

مجموعه جداول انتخاب شده از CLSI M100 33th ed., 2023 برای میکروارگانیسم های اولویت دار در برنامه کشوری مهار مقاومت میکروبی بر اساس راهنمای سازمان جهانی بهداشت (GLASS)

ويرايش هفتم سال ۱۴۰۲

کمیته تخصصی میکروب شناسی آزمایشگاه مرجع سلامت وزارت بهداشت، درمان و آموزش پزشکی



بسمه تعالى

این سند با هدف استفاده آزمایشگاههای بیمارستانهای **منتخب** در برنامه کشوری مهار مقاومت میکروبی تهیه شده است ولی در سایر آزمایشگاههای پزشکی غیر منتخب، با در نظر گرفتن راهنماهای مربوط به مرکز درمانی/ بیمارستان برای انجام آزمایش و گزارشدهی آزمایش تعیین حساسیت ضدمیکروبی، نیز می تواند استفاده شود.

آزمایشگاههای بیمارستانهای <mark>منتخب</mark> برای آزمایش و گزارشدهی میکروارگانیسمهای بیماریزای اولویتدار باید مطابق جدول زیر عمل نمایند.

Target pathogens	Specimens			
Acinetobacter spp.				
Escherichia coli				
Streptococcus pneumoniae				
Salmonella spp.	Blood			
Staphylococcus aureus				
Klebsiella pneumoniae				
Enterococcus spp.				
Pseudomonas aeruginosa				
Escherichia coli				
Klebsiella pneumoniae	L Inin e			
Enterococcus spp.	Urine			
Pseudomonas aeruginosa				
Salmonella spp.	Stool			
Shigella spp.	51001			

GLASS target pathogens and specimen types



ویرایش هفتم این سند جایگزین ویرایش قبلی (ویرایش ششم – سال ۱۴۰۱) می باشد. تغییرات عمده در ویرایش هفتم این سند در جدول زیر فهرست شده است. تغییرات کوچک یا ویراستاری و توضیحات، با حروف پررنگ نوشته شده است.

Overview of Changes

Table	Changes
Escherichia coli	Added:
	 Levofloxacin disk diffusion breakpoints (page 6)
	Revised:
	 Gentamicin, and amikacin disk diffusion breakpoints (page 6)
Klebsiella pneumonia	Added:
	 Levofloxacin disk diffusion breakpoints (page 10)
	Revised:
	 Gentamicin, and amikacin disk diffusion breakpoints (page 9)
Salmonella spp.	Added:
	 Levofloxacin MIC breakpoints (page 12)
	 Imipenem, meropenem and tetracycline disk diffusion
	breakpoints (page 12)
Shigella spp.	Added:
	 Levofloxacin, imipenem, meropenem and tetracycline disk
	diffusion breakpoints (pages 13-14)
Tests for Extended-	Added:
Spectrum β-Lactamases in	• Note (page 15)
Escherichia coli, Klebsiella	
pneumonia, Salmonella spp.	
and Shigella spp.	
Pseudomonas aeruginosa	Revised:
	• Piperacillin-tazobactam and tobramycin disk diffusion
	breakpoints (pages 17-18)
	Urine designation for amikacin (page 18)
	Gentamicin disk diffusion breakpoints
Acinetobacter spp.	Added:
	• Comment (d) (page 19)
Staphylococcus aureus	
	Levofloxacin disk diffusion breakpoints (page 23)
Enterococcus spp.	
	 Levotloxacin disk dittusion breakpoints (page 25)



Note: Intermediate ranges denoted with a "^" for the applicable antimicrobial agents in the drug groups are based on the known ability of these agents to concentrate in the urine.

Escherichia coli					
Antimicrobial Agent	Disk I Content	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm			Comments
		S	Ι	R	
PENICILLINS				<u> </u>	
Ampicillin	10 μg	≥17	14–16^	≤ 13	 (a) Results of ampicillin testing can be used to predict results for amoxicillin. (b) Breakpoints are based on an ampicillin dosage regimen of 2 g parenterally administered every 4–6 h or an amoxicillin dosage regimen of 1-2 g parenterally administered every 6 h. (c) Breakpoints when oral ampicillin is used only for therapy of salmonellosis, shigellosis, or uncomplicated UTIs due to <i>E. coli</i> and <i>P. mirabilis</i> are based on an amoxicillin dosage regimen of 500 mg orally administered every 6 h or an amoxicillin dosage regimen of 250 mg orally administered every 8 h or 500 mg every 12 h.
CEPHEMS					
Cefazolin (PARENTERAL)	30 µg	≥23	20–22	≤ 19	Breakpoints when cefazolin is used for therapy of infections other than uncomplicated UTIs due to <i>E. coli</i> , <i>K.</i> <i>pneumoniae</i> & <i>P. mirabilis</i> . Breakpoints are based on a dosage regimen of 2 g administered every 8 h.
Cefazolin (PARENTERAL) (urine)	30 µg	≥ 15	-	≤ 14	 (a) Report only on organisms isolated from the urinary tract. (b) Breakpoints when cefazolin is used for therapy of uncomplicated UTIs due to <i>E. coli</i>, <i>K. pneumoniae & P. mirabilis</i>. Breakpoints are based on a dosage regimen of 1 g administered every 12 h.



