>Urine:</u> Early morning clean-voided specimens are preferred. Specimens should be submitted daily for at least 3 days. Refrigeration prior to processing is necessary. Direct smears are not prepared on urine specimens. Pooled specimens are not desirable because of excessive dilution, higher contamination and difficulty in concentrating.

<u>Tissue and Body Fluids:</u> All body fluids, exudates, and tissue should be submitted in sterile containers. Large amounts are preferable. Small amounts of exudates may require moistening with sterile saline and tissues may be sent in a small amount of sterile saline.

<u>Gastric Lavages:</u> Optimal time for gastric lavage is early in the morning before meals. The objective of gastric lavage is to obtain sputum that may have been swallowed during the night. The specimen should be obtained at least 8 hours after the patient has eaten or taken oral drugs. The procedure should be limited to senile, non-ambulatory patients, children younger than 3 years of age, and patients who fail to produce sputum by aerosol induction. Specimens should be collected as a series of three specimens collected on separate days.

A preliminary report is received from the state within 1-2 days. A final report, if negative, is issued at 8 weeks. Positive interim reports from DNA probes may be available in 3 days. Positive culture results will take 2-4 weeks.

The patient's physician is notified immediately if any report is positive for AFB.

The reference laboratory will perform sensitivities on *M. tuberculosis* isolates only.

In July 2016, the microbiology laboratory added a new molecular test on the Cepheid Gene Xpert for the rapid detection of *Mycobacterium tuberculosis* complex (MTB) and rifampin resistance associated mutations of the rpoB gene from both raw induced or expectorated sputum specimens. This MTB/RIF assay does not differentiate between the species of MTB-complex (*M. tuberculosis, M. bovis, M. africanum, M. canettii, M. microti, M. caprae, M. pinnipedi, M. mungi,* and *M. orygis*).

This assay is intended for use with specimens from patients for whom there is clinical suspicion of TB and who have received no more than 3 days of therapy of anti-tuberculosis therapy. This test is intended as an aid in the diagnosis of pulmonary tuberculosis when used in conjunction with clinical and other laboratory findings.

A result of "MTB not detected" from two sputum specimens is highly predictive of the absence of *M. tuberculosis* complex and can be used as an aid in the decision of whether continued airborne infection isolation (AII) is warranted in patients with suspected pulmonary tuberculosis.

The test name is "AFB by PCR with culture (AFBPCR). The test takes 2.5 hours to complete. Testing will be performed on the day shift only. This assay is not FDA approved for bronchial washings (BW's) or broncho-alveolar lavage (BAL) specimens.

A negative test result does not exclude the possibility of isolating MTB-complex from the sputum sample. An un-concentrated, non-fluorescent stain will be performed in-house on any sputum with AFBPCR ordered.

#### Virology

**Virus detection-** There are four general ways in which viral infections can be detected: culture, direct viral antigen detection, serology, or nucleic acid detection.

**Culture:** Specimens for culture will be sent to the reference laboratory. The site of collection must be noted for the reference laboratory to perform testing. The specimen should preferably be submitted in viral transport media (a sterile container is acceptable) and delivered to the laboratory as soon as possible.

Influenza A/B and RSV by PCR: PCR testing for Influenza A, Influenza B, and RSV (subtypes A and B) is performed in-house. The specimen requirement is a nasopharyngeal swab in viral media (UTM kit). The specimen should be refrigerated if there is more than a 24 hour delay in delivery to the laboratory. The assay takes 30 minutes to perform. Testing of inpatients and ED patients for Influenza/RSV are available as a combination panel only. Note: The option to order individual tests for influenza A/B and RSV is available for outpatients only.

Rapid antigen based tests are no longer performed in the laboratory for Respiratory Syncytial Virus (RSV) and Influenza A/B. All influenza and RSV testing is performed using PCR on a nasopharyngeal swab in viral media (UTM or M4 kit).

**Sars-Cov-2:** Testing is available on multiple platforms for Sars-Cov-2. The specimen requirements are for nasopharyngeal swab in VTM, UTM, or M4 saline can only be used for the Diasorin and Gene xpert assays at this time).

Respiratory Pathogen Panel: The laboratory offers in-house qualitative testing using PCR for the detection of: Influenza A, Influenza A (subtype H1), Influenza A (subtype H1N1, 2009), Influenza A (subtype H3), Influenza B, Parainfluenza 1, Parainfluenza 2, Parainfluenza 3, Parainfluenza 4, RSV A, RSV B, Rhinovirus/Enterovirus, Coronavirus, SARS-CoV-2, Adenovirus, human Metapneumovirus, *Mycoplasma pneumoniae*, and *Chlamydia pneumoniae* from nasopharyngeal swabs in viral transport Media. Other specimen types have not been validated.

Results are reported as Detected or Not Detected. A specimen yielding a negative result may contain respiratory viruses or bacteria other than those included in the assay. A negative test result does not exclude the possibility of infection because test results may be affected by improper specimen collection, concurrent antibiotic therapy or the number of organisms in the specimen which may be below the sensitivity of the test.

The test takes under 2 hours to complete. Specimens collected from inpatients and patients in the Emergency Department will be performed stat. Specimens received from clinics will be performed on a routine basis ensuring results in less than 24 hours.

**Rotavirus Detection:** Rotavirus is a major cause of acute gastroenteritis, especially in children 6 to 24 months in age. In addition, rotavirus infections can produce severe illness as well as asymptomatic infection in adults. The incubation period of rotavirus infection is usually one to three days, followed by gastroenteritis with an average duration of five to eight days. Virus titers in stool reach a maximum shortly after the onset of illness, then decline. Due to inadequacies in existing culture methods, human rotavirus is not routinely isolated from rotavirus-containing specimens.

The Enteric Pathogens (EP) molecular assay detects Rotavirus A in liquid or soft stool. The specimen requirement is a stool specimen in a clean container or in Cary Blair preservative.

**Norovirus Detection:** Noroviruses are highly contagious and cause on average 19-21 million cases of acute gastroenteritis each year ranking norovirus in the top five pathogens for enteric illnesses.

The Enteric Pathogens (EP) molecular assay detects Norovirus GI and GII in liquid or soft stool. The specimen requirement is a stool specimen in a clean container or in Cary Blair preservative.

#### Mycology

**Specimen Collection:** The ideal specimens for fungal isolation are either tissue, sterile body fluid, or blood. If a tissue specimen is to be tested for the presence of fungi, it is important that part of the specimen is sent to the microbiology laboratory **before** the specimen is fixed in formalin for histological examination. Blood to be tested for fungus should be added to a separate blood culture bottle that can be held for 30 days.

**Timing of Reports:** Moulds may take 3-4 weeks to grow, whereas yeasts grow rather rapidly and can usually be identified within 3-5 days. Specimens will be finalized at 4 weeks.

**Susceptibility Testing:** Susceptibility testing is performed only upon special request. If susceptibility testing is needed, please contact the microbiology laboratory at 505-6177. If susceptibility testing is needed for mould, a list of antifungals for testing must be specified for the reference laboratory to perform testing.

**Cryptococcus Antigen Testing in CSF/serum:** The laboratory offers a highly sensitive rapid lateral flow assay that can be performed on serum or CSF. This assay is more sensitive than culture, India ink, latex agglutination and enzyme immunoassay (EIA). The assay will detect antigens for *Cryptococcus* species complex (*Cryptococcus neoformans and Cryptococcus gattii*). This test is not meant to be used as a screening test for the general population. It should only be done when clinical evidence suggests the diagnosis of cryptococcal disease. Fungal culture will be performed on all CSF's submitted for cryptococcal antigen testing.

#### **Parasitology**

**Ova and Parasites:** An O&P test should be ordered on patients presenting with a history of chronic diarrhea (> 10 days). It is **not** appropriate to order an O&P test if the patient develops diarrhea while in the hospital. Due to the low incidence of most parasitic infections in the United States and Kansas, stool specimens for Ova and Parasite are routinely tested only for *Giardia lamblia* and *Cryptosporidium* 

parvum through an EIA test. However, if the patient has a travel history that includes regions of the world where parasitic infections are endemic, microscopic evaluation for ova and parasites can be ordered. Please contact the microbiology lab if this is the case (505-6177). These specimens will then be submitted to the reference lab (ARUP) for further testing.

**Timing of Reports:** A *Giardia/Cryptosporidium* antigen test is available every day of the week and is offered on all shifts.

**Malaria Exam:** Specimens submitted for malaria exam are tested using a Rapid immunochromatographic assay. Malaria smears are sent to the reference laboratory, however, malaria may be found by the hematology department from the differential smear.

#### References:

University of Nebraska Antimicrobial and Clinical Microbiology Guide Book, 2<sup>nd</sup> Edition, 2010, Omaha, NE.

Guidance for Control of Infections with Carbapenem-Resistant or Carbapenemase-Producing *Enterobacteriaceae* in Acute Care Facilities, CDC MMWR, 58(10);256-260, March 20, 2009, http://www.cdc.gov/mmwr/preview/mmwrhtml/mm5810a4.htm.

<u>Bailey and Scott's Diagnostic Microbiology</u>, 15<sup>th</sup> Edition, Mosby Inc. St. Louis, Missouri, 2022.

Manual of Clinical Microbiology, 11th Ed., Washington, D.C., ASM Press, 2015.

Performance Standards for Antimicrobial Susceptibility Testing; Thirty-second Edition Informational Supplement, M100-S, Clinical and Laboratory Standards Institute, January 2022.

# LMH Antimicrobial Formulary January 2023

(Key: \$: <\$25; \$\$: \$26-\$100; \$\$\$: \$101-\$150; \$\$\$\$: >\$150)

## Antifungals

Azole Antifungals		
Fluconazole 200 mg premix (DIFLUCAN ) 200 mg 100 mL Bag	\$	
Fluconazole 400 mg premix (DIFLUCAN ) 400 mg 200 mL Bag	\$	
Fluconazole susp (DIFLUCAN ) 200 mg 5 mL suspension	\$	
Fluconazole tab (DIFLUCAN) 50 mg 1 tablet	\$	
Fluconazole tab (DIFLUCAN ) 100 mg 1 tablet	\$	
Itraconazole (SPORANOX ) 100 mg 1 capsule	\$	
Ketoconazole (NIZORAL ) 200 mg 1 tablet	\$	
Voriconazole (VFEND ) 50 mg 1 tablet	\$	
Voriconazole (VFEND ) 200 mg 1 tablet	\$\$	
Voriconazole inj (VFEND) 200mg 20 mL injection	\$\$	
Echinocandins		
Micafungin (MYCAMINE ) 50 mg 5 mL injection	\$\$\$	
Micafungin (MYCAMINE) 100 mg 5 mL injection	\$\$\$\$	
Misc Antifungals		
Betameth/mupir/miconazole oint(- )	\$\$	
Terbinafine (LamISIL ) 250 mg 1 tablet		
Polyenes		
Amphotericin B Inj (AMPHOTERICIN B ) 50 mg 1 vial injection	\$	
Amphotericin B LIPOSOMAL(Ambisome ) 50 mg 12.5 mL injection	\$\$\$\$	
Amphotericin B Inj (ABELCET) 100 mg 20 mL injection		
Anti-infectives		
Aminoglycosides		
	\$	
Amikacin Inj. (AMIKACIN SULFATE ) 500 mg 2 mL inj		
Gentamicin (baby/peds) PF (AENTAMICIN) 20 mg 2 mL inj		
Gentamicin (GARAMYCIN ) 80 mg 2 mL injection	\$	
Gentamicin (GENTAMICIN) 40 mg 1 mL inj	\$	
-		

Gentamicin (GENTAMICIN SULFATE, INJECTABLE ) 40 mg 1 mL inject	ion \$
Neomycin (NEOMYCIN SULFATE ) 500 mg 1 tablet	\$
Tobramycin Inhalation (TOBI ) 300 mg 5 mL Inhaler	\$
Tobramycin (NEBCIN ) 40 mg 1 mL injection	\$
Tobramycin (NEBCIN ) 80 mg 2 mL injection  Carbapenems	\$
•	\$\$
Ertapenem (INVanz ) 1,000 mg 1 vial injection	
ImiPENem-cilastatin (PRIMAXIN IV ) 250 mg 1 vial injection	\$\$
ImiPENem-cilastatin (PRIMAXIN IV ) 500 mg 1 vial injection	\$\$
MEROpenem (MERREM ) 500 mg 1 vial injection	\$\$
MEROpenem (MERREM ) 1,000 mg 1 vial injection	\$\$
Cephalosporins	
First generation Cephalosporins	
Cefadroxil (DURICEF ) 500 mg 1 capsule	\$
CeFAZolin duplex bag (ANCEF ) 1,000 mg 50 mL duplex bag	\$
CeFAZolin premix bag (ANCEF ) 1,000 mg 50 mL Bag	\$
CeFAZolin vial (ANCEF ) 1,000 mg 1 vial inj	\$
Cephalexin 200ml susp (KEFLEX ) 250 mg 5 mL suspension	\$
Cephalexin susp (KEFLEX ) 125 mg 5 mL suspension	\$
Cephalexin (KEFLEX ) 500 mg 1 capsule	\$
Second generation Cephalosporins	
CefoTEtan vial(CEFOTAN ) 1,000 mg 1 vial injection	\$
CefoTEtan vial (CEFOTAN ) 2,000 mg 1 vial injection	\$\$
CefOXItin duplex (MeFOXin ) 1,000 mg 50 mL duplex bag	\$
CefOXItin duplex (MeFOXin ) 2,000 mg 50 mL duplex bag	\$
CefOXitin (MeFOXin ) 1,000 mg 1 vial inj	\$
CefOXitin (MeFOXin ) 2,000 mg 1 vial inj	\$\$
CeFUROxime (CEFTIN) 250 mg 1 tablet	\$
CeFUROxime (ZINACEF ) 1,500 mg 1 vial injection	\$
Third generation Cephalosporins	
Cefixime (SUPRAX ) 400 mg 1 tablet	\$
Cefdinir (OMNICEF) 300 mg 1 tablet	\$