Escherichia coli (continued)								
Cefazolin (ORAL) (surrogate test for oral cephalosporins & uncomplicated UTIs) (urine)	30 µg	≥15	_	≤ 14	 (a) Report only on organisms isolated from the urinary tract. (b) Breakpoints are for cefazolin when used as a surrogate test to predict results for the oral agent cefaclor, cefdinir, cefpodoxime, cefprozil, cefuroxime, cephalexin, and loracarbef when used for therapy of uncomplicated UTIs due to <i>E. coli, K. pneumoniae</i>, and <i>P. mirabilis</i>. Cefazolin as a surrogate may overcall resistance to cefdinir, cefpodoxime, and cefuroxime. If cefazolin tests resistant, test these drugs individually if needed for therapy. 			
Cefepime	30 µg	≥ 25	19–24	≤ 18	The breakpoint for susceptible is based on a dosage regimen of 1 g administered every 12 h. The Breakpoint for SDD* is based on dosing regimens that result in higher cefepime exposure, either higher doses or more frequent doses or both, up to approved maximum dosing regimens. *SDD: Susceptible-Dose Dependent			
Cefotaxime <u>or</u>	30 µg	≥26	23–25^	≤ 22	Breakpoints are based on a dosage regimen of 1 g administered every 24 h for ceftriaxone and 1 g administered every 8 h for cefotaxime.			
ceftriaxone	30 µg	≥23	20–22^	≤19				
Ceftazidime	30 µg	≥21	18–20^	≤17	Breakpoints are based on a dosage regimen of 1 g administered every 8 h.			
CARBAPENEMS								
Imipenem	10 µg	≥23	20–22^	≤ 19	Breakpoints are based on a dosage regimen of 500 mg administered every 6 h or 1 g every 8 h.			
Meropenem	10 µg	≥23	20–22*	≤19	Breakpoints are based on a dosage regimen of 1 g administered every 8 h.			



Escherichia coli (continued) LIPOPEPTIDES WARNING: Clinical and PK/PD data demonstrate colistin and polymyxin B have limited clinical efficacy, even if an intermediate result is obtained. Alternative agents are strongly preferred. Colistin and polymyxin B should be used in combination with one or more active antimicrobial agents. Consultation with an infectious diseases specialist is recommended. Colistin (a) Colistin (methanesulfonate) should be given with a loading dose and maximum or polymixin B renally adjusted dose. (b) Polymixin B should be given with a loading dose and maximum recommended dose. (c) When colistin or polymixin B is given systemically, neither is likely to be effective for pneumonia. (d) For colistin, broth microdilution, CBDE*, and CAT** MIC methods are acceptable. For polymixin B, broth microdilution is the only approved method. Disk diffusion and gradient diffusion methods should not be performed (see Table 3D, Pages 174-179). *CBDE: Colistin Broth Disk Elution **CAT: Colistin Agar Test **Interpretive Categories and MIC** Breakpoints, µg/mL S Ι R $\geq \overline{4}$ < 2 -AMINOGLYCOSIDES 10 µg Gentamicin **≥18** 15-17^ **≤**14 Breakpoints are based on a dosage regimen of 7 mg/kg parenterally administered every 24 h. Breakpoints are based on a dosage regimen Amikacin 30 µg > 20 17-19^ < 16 of 15 mg/kg parenterally administered every 24 h.

FLUOROQUINOLONES					
Ciprofloxacin <u>or</u>	5 µg	≥26	22-25^	≤21	(a) Breakpoints for ciprofloxacin are based on a dosage regimen of 400 mg IV or 500 mg orally administered every 12 h.
levofloxacin	5 µg	≥21	17-20^	≤16	(b) Breakpoints for levofloxacin are based on a dosage regimen of 750 mg administered every 24 h.



Escherichia coli (continued)							
FOLATE PATHWAY INHIBITORS							
Trimethoprim- sulfamethoxazole	1.25/23.75	≥16	11-15	≤ 10			
	μg						
NITROFURANS							
Nitrofurantoin	300 µg	≥17	15–16	≤14	Report only on organisms isolated from the		
(urine)					urinary tract.		



Klebsiella pneumoniae						
Antimicrobial Agent	Disk Content	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm			Comments	
CEPHEMS		~				
Cefazolin (PARENTERAL)	30 µg	≥ 23	20–22	≤ 19	Breakpoints when cefazolin is used for therapy of infections other than uncomplicated UTIs due to <i>E. coli</i> , <i>K.</i> <i>pneumoniae</i> & <i>P.mirabilis</i> . Breakpoints are based on a dosage regimen of 2 g administered every 8 h.	
Cefazolin (PARENTERAL) (urine)	30 µg	≥ 15	-	≤ 14	 (a) Report only on organisms isolated from the urinary tract. (b) Breakpoints when cefazolin is used for therapy of uncomplicated UTIs due to <i>E. coli</i>, <i>K. pneumoniae & P. mirabilis</i>. Breakpoints are based on a dosage regimen of 1 g administered every 12 h. 	
Cefazolin (ORAL) (surrogate test for oral cephalosporins & uncomplicated UTI) (urine)	30 µg	≥ 15	_	<i>≤</i> 14	 (a) Report only on organisms isolated from the urinary tract. (b) Breakpoints are for cefazolin when cefazolin results are used to predict results for the oral agents cefaclor, cefdinir, cefpodoxime, cefprozil, cefuroxime, cephalexin, and loracarbef when used for therapy of uncomplicated UTIs due to <i>E. coli, K. pneumoniae,</i> and <i>P. mirabilis.</i> Cefazolin as a surrogate may overcall resistance to cefdinir, cefpodoxime, and cefuroxime. If cefazolin tests resistant, test these drugs individually if needed for therapy. 	
Cefepime	30 µg	≥ 25	19–24	≤ 18	The Breakpoint for susceptible is based on a dosage regimen of 1 g every 12 h. The Breakpoint for SDD* is based on dosing regimens that result in higher cefepime exposure, either higher doses or more frequent doses or both, up to approved maximum dosing regimens. *SDD: Susceptible-Dose Dependent	
Cefotaxime or	30 µg	≥26	23–25^	≤22	Breakpoints are based on a dosage regimen of 1 g administered every 24 h for ceftriaxone and 1 g administered every 8 h	
ceftriaxone	30 µg	≥23	20-22^	≤19		



Klebsiella pneumonia	(continu	ed)			
Ceftazidime	30 µg	≥21	18–20^	≤17	Breakpoints are based on a dosage regimen of 1 g administered every 8 h.
CARBAPENEMS					
Imipenem	10 µg	≥23	20-22^	≤19	Breakpoints are based on a dosage regimen of 500 mg administered every 6 h or 1 g every 8 h.
Meropenem	10 µg	≥23	20–22^	≤19	Breakpoints are based on a dosage regimen of 1 g administered every 8 h.
LIPOPEPTIDES					
WARNING: Clinical and PK/P if an intermediate result is obta be used in combination with or specialist is recommended.	PD data demo ined. Alterna ne or more ac	nstrate tive age ctive ant	colistin and nts are stro timicrobial	d polyn ongly p l agents	nyxin B have limited clinical efficacy, even referred. Colistin and polymyxin B should s. Consultation with an infectious diseases
Colistin <u>or</u> polymixin B					(a) Colistin (methanesulfonate) should be given with a loading dose and maximum renally adjusted dose.(b) Polymixin B should be given with a loading dose and maximum recommended doses.(c) When colistin or polymixin B is given systemically, neither is likely to be effective for pneumonia.(d) For colistin, broth microdilution, CBDE*, and CAT** MIC methods are acceptable. For polymixin B, broth microdilution is the only approved method. Disk diffusion and gradient diffusion methods should not be performed (see Table 3D, Pages 174-179).*CBDE: Colistin Broth Disk Elution **CAT: Colistin Agar TestInterpretive Categories and MIC Breakpoints, µg/mLSIR- ≤ 2 ≥ 4
AMINOGLYCOSIDES					
Gentamicin	10 µg	≥18	15-17^	≤14	Breakpoints are based on a dosage regimen of 7 mg/kg parenterally administered every 24 h.
Amikacin	30 µg	≥ 20	17–19^	<u>≤16</u>	Breakpoints are based on a dosage regimen of 15 mg/kg parenterally administered every 24 h.