Cefdinir (OMNICEF) 250mg 5 mL suspension	\$
Cefpodoxime susp (VANTIN ) 50 mg 5 mL suspension	\$
Cefpodoxime (VANTIN ) 200 mg 1 tablet	\$
CefTAZidime (FORTAZ ) 1,000 mg 1 vial injection	\$
CefTAZidime/Avibactam (AVYCAZ) 2,500 mg 1 vial injection	\$\$\$\$
CefTRIAXone premix (ROCEPHIN ) 1,000 mg 50 mL Bag	\$
CefTRIAXone premix (ROCEPHIN ) 2,000 mg 50 mL Bag	\$
CefTRIAXone duplex (ROCEPHIN ) 1,000 mg 50 mL duplex bag	\$\$
CefTRIAXone duplex (ROCEPHIN ) 2,000 mg 50 mL duplex bag	\$\$\$
CefTRIAXone vial (ROCEPHIN ) 500 mg 1 vial inj	\$\$
CefTRIAXone vial (ROCEPHIN ) 1,000 mg 1 vial inj	\$\$
CefTRIAXone vial (ROCEPHIN ) 2,000 mg 1 vial inj	\$\$\$
Fourth generation Cephalosporins	
CefePIME (MAXIPIME ) 1 g 1 vial inj	\$
CefePIME (MAXIPIME ) 2 g 1 vial inj	\$\$
	\$\$
CefePIME (MAXIPIME ) 2 g 1 vial inj	\$\$ \$\$
CefePIME (MAXIPIME ) 2 g 1 vial inj  Glycylcyclines	
CefePIME (MAXIPIME ) 2 g 1 vial inj  Glycylcyclines  Tigecycline (TYGACIL ) 50 mg 1 vial IV Piggyback	
CefePIME (MAXIPIME ) 2 g 1 vial inj  Glycylcyclines  Tigecycline (TYGACIL ) 50 mg 1 vial IV Piggyback  Leprostatics	\$\$

Clindamycin premix bag (CLEOCIN) 600 mg 50 mL Bag	\$
Clindamycin premix bag (CLEOCIN ) 900 mg 50 mL Bag	\$
Clindamycin susp (CLEOCIN PEDIATRIC ) 75 mg 5 mL suspension	\$
Clindamycin (CLEOCIN PHOSPHATE ) 300 mg 2 mL inj	\$
Clindamycin (CLEOCIN PHOSPHATE ) 600 mg 4 mL injection	\$
Clindamycin (CLEOCIN PHOSPHATE ) 900 mg 6 mL injection	\$
Macrolides	
Azithromycin (ZITHROMAX ) 100 mg 5 mL suspension	\$
Azithromycin(ZITHROMAX) 200 mg 5 mL suspension	\$
Azithromycin(ZITHROMAX) 250 mg 1 tablet	\$
Erythromycin susp (Eryped 200 ) 200 mg 5 mL Susp	\$
Erythromycin (- ) 500 mg 10 mL inj	\$
Erythromycin(ERY-TAB ) 250 mg 1 tab DR Tab	\$
, , , , , , , , , , , , , , , , , , , ,	
Fidaxomicin(Dificid ) 200 mg 1 tablet	\$\$\$\$
Fidaxomicin(Dificid ) 200 mg 1 tablet  Miscellaneous antibiotics	\$\$\$\$
	\$\$\$\$ \$
Miscellaneous antibiotics	
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM ) 500 mg injection	\$
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM ) 500 mg injection  Aztreonam Inj (AZACTAM ) 1,000 mg 1 vial injection	\$ \$\$
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM ) 500 mg injection  Aztreonam Inj (AZACTAM ) 1,000 mg 1 vial injection  Aztreonam Inj (AZACTAM ) 2,000 mg 1 vial injection	\$ \$\$ \$\$
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM ) 500 mg injection  Aztreonam Inj (AZACTAM ) 1,000 mg 1 vial injection  Aztreonam Inj (AZACTAM ) 2,000 mg 1 vial injection  Bacitracin irrigation (- ) 0 250 mL Irrigation  Bacitracin (BACITRACIN ) 50,000 unit 1 vial injection	\$ \$\$ \$\$ \$\$
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM ) 500 mg injection  Aztreonam Inj (AZACTAM ) 1,000 mg 1 vial injection  Aztreonam Inj (AZACTAM ) 2,000 mg 1 vial injection  Bacitracin irrigation (- ) 0 250 mL Irrigation  Bacitracin (BACITRACIN ) 50,000 unit 1 vial injection  Ceftaroline	\$ \$\$ \$\$ \$
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM) 500 mg injection  Aztreonam Inj (AZACTAM) 1,000 mg 1 vial injection  Aztreonam Inj (AZACTAM) 2,000 mg 1 vial injection  Bacitracin irrigation (-) 0 250 mL Irrigation  Bacitracin (BACITRACIN) 50,000 unit 1 vial injection  Ceftaroline  Chloramphenicol (CHLOROMYCETIN) 1,000 mg 1 vial injection	\$ \$\$ \$\$ \$
Miscellaneous antibiotics  Aztreonam Inj (AZACTAM) 500 mg injection  Aztreonam Inj (AZACTAM) 1,000 mg 1 vial injection  Aztreonam Inj (AZACTAM) 2,000 mg 1 vial injection  Bacitracin irrigation (-) 0 250 mL Irrigation  Bacitracin (BACITRACIN) 50,000 unit 1 vial injection  Ceftaroline  Chloramphenicol (CHLOROMYCETIN) 1,000 mg 1 vial injection  Colistimethate (-) 150 mg 1 vial injection	\$ \$\$ \$ \$ \$

	Linezolid premix bag (ZYVOX	) 600 mg	300 mL Bag	\$
	Linezolid (ZYVOX ) 600 mg	ı 1 table	t	\$\$
	Oritavancin (KIMYRSA) 1,2	00 mg 40	o mL inj	\$\$\$\$
	Pentamidine INHALATION (N	IEBUPENT)	300 mg 1 vial susp	\$\$
	Sulfamethoxazole-trimethop	rim DS (BAC	TRIM DS) 16omg 1 tablet	\$
	Sulfamethoxazole-trimethopi	rim inj (BAC	TRIM) 160 mg 10 mL inj	\$
	Sulfamethoxazole-trimethop	rim SS (BAC	TRIM) 8omg 1 tablet	\$
	Sulfamethoxazole-trimethop	rim susp (SU	LFATRIM) 40mg 5 mL Susp	\$
	Vancomycin liq (VANCOMYCI	N HCL) 2	250 mg 5 mL liq	\$
	Vancomycin vial(VANCOMYC	IN HCL)	500 mg 5 mL inj	\$
	Vancomycin vial(VANCOMYC	IN HCL) 1,	000 mg 10 mL inj	\$
	Vancomycin premix (VANCON	MYCIN HCL)	1,000 mg 200 mL inj	\$
	Vancomycin premix (VANCON	MYCIN HCL)	1,250 mg 250 mL inj	\$
	Vancomycin premix (VANCON	MYCIN HCL)	1,500 mg 300 mL inj	\$
	Vancomycin premix (VANCON	MYCIN HCL)	1,750 mg 350 mL inj	\$\$
	Vancomycin premix (VANCON	MYCIN HCL)	2,000 mg 400 mL inj	\$\$
Pe	nicillins			
	Aminopenicillins			
	Amoxicillin susp (TRIMOX )	125 mg	5 mL suspension	\$
	Amoxicillin susp (TRIMOX )	250 mg	5 mL suspension	\$
	Amoxicillin susp (TRIMOX )	400 mg	5 mL suspension	\$
	Amoxicillin (AMOXICILLIN )	250 mg	1 capsule	\$
	Amoxicillin (AMOXICILLIN )	500 mg	1 capsule	\$
	Ampicillin vial (AMPICILLIN )	500 mg	2 mL injection	\$
	Ampicillin vial (AMPICILLIN )	1 <b>,</b> 000 mg	3.5 mL injection	\$
	Ampicillin vial (AMPICILLIN )	2 <b>,</b> 000 mg	1 vial injection	\$
_	Beta-lactamase inhibitors			