Klebsiella pneumonia (continued)							
FLUOROQUINOLONES							
Ciprofloxacin	5 µg	≥26	22-25^	≤21	(a) Breakpoints for ciprofloxacin are based on a dosage regimen of 400 mg IV or 500 mg orally administered every 12 h.		
levofloxacin	5 µg	≥21	17-20^	≤16	(b) Breakpoints for levofloxacin are based on a dosage regimen of 750 mg administered every 24 h.		
FOLATE PATHWAY INHIBITORS							
Trimethoprim-	1.25/ 23.75	≥16	11–15	≤ 10			
sulfamethoxazole	μg						
NITROFURANS							
Nitrofurantoin	300 µg	≥ 17	15-16	≤ 14	Report only on organisms isolated from		
(urine)					the urinary tract.		



* When fecal isolates of *Salmonella* spp. are tested, only ampicillin, a fluoroquinolone, and trimethoprimsulfamethoxazole should be reported routinely. For extraintestinal isolates of *Salmonella* spp., a thirdgeneration cephalosporin should be tested and reported, and chloramphenicol may be tested and reported if requested. Susceptibility testing is indicated for typhoidal *Salmonella* (*S. enterica* ser. Typhi and *S. enterica* ser. Paratyphi A–C) isolated from extraintestinal and intestinal sources. Routine susceptibility testing is not indicated for nontyphoidal *Salmonella* spp. isolated from intestinal sources

Salmonella spp.							
Antimicrobial Agent	Disk Content	Interp and B near	Interpretive Categories and Zone Diameter Breakpoints,		Comments		
		S	Ι	R			
PENICILLINS							
Ampicillin	10 μg	≥17		≤ 13	 (a) Results of ampicillin testing can be used to predict results for amoxicillin. (b) Breakpoints are based on an ampicillin dosage regimen of 2 g parenterally administered every 4–6 h. (c) Breakpoints when oral ampicillin is used only for therapy of salmonellosis, shigellosis, or uncomplicated UTIs due to <i>E. coli</i> and <i>P. mirabilis</i> are based on an ampicillin dosage regimen of 500 mg orally administered every 6 h or an amoxicillin dosage regimen of 250 mg orally administered every 8 h or 500 mg every 12 h. 		
CEPHEMS							
Ceftriaxone	30 µg	$\geq \overline{23}$	_	≤ <u>19</u>	(a) Breakpoints are based on a dosage regimen of 1 g administered every 24 h for ceftriaxone		
ceftazidime (For extraintestinal isolates)	30 µg	≥21	_	≤17	(b) Breakpoints are based on a dosage regimen of 1 g administered every 8 h.		