Amoxicillin-clavulanate susp(AUGMENTIN ) 400 mg 5 mL suspension	
Amoxicillin-clavulanate susp(AUGMENTIN ) 600 mg 5 mL suspension	\$
Amoxicillin-clavulanate (AUGMENTIN ) 500 mg 1 tablet	\$
Amoxicillin-clavulanate (AUGMENTIN ) 875 mg 1 tablet	\$
Ampicillin-sulbactam (UNASYN ) 1,500 mg 1 vial injection	\$
Ampicillin-sulbactam (UNASYN) 3,000 mg 1 vial injection	\$
Piperacillin-tazo premix (ZOSYN ) 4.5 g 50 mL Bag	\$
Piperacillin-tazo (ZOSYN ) 2.25 g 10 mL inj	\$
Piperacillin-tazo (ZOSYN ) 3.375 g 10 mL inj	\$
Piperacillin-tazo (ZOSYN ) 4.5 g 10 mL inj	\$
Natural penicillins	
Penicillin (IM ONLY)(BICILLIN L-A ) 1.2 MU 2 mL LA inj	\$\$\$
Penicillin G potassium vial 1 MU 2 mL inj	\$ \$
Penicillin G potassium premix 5 MU 100 mL premix bag	\$\$
Penicillin G sodium vial (- ) 5 MU 10 mL inj	\$\$
Penicillin VK tab (- ) 250 mg 1 tablet	\$
Penicillin VK tab (-) 500 mg 1 tablet	\$
Penicillinase resistant penicillins	•
Dicloxacillin (DICLOXACILLIN SODIUM ) 250 mg 1 capsule	<b>d</b>
Nafcillin (- ) 1,000 mg 10 mL injection	\$
Nafcillin (NAFCIL ) 2,000 mg 10 mL injection	\$
Quinolones	
Ciprofloxacin premix bag (CIPRO I.V.) 200 mg 100 mL Bag	\$
Ciprofloxacin premix bag (CIPRO I.V.) 400 mg 200 mL Bag	\$
Ciprofloxacin (CIPRO ) 250 mg 1 tablet	\$
Ciprofloxacin (CIPRO ) 500 mg 1 tablet	\$
Ciprofloxacin (CIPRO ) 500 mg 5 mL Susp	\$
Levofloxacin premixed bag (LEVAQUIN ) 500 mg 100 mL Bag	\$
Levofloxacin premixed bag (LEVAQUIN ) 750 mg 150 mL Bag	\$
Levofloxacin (LEVAQUIN ) 250 mg 1 tablet	\$

Levofloxacin (LEVAQUIN ) 500 mg 1 tablet	\$
Levofloxacin (LEVAQUIN ) 750 mg 1 tablet	\$\$\$\$
Moxifloxacin (AVELOX ) 400 mg 1 tablet	\$\$\$
Moxifloxacin (AVELOX ) 400 mg 250 mL Bag	\$\$\$
Sulfonamides	
SulfaDIAZINE (- ) 500 mg 1 tablet	\$
SulfaSALAzine (Azulfadine ) 500 mg 1 tablet	\$
Tetracyclines	
Demeclocycline (DECLOMYCIN ) 150 mg 1 tablet	\$
Doxycycline inj (VIBRAMYCIN) 100 mg 1 vial injection	\$
Doxycycline susp (VIBRAMYCIN) 25 mg 5 mL suspension	\$
Doxycycline (VIBRAMYCIN) 100 mg 1 tablet	\$
Minocycline (MINOCIN ) 50 mg 1 capsule	\$\$
Minocycline (MINOCIN ) 100 mg 1 vial injection	\$\$
Urinary anti-infectives	
Fosfomycin (MONUROL ) 3 g suspension	\$
Nitrofurantoin monohydrate (MACROBID ) 100 mg 1 capsule	\$
Nitrofurantoin macrocrystals (MACRODANTIN ) 50 mg 1 capsule	\$
Antimalarial agents	
Pyrimethamine (DARAPRIM ) 25 mg 1 table	\$
Anti-tuberculosis agents	
Nicotinic acid derivatives	
Pyrazinamide (- ) 500 mg 1 tablet	\$
Rifamycin derivatives	
Rifampin inj (RIFADIN IV ) 600 mg 10 mL inj	\$
Rifampin (RIFADIN ) 300 mg 1 capsule	\$

Antiviral agents	
Adamantane antivirals	
Amantadine syrup (Symmetrel ) 50 mg 5 mL Syrup	\$
Amantadine (symmetrel ) 100 mg 1 capsule	<b>≯</b> \$
Antiviral chemokine receptor antagonist	
Maraviroc (Selentry ) 300 mg 1 tablet	\$
Antiviral combinations	
Emtricitabine-Tenofovir (TRUVADA ) 200/300 mg 1 tablet	\$\$
	<b>ት</b> ት
Integrase strand transfer inhibitor  Raltegravir (ISENTRESS ) 400 mg 1 tablet	\$
Kaltegravii (i3Ervi KE33) 400 iiig T tablet	,
Miscellaneous antivirals	
Miscellaneous antivirals  Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection	\$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection	\$\$\$\$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection	-
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection Palivizumab (SYNAGIS) 100 mg 1 mL injection  Neuraminidase inhibitors	\$\$\$\$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection Palivizumab (SYNAGIS) 100 mg 1 mL injection	\$\$\$\$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection Palivizumab (SYNAGIS) 100 mg 1 mL injection  Neuraminidase inhibitors Oseltamivir (TAMIFLU ) 60 mg 5 mL suspension	\$\$\$\$ \$\$\$\$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection Palivizumab (SYNAGIS) 100 mg 1 mL injection  Neuraminidase inhibitors  Oseltamivir (TAMIFLU ) 60 mg 5 mL suspension Oseltamivir (TAMIFLU ) 75 mg 1 capsule	\$\$\$\$ \$\$\$\$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection Palivizumab (SYNAGIS) 100 mg 1 mL injection  Neuraminidase inhibitors Oseltamivir (TAMIFLU ) 60 mg 5 mL suspension Oseltamivir (TAMIFLU ) 75 mg 1 capsule  NNRTIS	\$\$\$\$ \$\$\$\$ \$
Foscarnet (FOSCAVIR ) 6,000 mg 250 mL injection Palivizumab (SYNAGIS ) 50 mg 0.5 mL injection Palivizumab (SYNAGIS) 100 mg 1 mL injection  Neuraminidase inhibitors Oseltamivir (TAMIFLU ) 60 mg 5 mL suspension Oseltamivir (TAMIFLU ) 75 mg 1 capsule  NNRTIS  Efavirenz (Sustiva ) 600 mg 1 tablet	\$\$\$\$ \$\$\$\$ \$

Zidovudine (RETROVIR ) 200 mg 20 mL injection	\$
Zidovudine (RETROVIR ) 50 mg 5 mL Syrup	\$
Purine nucleosides	
	_
Acyclovir susp (ZOVIRAX ) 800 mg 20 mL Susp	\$
Acyclovir (ZOVIRAX ) 200 mg 1 capsule	\$
Acyclovir (ZOVIRAX) 800 mg 1 tablet	\$
Cidofovir (VISTIDE ) 375 mg 5 mL injection	\$\$\$\$
Famciclovir (FAMVIR) 250 mg 1 tablet	\$
Famciclovir (FAMVIR ) 500 mg 1 tablet	\$
Ganciclovir (CYTOVENE ) 500 mg 1 vial injection	\$\$

1 capsule

1 tablet

1 Inhaler

\$\$

\$\$\$\$

\$

500 mg

500 mg

6 g

LMH DEPARTMENT OF PHARMACY			
ANTI-INFECTIVES RENAL DOSING CHART			
ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS

Ganciclovir (CYTOVENE )

ValACYclovir (VALTREX )

Ribavirin Inh soln (VIRAZOLE )

A - da tana	F /I - IV 00 h	25.40	F
Acyclovir <b>IV</b>	5 mg/kg IV Q8 hours	25-49:	Frequency to Q12 hours
		10-24:	Frequency to Q24 hours
	CNS infections:	<10	50% normal dose IV Q24
Use <b>ideal body weight</b>	10 mg/kg IV q8 hours		hours
		HD	Q24 hours schedule dose to
			be given after dialysis
Acyclovir PO	Dose and Renal Adjustments Va	ry by Indic	ation. Please refer to
·	appropriate drug reference.	• •	
Amikacin	Extended Interval Dosing:	****	Extended Interval Dosing:
	15 mg/kg IV Q24 hours	40-59:	15 mg/kg IV Q36 hours
Note: Consultation with ID	Note: draw level 6-14 hours	30-39:	15 mg/kg IV Q48 hours
& Pharmacy recommended.	after 1st dose	<30	AVOID, use conventional
			dosing
	Conventional Dosing:	****	Conventional Dosing:
	5-7.5 mg/kg/dose Q8 hours	40-60:	5-7.5 mg/kg Q12 hours
	Note: Peak & trough levels	20-39:	5-7.5 mg/kg Q24 hours
	should be monitored	<20:	5-7.5 mg/kg then monitor
			level
		HD:	5-7.5 mg/kg Q48-72h,
			monitor level prior to HD to
			determine dosing needs
Amoxicillin	250-500 mg PO Q8 hours	10-30:	500 mg PO Q12 hours
		<10:	500 mg PO Q24 hours
		HD:	500 mg PO q24 hours
			schedule dose to be given
			after dialysis
Amoxicillin/ Clavulanate	875 mg PO Q12 hours	10-30:	500/125 mg PO Q12 hours
		<10:	250 mg PO Q12 hours
	500 mg PO Q8 hours	HD:	250-500 mg PO q24 hours
			schedule dost to be given
			after dialysis
	1000/62.5 mg PO q12h (XR	< 30	XR formulation NOT
	formulation)		recommended with CrCl < 30
ANTI-INFECTIVE	NORMAL DOSE	CRCL in	RENAL DOSE ADJUSTMENTS
	-	ml/min	
Ampicillin IV	Continuous Infusion:	****	Continuous Infusion
i .		i .	
	8-12 g IV over 24	30-49:	8 g IV over 24 hours

	Conventional Dosing:	<10: HD: ****	· ·
	1-2 g IV Q4 hours	30-49:	9
	Note: 2g dose is	10-29:	1-2 g IV Q12 hours
	recommended for	<10:	1-2 g IV Q12 hours
	endocarditis, meningitis and bacteremia	HD:	1-2 g IV Q12 hours
Ampicillin/	1.5 g IV Q6 hours	30-49:	1.5-3 g IV Q8 hours
Sulbactam		15-29:	1.5-3 g IV Q12 hours
		5-14:	1.5-3 g IV Q24 hours
		HD:	1.5-3 g IV Q24 hours – GIVE AFTER HD
Azithromycin	Usual: 500mg PO x 1, then 250 or 500 mg PO Q24 hours		hours x 4-10 days r 1 gm PO as a single dose
	Note: Dose and duration vary by	=	_
	reference .		
Aztreonam	1-2 g IV Q8 hours	10-29:	50% of usual dose at usual interval
	Interval to q6 hours for severe infections (esp. P.aeruginosa)	<10:	25% of usual dose at usual interval
		HD:	500 mg IV Q12 hours
Cefadroxil PO	1000 mg PO Q12 hours	25-50:	500 mg PO Q12 hours
	Indication based alternative	10-25:	500 mg PO Q24 hours
	dosing: Please see appropriate reference	<10:	500 mg PO Q36 hours
Cefazolin	1 g IV Q8 hours	10-30:	Frequency to Q12 hours
		<10:	•
	Note: >= 80 kg or severe	HD:	'
	infections may use 2 g		schedule dose to be given after dialysis
			arter dialysis
ANTI-INFECTIVE	NORMAL DOSE	CRCL in	RENAL DOSE ADJUSTMENTS
0.6	1 11/06/10	ml/min	1 11 11 11
Cefepime	1 g IV Q6-12 hours	30-49:	1 g IV Q12 hours
		10-29:	1 g IV Q24 hours
		<10:	1 g IV Q24 hours

	CNS infections or febrile neutropenia 2 g IV Q8 hours	***** 30-49: 10-29: <10: HD:	HD CNS Infections or febrile neutropenia 2 g IV Q12 hours 2 g IV Q24 hours 1 g IV Q24 hours
Cefixime	400 mg PO Q24 hours or 200 mg PO Q12 hours Note: alternate dosing available for STD, please see appropriate drug information reference.	21-60: <20 HD:	50% dose PO Q24 hours
Cefotetan	1 g IV Q12 hours Note:>=80 kg may use 2 g	10-30: <10: HD:	Frequency to Q48 hours
Cefoxitin	1 g IV Q6 hours	10-49:	<b>J</b>
Cefpodoxime	Note:>=80 kg may use 2 g 200-400 mg PO Q12 hours	<10: <30: HD:	200-400 mg PO Q24 hours 200-400 mg PO 3 x week after HD
Ceftaroline	600 mg IV Q12 hours	31-50: 15-30: <15: HD:	300 mg IV Q12 hours 200 mg IV Q12 hours
ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS
Ceftazidime	1 g IV Q8 hours  CNS infections: Use 2g dose with same frequency	30-49: <30: HD:	1 g IV Q24 hours

Ceftriaxone	Meningitis or CNS Infection: 2 g Q12 hours For all indications other than CN 1 g IV Q24 hours		No renal adjustments needed	
Cefuroxime IV	750mg-1.5g IV Q8 hours Note: Frequency can be increased to Q6 hours for life threatening infections	1	10-20: <10: HD:	750 mg IV Q12 hours 750 mg IV Q24 hours 750 mg IV after each dialysis session
Cefuroxime <b>PO</b> tab	250-500 mg PO Q12 hours Note: tablet dosing not equivalent to suspension, we do not carry suspension		<10: HD:	250-500 mg Q24 hours 250-500 mg PO q24 hours schedule dost to be given after dialysis
Cephalexin	500 mg PO Q6 hours	1	l0-30: <10: HD:	500 mg PO Q8 hours 500 mg PO Q12 hours 250-500 mg PO q24 hours schedule dost to be given after dialysis
Ciprofloxacin IV	400 mg IV Q12 hours HAP or VAP: 400 mg IV Q8 hours		<30 HD:	200-400 mg IV Q24 hours 200 mg IV Q24 hours schedule dose to be given after dialysis
Ciprofloxacin <b>PO</b>	500 mg PO Q12 hours	1	l0-29: <10: HD:	250 mg PO Q12 hours 250 mg PO Q24 hours 250 mg PO Q24 hours schedule dose to be given after dialysis
Clarithromycin	500 mg PO Q12 hours	1	l0-29: <10: HD:	250 mg PO Q12 hours 250 mg PO Q24 hours 250 mg PO Q24 hours schedule dose to be given after dialysis
ANTI-INFECTIVE	NORMAL DOSE		CL in /min	RENAL DOSE ADJUSTMENTS
Clindamycin IV	600 mg IV Q8 hours, 900 mg IV hours for necrotizing facilitis  Note: For patients >100 kg 900 lower to the second	/ Q8 No rei		nal adjustment needed