FLUOROQUINOLONESThe preferred test for assessing fluroquinolone susceptibility or resistance in Salmonella spp. is a ciprofloxacin MIC test. A levofloxacin or ofloxacin MIC test can be performed if either agent, respectively, is the fluoroquinolone of choice in a specific facility.Ciprofloxacin $5 \ \mu g$ ≥ 31 $ \leq 20$ Isolates of Salmonella spp. that test not susceptible to ciprofloxacin, levofloxacin, offloxacin, or pefloxacin may be associated with clinical failure or delayed response in fluoroquinolone-treated patients with salmonellosis.or $ -$ levofloxacin $ -$ reordination $ -$ levofloxacin $ -$ levofloxacin $ -$ levofloxacin $ -$ reordination $ -$ reordination <td< th=""></td<>
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$ \begin{array}{ c c c c } Ciprofloxacin & 5 \ \mu g & \geq 31 \\ or & 5 \ \mu g & \geq 31 \\ or & - & - & - \\ \hline \\ evofloxacin & - & - & - \\ \hline \\ evofloxacin & - & - & - \\ \hline \\ evofloxacin & - & - & - \\ \hline \\ evofloxacin & - & - & - \\ \hline \\ \hline \\ evofloxacin & - & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - & - & - & - & - & - \\ \hline \\ FOLATE PATHWAY INHIBITORS & - & - & - & - & - & - & - & - & - & $
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levofloxacinsalmonellosis.Interpretive Categories and MIC Breakpoints, µg/mLBreakpoints, µg/mLImage: Book state s
levofloxacin- - - - -Interpretive Categories and MIC Breakpoints, $\mu g/\mu L$ For LevofloxacinIterpretive PATHWAY INHIBITORS- FOLATE PATHWAY INHIBITORS- Trimethoprim- sulfamethoxazole μg 1.25/23.75 μg ≥ 16 11-15 ≤ 10 ≤ 10 PHENICOLSChloramphenicol Azithromycin30 μg ≥ 18 13-17 ≤ 12 ≤ 12 (a) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily.CARBAPENEMS ≤ 12 (a) S. enterica request if antimicrobial agents in other tiers are not ontimal because of various factors*
levofloxacin- - - - -Breakpoints, $\mu g/mL$ For LevofloxacinBIRSIR ≤ 0.12 - ≥ 2 FOLATE PATHWAY INHIBITORSTrimethoprim- sulfamethoxazole $1.25/23.75$ ≥ 16 $11-15$ ≤ 10 $ \geq 2$ PHENICOLSChloramphenicol $30 \ \mu g$ ≥ 18 $13-17$ ≤ 12 $ -$ Azithromycin $15 \ \mu g$ ≥ 13 $ \leq 12$ (a) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of $500 \ m g$ administered daily.CARBAPENEMSCARBAPENEMSCARBAPENEMSThese antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not ontimal because of various factors*
levofloxacin $ -$ <
SIR ≤ 0.12 $ \geq 2$ FOLATE PATHWAY INHIBITORSTrimethoprim- sulfamethoxazole $1.25/23.75$ ≥ 16 $11-15$ ≤ 10 $ \geq 2$ PHENICOLSChloramphenicol $30 \ \mu g$ ≥ 18 $13-17$ ≤ 12 $ -$ MACROLIDSAzithromycin $15 \ \mu g$ ≥ 13 $ \leq 12$ (a) <i>S. enterica</i> ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily.CARBAPENEMSThese antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not ontimal because of various factors*
FOLATE PATHWAY INHIBITORS ≤ 0.12 $ \geq 2$ Trimethoprim- sulfamethoxazole $1.25/23.75$ ≥ 16 $11-15$ ≤ 10 $ \geq 2$ PHENICOLS μ g 1 $1-17$ ≤ 10 $ -$ Chloramphenicol 30μ g ≥ 18 $13-17$ ≤ 12 $ -$ <t< td=""></t<>
FOLATE PATHWAY INHIBITORSTrimethoprim- sulfamethoxazole $1.25/23.75$ ≥ 16 $11-15$ ≤ 10 pHENICOLSChloramphenicol $30 \ \mu g$ ≥ 18 $13-17$ ≤ 12 MACROLIDSAzithromycin $15 \ \mu g$ ≥ 13 $ \leq 12$ (a) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily.CARBAPENEMSThese antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not ontimal because of various factors*
$\begin{array}{c c c c c c c c c c c c c c c c c c c $
sulfamethoxazoleµgImage: constraint of the second s
PHENICOLSChloramphenicol $30 \ \mu g$ ≥ 18 $13-17$ ≤ 12 MACROLIDSAzithromycin $15 \ \mu g$ ≥ 13 $ \leq 12$ (a) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily.CARBAPENEMSThese antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not ontimal because of various factors*
$\begin{array}{c c c c c c c c c c c c c c c c c c c $
MACROLIDSAzithromycin $15 \ \mu g$ ≥ 13 $ \leq 12$ (a) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily.CARBAPENEMSThese antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not ontimal because of various factors*
Azithromycin $15 \ \mu g$ ≥ 13 $ \leq 12$ (a) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily.CARBAPENEMSThese antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not optimal because of various factors*
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These antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in other tiers are not optimal because of various factors*
- OTHER TIERS ARE NOT ODDITINAL DECAUSE OF VALIOUS LACIOUS
$\frac{10 \text{ ug}}{23} > 23 \qquad \qquad \leq 10 \text{ Breakpoints are based on a decage}$
$\frac{10 \ \mu g}{10 \ \mu g} = \frac{2.5}{2.5} = \frac{5.17}{5} \frac{\text{Dreakpoints are based on a dosage}}{\frac{10 \ \mu g}{10 \ \mu g}}$
h or 1 g every 8 h
Meronenem $10 \mu \sigma > 23 - <19$ Rreaknoints are based on a dosage
regimen of 1 g administered every 8 h.
TETRACYCLINES
These antimicrobial agents may warrant testing and reporting by clinical request if antimicrobial agents in
other tiers are not optimal because of various factors*
Tetracycline $30 \ \mu g \ge 15 \ 12-14 \ \le 11 \ 0 \ rganisms$ that are susceptible to
Tetracycline $30 \ \mu g \ge 15$ $12-14 \le 11$ Organisms that are susceptible to tetracycline are also considered
Tetracycline $30 \ \mu g$ ≥ 15 $12-14$ ≤ 11 Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and
Tetracycline $30 \ \mu g$ ≥ 15 $12-14$ ≤ 11 Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and minocycline. However, some organisms
Tetracycline $30 \ \mu g$ ≥ 15 $12-14$ ≤ 11 Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and minocycline. However, some organisms that are intermediate or resistant to
Tetracycline $30 \ \mu g$ ≥ 15 $12-14$ ≤ 11 Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and minocycline. However, some organisms that are intermediate or resistant to tetracycline may be susceptible to
Tetracycline $30 \ \mu g$ ≥ 15 $12-14$ ≤ 11 Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and minocycline. However, some organisms that are intermediate or resistant to tetracycline may be susceptible to doxycycline, minocycline, or both.

سالمونلاً و شَيْكَلا مي باشند.



*When fecal isolates of *Shigella* spp. are tested, only ampicillin, a fluoroquinolone, and trimethoprimsulfamethoxazole should be reported routinely. **Data regarding whether amoxicillin should be used to treat shigellosis are conflicting. When reporting ampicillin results, state that treatment of shigellosis with amoxicillin might have poorer efficacy compared with treatment with ampicillin. Susceptibility testing is indicated for all** *Shigella* **isolates.**

Shigella spp.					
Antimicrobial Agent	Disk Content	Interp and B near	retive Cate Zone Dian Breakpoints rest whole	egories ieter s, mm	Comments
		S	I	R	
PENICILLINS			<u> </u>		
Ampicillin	10 μg	≥ 17	_	≤ 13	 (a) Results of ampicillin testing can be used to predict results for amoxicillin. (b) Breakpoints are based on an ampicillin dosage regimen of 2 g parenterally administered every 4-6 h. (c) Breakpoints when oral ampicillin is used only for therapy of salmonellosis, shigellosis, or uncomplicated UTIs due to <i>E. coli</i> and <i>P. mirabilis</i> are based on an ampicillin dosage regimen of 500 mg orally administered every 6 h or an amoxicillin dosage regimen of 250 mg orally administered every 8 h or 500 mg every 12 h.
CEPHEMS					· · · · · ·
Ceftriaxone <u>or</u>	30 µg	≥23	_	≤19	Breakpoints are based on a dosage regimen of 1 g administered every 24 h for ceftriaxone
ceftazidime	30 µg	≥21	_	≤17	Breakpoints are based on a dosage regimen of 1 g administered every 8 h.
(Only for ciprofloxacin					
FLUOROOUINOLONES	<u> </u>				
Ciprofloxacin or	5 µg	≥26	-	≤21	Breakpoints for ciprofloxacin are based on a dosage regimen of 400 mg IV or 500 mg orally administered every 12 h.
levofloxacin	5 µg	≥21	_	≤16	Breakpoints for levofloxacin are based on a dosage regimen of 750 mg administered every 24 h.



Shigella spp. (conti	nued)									
FOLATE PATHWAY INHIBITORS										
Trimethoprim-	1.25/23.75	≥16	11–15	≤ 10						
sulfamethoxazole	μg									
MACROLIDES										
Azithromycin	15 μg	≥ 16	11-15	≤ 10	 (a) <i>Shigella</i> spp. only: azithromycin disk diffusion zones can be hazy and difficult to measure, especially <i>S. sonnei</i>. If an isolate has a zone of inhibition that is difficult to measure, an MIC method is recommended. Media source may affect the clarity of the end points for disk diffusion tests. (b) Breakpoints are based on a dosage regimen of 500 mg administered daily. 					
CARBAPENEMS										
These antimicrobial agents	may warran	t testing	and repor	ting by	clinical request if antimicrobial agents in					
other tiers are not optimal	because of va	rious fac	ctors*							
Imipenem	10 µg	≥23	_	≤19	Breakpoints are based on a dosage regimen of 500 mg administered every 6 h or 1 g every 8 h.					
Meropenem	10 µg	≥23	-	≤19	Breakpoints are based on a dosage regimen of 1 g administered every 8 h.					
TETRACYCLINES										
These antimicrobial agents	may warrant	t testing	and repor	ting by o	clinical request if antimicrobial agents in					
other tiers are not optimal	because of va	rious fac	ctors*	1						
Tetracycline	30 µg	≥15	12–14	≤11	Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and minocycline. However, some organisms that are intermediate or resistant to tetracycline may be susceptible to doxycycline minocycline or both					

سالمونلا و شيكلا مي بأشند.



Tests for Extended-Spectrum β-Lactamases in *Escherichia coli*, *Klebsiella pneumonia*, *Salmonella* spp and *Shigella* spp.