Clindamycin <b>PO</b>	150-450 mg PO q6-8 hours (vari by indication, please use appropriate reference)			For patients >100 kg 450 mg S hours may be used
Daptomycin	4-6 mg/kg IV Q24 hours Note: Not to be used for infections in the lungs; 4 mg/kg for UTI/SSSTI		<30: HD:	4-6 mg/kg IV Q48 hours 4-6 mg/kg IV Q48 hours schedule dose to be given after dialysis
Dicloxacillin	125-500mg PO Q6 hours		No ad	justment needed
Doxycycline IV or PO	100mg Q12 hours		No ad	justment needed
Ertapenem	1000 mg IV Q24 hours		<30: HD:	500 mg IV Q24 hours 500 mg IV Q24 hours schedule dose to be given after dialysis
Erythromycin IV	500-1000 mg IV Q6 hours		<10:	50% PO/IV dose at same interval
Erythromycin <b>PO</b>	Base: 250-500mg PO Q6-12 h EES: 400-800mg PO Q6-12 h	HD/0	CAPD:	Dose the same as CrCl <10 ml/min
Ethambutol		k	****	Initial Treatment:
	Initial: 15 mg/kg PO Q24 hrs Retreatment: 25 mg/kg PO	1	.0-50:	15 mg/kg PO Q24-36 hours
	Q24 hours x 60 days, then 15		<10:	15 mg/kg PO Q48 hours
	mg/kg/day Note: Should not be used		HD:	Administer dose after dialysis
	alone; Monthly eye exams	*	****	Retreatment:
	recommended on high dose regimen; max dose 2.5 gm	1	.0-50:	25 mg/kg PO Q24-36 hours
			<10: HD:	25 mg/kg PO Q48 hours Administer dose after dialysis
Famciclovir	250-500 mg PO Q8 hours	4	10-59:	same dose Q12 hours
	500 mg PO Q8 hours for VZV	2	20-39:	same dose Q24 hours
	Note: no clear adjustment		<20:	50% dose Q24 hours
	recommendations in CAPD		HD:	50% after each dialysis session
Fidaxomicin	200 mg PO Q12 hours			nent needed
ANTI-INFECTIVE	NORMAL DOSE	CRCL in		RENAL DOSE ADJUSTMENTS
		ml/	min	
Fluconazole IV or PO	*Invasive candidiasis: 800 mg (12 mg/kg) load x1, then 400	1	.0-29:	Load x1, then 50% PO/IV dose Q24 hours
	mg (6 mg/kg) PO/IV Q24 hours		<10:	Load x1, then 25% PO/IV dose Q24 hours

		HD: CAPD:	Load x1, then 400 mg (6 mg/kg) after HD 3x/weekly 50% PO/IV Q24 hours
	*Esophageal/oropharyngeal candidiasis: 200 mg PO/IV Q24 hours for esophageal;	<30:	50% PO/IV Q 24 hours
	100 mg Q24 hours for oropharyngeal	HD:	100% PO/IV after each dialysis session
	Note: Some indications may	CAPD:	50% PO/IV Q24 hours
	require different dosing, please refer to appropriate drug reference		
Fosfomycin <b>PO</b>	Uncomplicated cystitis: 3 gm	<50:	Same dose
	x1 Complicated cystitis: 3 gm Q48 hours	<50:	3 gm Q72 hours
Ganciclovir IV	Induction:	****	Induction:
	5 mg/kg IV Q12 hours	50-69:	2.5 mg/kg IV Q12 hours
		25-49:	2.5 mg/kg IV Q24 hours
	Maintenance:	10-24:	1.25 mg/kg IV Q24 hours
	5 mg/kg IV Q24 hours	<10:	1.25 mg/kg IV 3 x week
		HD:	1.25 mg/kg IV 3 x week schedule dose to be given after hemodialysis
			i arter nemodiarvaia
		****	•
			Maintenance:
		50-69:	Maintenance: 2.5 mg/kg IV Q24 hours
			Maintenance:
		50-69: 25-49:	Maintenance: 2.5 mg/kg IV Q24 hours 1.25 mg/kg IV Q24 hours
		50-69: 25-49: 10-24:	Maintenance: 2.5 mg/kg IV Q24 hours 1.25 mg/kg IV Q24 hours 0.625 mg/kg IV Q24 hours
		50-69: 25-49: 10-24: <10:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week  schedule dose to be given
ANTI INICCTIVE	NODMAL DOCE	50-69: 25-49: 10-24: <10: HD:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week  schedule dose to be given after hemodialysis
ANTI-INFECTIVE	NORMAL DOSE	50-69: 25-49: 10-24: <10: HD:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week  schedule dose to be given
ANTI-INFECTIVE Ganciclovir PO	NORMAL DOSE  1000 mg PO Q8 hours	50-69: 25-49: 10-24: <10: HD:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week  schedule dose to be given after hemodialysis
		50-69: 25-49: 10-24: <10: HD:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week  schedule dose to be given after hemodialysis  RENAL DOSE ADJUSTMENTS
		50-69: 25-49: 10-24: <10: HD:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week schedule dose to be given after hemodialysis  RENAL DOSE ADJUSTMENTS  1500 mg PO Q24h or 500
		50-69: 25-49: 10-24: <10: HD: CRCL in ml/min 50-69:	Maintenance:  2.5 mg/kg IV Q24 hours  1.25 mg/kg IV Q24 hours  0.625 mg/kg IV Q24 hours  0.625 mg/kg IV 3 x week  0.625 mg/kg IV 3 x week schedule dose to be given after hemodialysis  RENAL DOSE ADJUSTMENTS  1500 mg PO Q24h or 500 mg PO Q8h

		<10 HD:	500 mg PO 3 x week 500 mg PO 3 x week schedule dose to be given afer hemodialysis	
Gentamicin	Extended Interval Dosing:	****	Extended Interval Dosing:	
	7 mg/kg IV Q24 hours	40-59:	7 mg/kg IV Q36 hours	
Note: Consultation with	Note: draw level 6-14 hours	30-39:	7 mg/kg IV Q48 hours	
Note: Consultation with Pharmacy recommended	after 1st dose	<30	AVOID, use conventional dosing	
		****	Conventional Dosing:	
Exclusion to Extended	Conventional Dosing:	40-60:	1-2.5 mg/kg Q12 hours	
Interval: Pregnancy, breastfeeding, burns (>20%	1-2.5 mg/kg/dose Q8 hours Note: Peak & trough levels	20-39:	1-2.5 mg/kg Q24 hours	
body), ascites,	monitored, consult pharmacy	<20:	1-2.5 mg/kg then monitor	
Enterococcoal endocarditis,		LID.	level 2-3 mg/kg Q48-72h,	
HD or CrCl < 20 ml/min		HD:	monitor level prior to HD to	
			determine dosing needs	
Imipenem/ Cilastatin	250-1000mg IV Q6-8 hours	30-70:	Decrease daily dose by 50%	
			& divide Q6-8 hours (round	
			to nearest 250 mg)	
	Usually 500 mg IV Q6 hours	20-30:	Decrease daily dose by 63%	
	Note: max dose 50mg/kg/day		& divide q8-12 hours (round	
	or 4g/day		to nearest 250 mg)	
		6-20:	, , ,	
			& divide Q12 hours (round	
	Note: Dose varies by severity	<b>&lt;</b> 5:	to nearest 250 mg) Not recommended unless	
	of infection & organism	<b>\</b> 5.	HD being started	
	sensitivity	HD:	_	
	,		& divide q12 hours (round	
			to nearest 250 mg); dose	
			after dialysis on dialysis day	
Isoniazid	300 mg PO Q24 hours (5 mg/kg	PO Q24 hou	rs); max of 300 mg;	
	HD/CAPD: dose after dialysis			
ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS	
Itraconazole	200 mg PO Q8-24 hours (histo/o	-	load with 200 mg Q8 hours	
	x2 days, then 200 mg Q12 hours	5)		

	Notes: avoid PPIs/H2 blockers; suspension on empty stomach; capsules with meal or acidic drink; consider monitoring SS trough after 5-7 days (>1 mg/dL, sum of itraconzole and hydroxy-itraconazole)				
Ketoconazole	200-400 mg PO Q24 hours  Note: Dose and frequency vary by indications. Please see appropriate drug information reference.				
Levofloxacin IV or PO	Pneumonia/Pseudomonas:		20-49:	Pneumonia /Pseudomonas: 750mg Q48 hours	
	750 mg Q24 hours		10-19:	Other: 500 mg load then 250 mg Q24h Pneumonia /Pseudomonas: 500 mg Q48 hours	
	Other Indications: 500 mg Q24 hours		<10:	Other: 500 mg load then 250 mg Q48h Pneumonia /Pseudomonas: 250-500 mg Q48h Other: 250 mg Q48 hours	
			HD:	Pneumonia /Pseudomonas: 250-500 mg Q48h Other: 250 mg Q48 hours	
Linezolid IV or PO	600 mg Q12 hours		No ad	iustment needed	
Meropenem	500 mg IV Q6 hours		25-49:	500 mg IV Q8 hours (UTI: 500 mg Q12 hours; meningitis, etc: 2 gm Q 12 hours)	
	UTI: 500 mg Q8 hours	:	10-24:	500 mg IV Q12 hours (UTI: 250 mg Q12 hours; meningitis, etc: 1 gm Q12 hours)	
	Meningitis, CF, meropenem MIC of 4 mg/dL: 2 gm Q8 hours	<1	.0/HD:	500 mg IV Q24 hours (UTI: 500 mg Q24 hours; meningitis, etc: 1 gm Q24 hours)	
		HD,	/CAPD	dose as CrCl <10, given after dialysis on dialysis days	
ANTI-INFECTIVE	NORMAL DOSE	_	CL in /min	RENAL DOSE ADJUSTMENTS	
Metronidazole IV or PO	500 mg Q12 hours if C difficile is not suspected	<10:		Consider 50% at same interval if >14 day duration	

	500 mg Q8 hours if C difficile is suspected	HD/CAPD	Give after dialysis on dialysis days
Micafungin IV	100 mg IV Q24 hours	No adjustn	nent needed
Minocycline <b>PO</b>	100 mg PO Q12 hours	<10:	100 mg Q24 hours
		HD:	100 mg Q24 hours
Nafcillin	Continuous Infusion: 8-12 g/da Note: 12 g/day recommended for endocarditis  Intermittent Dosing: 2 g IV Q4-6 hours	or	justment needed
Nitrofurantoin	100 mg PO Q12 hours	<60, HD/CAPD:	<50, HD/CAPD: Use is not recommended - will not
		1127 67 (1 2 .	reliably reach useful
			concentrations in urine and
			will have increased risk of
			toxicity.
Oseltamivir	Treatment:	<30:	Prophylaxis: 75 mg PO Q48
	75 mg PO Q12 hours		hours
	Prophylaxis: 75 mg PO Q24 hours	HD:	Treatment/Prophylaxis: 30 mg PO x 1 before HD then after every other dialysis session x 3 doses
Penicillin G IV	Continuous Infusion:	****	Continuous Infusion
	18-24 million units/day	10-50:	12-18 million units/day (75% IV at same time interval)
		<10	6-12 million units/day (75% IV at same time interval)
		HD:	Dose as CrCl <10, given after dialysis on dialysis days
		CAPD:	Dose a CrCl <10.
	Intermittent Dosing: 2-3 million units IV Q4-6 hours	****	Intermittent Dosing
		10-50:	1-2 million units/day IV Q4- 6 hours

	Note: Doses varies by indication. Please refer to appropriate drug reference.	<10:	hours
ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS
Penicillin VK <b>PO</b>	500 mg PO Q6 hours Note: Doses varies by indication. Please refer to appropriate drug reference.	10-50: <10: HD:	500 mg PO Q8 hours 500 mg PO Q12 hours Give dose after dialysis on dialysis days
Piperacillin/ Tazobactam	Pneumonia/Pseudomonas: 4.5 g IV Q6 hours  Other Indications: 3.375 g IV Q6 hours or 4.5 g IV Q8 hours	20-39: <20: HD:	IV Q6 hours Other: 2.25 g IV Q6 hours Pna/Pseudomonas: 2.25 g IV Q6 hours Other: 2.25 g IV Q8 hours
Quinipristin/ Dalfopristin	7.5 mg/kg IV Q8-12 hours Note due to cost and to maintain sus	=	
Rifampin	Doses vary by indication. Please reference.  Note: Should not be used alone;		
Rimantadine	100mg PO Q12 hours	<30:	100 mg PO Q24 hours
Terbinafine	500 mg PO Q12 hours 250 mg PO Q24 hours	<50:	Use not recommended
ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS

Note: Consultation with Pharmacy recommended.	Extended Interval Dosing:  7 mg/kg IV Q24 hours Note: draw level 6-14 hours after 1st dose  Conventional Dosing:  1-2.5 mg/kg/dose Q8 hours Note: Peak & trough levels monitored, consult pharmacy	*****  40-59: 30-39: <30  *****  40-60: 20-39: 10-20: <10: HD:	7 mg/kg IV Q48 hours AVOID, use conventional dosing Conventional Dosing (empiric, before levels): 1-2.5 mg/kg Q12 hours 1-2.5 mg/kg Q24 hours 1-2.5 mg/kg Q48 hours 1-2.5 mg/kg Q72 hours
Trimethoprim/ Sulfamethoxazole <b>PO</b> (1 Bactrim DS tablet = 160mg(TMP)/800(SMX)  Note: When changing from IV to PO use the same trimethoprim dose as IV	MRSA cellulitis (or Skin/skin structure infection/other infections):  1-2 DS tablets Q12 hours  Other Indications: 1 DS tablet Q12 hours	15-30: <15: HD:	hours or Decrease dose by 50% (ie. DS to SS)  AVOID or decrease dose 50% (ie. DS to SS) AND increase interval to Q48 hours
ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS
Trimethoprim/ Sulfamethoxazole <b>IV</b>	Pneumocystis treatment: 15-20 mg/kg/day in 3-4	15-30	50% of usual dose in 2-4 divided doses