NOTE: Following evaluation of PK/PD properties, limited clinical data, and MIC distributions, revised breakpoints for cefazolin, cefotaxime, ceftazidime, ceftizoxime, ceftriaxone, and aztreonam were published in January 2010 (M100-S20) and are listed in Table 2A. Cefuroxime (parenteral) was also evaluated; however, no change in breakpoints was necessary with the dosage. When using the current breakpoints, routine ESBL testing is not necessary before reporting results. If ESBL testing is performed, the results may be used to guide therapeutic management or for epidemiological or infection prevention purposes.

Some phenotypic ESBL tests have known limitations that affect sensitivity (eg, false-negative results due to the coproduction of an AmpC β -lactamase) and specificity (eg, false-positive results due to hyperproduction of non-ESBL β -lactamases combined with altered permeability). Genotypic methods are limited by the targets included in the assay (eg, most FDA-cleared ESBL assays target only blaCTX-M). Limitations of phenotypic and genotypic methods must be considered.

Breakpoints for drugs with limited availability in many countries (eg, moxalactam, cefonicid, cefamandole, and cefoperazone) were not evaluated. If considering use of these drugs for *E. coli*, *Klebsiella pneumoniae*, *Klebsiella oxytoca*, or *Proteus mirabilis*, ESBL testing should be performed. If isolates test ESBL positive, the results for moxalactam, cefonicid, cefamandole, and cefoperazone should be reported as resistant.



neumonia, Salmonella spectrum p-Lactamases in Escherichia coli, Kledslella pneumonia, Salmonella spp and Shigella spp. (continued)									
Test	Criteria for Performance of ESBL Test	ESBL Test							
Antimicrobial concentration	Cefpodoxime 10 µg or Ceftazidime 30 µg or Aztreonam 30 µg or Cefotaxime 30 µg or Ceftriaxone 30 µg (Testing more than one antimicrobial agent improves the sensitivity of ESBL detection.)	Ceftazidime 30 µg Ceftazidime-clavulanate 30/10 µg and Cefotaxime 30 µg Cefotaxime-clavulanate 30/10 µg (Testing necessitates using both cefotaxime and ceftazidime, alone and in combination with clavulanate.)							
Results	Cefpodoxime zone $\leq 17 \text{ mm}$ Ceftazidime zone $\leq 22 \text{ mm}$ Aztreonam zone $\leq 27 \text{ mm}$ Cefotaxime zone $\leq 27 \text{ mm}$ Ceftriaxone zone $\leq 25 \text{ mm}$ Zones above may indicate ESBL production.	$A \ge 5mm$ increase in a zone diameter for either antimicrobial agent tested in combination with clavulanate vs the zone diameter of the agent when tested alone = ESBL (eg, ceftazidime zone = 16; ceftazidime-clavulanate zone = 21).							
Reporting		For all confirmed ESBL-producing strains: If laboratories do not use current cephalosporin and aztreonam breakpoints, the test interpretation should be reported as resistant for all penicillins, cephalosporins, and aztreonam. If laboratories use current cephalosporin and aztreonam breakpoints, then test interpretations for these agents do not need to be changed from susceptible to resistant.							



Pseudomonas aerus	ginosa								
Antimicrobial Agent	Disk Content	Interpretive Categoriesand Zone DiameterBreakpoints,nearest whole mmSIR			Comments				
β-LACTAM/β-LACTAMA	SE INHIBIT	OR CO	MBINATIO	ONS	I				
Piperacillin-tazobactam	100/10 μg	≥22	18–21^ ≤ 17		Breakpoint dosage reg over 30 m intermedia prevent sr from ca interpretat	ts for susce imen of 4.5 g inutes or ov te are only to nall uncont using maj ions.	eptible are based on a g administered every 6 h er 3 h. Breakpoints for o provide a buffer zone to rolled technical factors for discrepancies in		
CEPHEMS									
Cefepime	30 µg	≥18	15-17^	≤14	Breakpoints administtere 12 h.	s are based o ed every 8 h	n a dosage regimen of 1 g or 2 g administtered every		
Ceftazidime	30 µg	≥ 18 15-17 [^] ≤ 14		Breakpoints are based on a dosage regimen of 1 g administered every 6 h or 2 g administered every 8 h.					
LIPOPEPTID					•				
WARNING: Clinical and P intermediate result is obtain in combination with one or recommended.	K/PD data de ned. Alternat more active	emonstra ive agen antimic	ate colistin its are stroi crobial agei	and pol ngly pro nts. Cor	lymyxin B ha eferred. Coli nsultation wi	ive limited c stin and poly th an infect	linical efficacy, even if an ymyxin B should be used ious diseases specialist is		
Colistin <u>or</u> polymixin B	-	-	-	-	 (a) Colistin with a loadidoses. (b) Polymiz dose and mathematical constraints of the system cally pneumonia. (c) When system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic CAT** Mathematical constraints of the system cally pneumonia. (d) For colistic Canadita constraints of the system cally pneumonia. (d) For colistic Canadita constraints of the system cally pneumonia. (e) Attractic Canadita constraints of the system cally pneumonia. (f) Attractic Canadita constraints of the system cally pneumonia. (g) Attractic Canadita constraints of the system constraints of the system cally pneumonia. (g) Attractic Canadita constraints of the system constrai	i (methanesung dose and ing dose and ing dose and aximum reconstruction of the construction of the constr	Ifonate) should be given maximum renally adjusted be given with a loading mmended doses. polymixin B is given likely to be effective for dicrodilution, CBDE*, and is are acceptable. For dicrodilution is the only diffusion and gradient ld not be performed (see 9). Disk Elution est s and MIC Breakpoints, /mL R		