Note: Dosing is based	divided doses	<15	AVOID if possible
on Trimethoprim component	Severe Infections: 8-10 mg/kg/day in 2-4 divided doses		Pneumocystis treatment: 15-20 mg/kg/dose Q48 hours or 7-10 mg/kg/day divided in 1-2 doses Severe Infections:
		HD:	8-10 mg/kg/dose Q48 hous or 4-5 mg/kg/day divided in 1- 2 doses Dose according to dosage for <15, schedule dose to be given after dialysis
Vancomycin	Loading Dose: 20 mg/kg IV x 1	40-60: 25-39:	15-20 mg/kg IV Q24 hours 15-20 mg/kg IV Q48 hours
	Maintenance:	<24:	Dosing adjustments per levels
Note: Consultation with Pharmacy recommended.	15-20 mg/kg IV Q8-12 hours	HD:	Dosing adjustments per levels
	Max Dose 2000 mg Round to nearest 250 mg Note: trough levels should be monitored on patient with expected duration > 3 days		

\*Dosing, therapeutic goals, and monitoring should be individualized for each patient.

Troughs of 15-20 mcg/mL are recommended for patients with MRSA bloodstream infections, endocarditis, meningitis, pneumonia, osteomyelitis, and septic arthritis.

ANTI-INFECTIVE	NORMAL DOSE	CRCL in ml/min	RENAL DOSE ADJUSTMENTS
Valacyclovir	250-500 mg PO Q8 hours		

	Note: Doso and from unner		Notal Bonal dosa
	Note: Dose and frequency		Note: Renal dose
	vary by indications. Please		adjustments vary by
	see appropriate drug		indication, please refer to
	information reference.		appropriate drug reference
	2g PO q12h	30-49:	1g PO q12h
		10-29:	1g PO q12h
		<10:	
	1g PO q8h	30-49:	
		10-29:	1g PO q24h
		<10:	500mg PO q24h
	1g PO q12h	30-49:	no adjustment
		10-29:	1g PO q24h
		<10:	500mg PO q24h
	1g PO q24h	30-49:	no adjustment
		10-29:	500mg PO q24h
		<10:	500mg PO q24h
	500mg PO q12h	30-49:	no adjustment
		10-29:	500mg PO q24h
		<10:	500mg PO q24h
	500mg PO q24h	30-49:	no adjustment
		10-29:	500mg PO q48h
		<10:	500mg PO q48h
		HD:	500mg PO q48h
		CAPD:	•
Voriconazole <b>PO</b>	Loading Dose: 400 mg PO x 1, th	l .	500mg PO q48h
Voriconazole PO	May require dose adjustment in	_	
		•	•
	Note: Consider weight base dos		patients using ADJ BW
	Note: Oral formulation is 95% b		
	Note: Screen for drug interactio	ns, dose may	r need adjustment
Mariana I. Dr	A althought a second		
Voriconazole <b>IV</b>	Active disease:		
	Loading dose of 6mg/kg q12h		
	x2 doses, then 4mg/kg 12h		
		1	ınction: no adjustment
	Prophylaxis:	necessary	
	200mg q12h (100mg q12h if		
	<40kg)	·	, accumulation of IV vehicle
		-	(cyclodextrin) – Switch to PO
	Therapeutic drug monitoring	or DC	
	is suggested. Voriconazole		

target trough at steady-state is 2 - 5.5 mg/L.	

Reference: Micromedex, Sanford Guide to Antimicrobial Therapy (42th edition)

## **Reportable Diseases**

REPORTABLE DISEASES IN KANSAS for health care providers, hospitals, and laboratories (K.S.A. 65-118, 65-128, 65-6001 - 65-6007, K.A.R. 28-1-2, 28-1-4, and 28-1-18. Changes effective as of 4/28/2006)

**■** - Indicates that a telephone report is required by law within four hours of *suspect* or *confirmed* cases to KDHE toll-free at 877-427-7317

① - Indicates that an isolates must be sent to: Division of Health and Environmental Laboratories

Forbes Field, Building #740, Topeka, KS 66620-0001

Phone: (785) 296-1633

Acquired Immune Deficiency Syndrome (AIDS)

Amebiasis

Anthrax 22

Arboviral disease (including West Nile virus, Western Equine encephalitis (WEE) and St. Louis encephalitis (SLE))

- indicate virus whenever possible

Botulism 🕾

Brucellosis

Campylobacter infections

Chancroid

Chlamydia trachomatis genital infection

Cholera 🕾

Cryptosporidiosis

Cyclospora infection

Diphtheria

Ehrlichiosis

Escherichia coli O157:H7 (and other shiga-toxin producing E. coli, also known as STEC) ①

Giardiasis

Gonorrhea

Haemophilus influenza, invasive disease

Hantavirus Pulmonary Syndrome

Hemolytic uremic syndrome, postdiarrheal

Hepatitis, viral (acute and chronic)

Hepatitis B during pregnancy

Human Immunodeficiency Virus (HIV) (includes Viral Load Tests)

Influenza deaths in children <18 years of age Legionellosis Leprosy (Hansen disease) Listeriosis Lyme disease Malaria Measles (rubeola) 🕾 Meningitis, bacterial 🕾 Meningococcemia ① 🕾 Mumps 🕾 *Pertussis* (whooping cough) 🕾 **Plague** (Yersinia pestis) Poliomyelitis 🕾 Psittacosis *Q Fever* (Coxiella burnetii) 🕾 Rabies, human and animal 🕾 Rocky Mountain Spotted Fever Rubella, including congenital rubella syndrome 🕾 Salmonellosis, including typhoid fever ① Severe Acute Respiratory Syndrome (SARS) ① 🕾 Shigellosis (1) Smallpox 22 Streptococcal invasive, drug-resistant disease from Group A Streptococcus or Streptococcus pneumoniae ①

Syphilis, including congenital syphilis

Tetanus

Toxic shock syndrome, streptococcal and staphylococcal

Transmissible Spongioform Encephalopathy (TSE) or prion disease (includes CJD)

Trichinosis

Tuberculosis, active disease ① 22

Tuberculosis, latent infection

Tularemia

Varicella (chickenpox)

Viral hemorrhagic fever 🕾

Yellow fever

## In addition, laboratories must report:

- Viral load results of reportable diseases
- ALL blood lead levels, as of 12/2002 (KCLPPP/ABLES)
- $\bullet$  CD4+ T-lymphocyte count < 500/  $\mu l$  or CD4+ T-lymphocytes  $<\!\!29\%$  of total lymphocytes

Outbreaks, unusual occurrence of any disease, exotic or newly recognized diseases, and suspect acts of terrorism should be reported within 4 hours by telephone to the Epidemiology Hotline: 877-427-7317

Mail or fax reports to your local health department and/or to:

KDHE Office of Surveillance and Epidemiology, 1000 SW Jackson, Suite 210, Topeka, KS 66612-1274 Fax: 877-427-7318 (toll-free)