Pseudomonas aeruginosa (continued)											
CARBAPENEMS											
Imipenem	10 µg	≥19	16-18^	≤15	Breakpoints for imipenem are based on a dosage regimen of 1 g administered every 8 h or 500 mg administered every 6 h.						
Meropenem	10 µg	≥19	16-18^	≤15	Breakpoints for meropenem are based on a dosage regimen of 1 g administered every 8 h.						
AMINOGLYCOSIDES											
Tobramycin	10 µg	≥19	13- 18^	≤ 12	 (a) Breakpoints are based on a dosage regimen of 7 mg/kg parenterally administered every 24 h. (b) Tobramycin does not predict susceptibility to gentamicin. 						
Amikacin (Urine)	30 µg	≥17	15–16^	≤ 14	 (a) Report only on organisms isolated from the urinary tract. (b) Breakpoints are based on a dosage regimen of 15 mg/kg parenterally administered every 24 h. 						
FLUOROQUINOLONES		-									
Ciprofloxacin	5 µg	≥25	19-24^	≤18	Breakpoints are based on a dosage regimen of 400 mg IV administered every 8 h.						



Acinetobacter spp.								
Antimicrobial Agent	Disk Content	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm				Comm	ents	
		S	I	R				
R-I ACTAM/R-I ACTAMA	SF INHIRIT	OR COI	MRINATI	ONS				
Ampicillin-sulbactam	10/10 µg	> 15	12-14	< 11				
	10,10 µB	_ 10						
Piperacillin-tazobactam	100/10 µg	≥21	18–20	≤17				
CEPHEMS		•						
Cefepime	30 µg	≥18	15-17	≤14				
Ceftazidime	30 µg	≥18	15-17	≤14				
CARBAPENEMS			l		L			
Imipenem	10 µg	$\geq 22 \qquad 19-21 \qquad \leq 18$			Breakpoints 500 mg adn	s are based or ninistered eve	a dosage regimen of ery 6 h.	
Meropenem	10 µg	$\geq 18 \qquad 15-17 \qquad \leq 14$			Breakpoints are based on a dosage regimen of 1 g administered every 8 h or 500 mg administered every 6 h.			
LIPOPEPTID								
WARNING: Clinical and P if an intermediate result is o be used in combination wit specialist is recommended.	K/PD data do obtained. Alte h one or mor	emonstra ernative re active	ate colistin agents are antimicrol	and pol strongly bial age	lymyxin B ha y preferred. nts. Consult	ave limited c Colistin and ation with a	linical efficacy, even polymyxin B should n infectious diseases	
Colistin	-	-	-	-	(a) Colistin (methanesulfonate) should be			
$\frac{\text{or}}{1}$					given with	a loading	dose and maximum	
polymixin B					renally adju	isted doses.	d he siven with a	
					loading do	se and maxi	in de given with a	
					doses.	se and max	initiani recommended	
					(c) When	colistin or p	olymixin B is given	
					systemically	y, the drug	is unlikely to be	
					effective fo	r pneumonia.	2	
					(d) The o	nly approve	d MIC methods is	
					broth micr	odilution. C	BDE*, CAT**, disk	
					diffusion, a	ind gradient	diffusion should not	
					be perform	<u>ied.</u>		
					*CBDE: Co	listin Broth I	Disk Elution	
					Interr	retive Cate	contest and MIC	
					Inter	Breaknoint	$s_{\rm ug/mI}$	
					S	I	R	
						<u>≤</u> 2	≥ <u>4</u>	



Acinetobacter spp. (continued)									
AMINOGLYCOSIDES									
Gentamicin	10 µg	≥15	13-14	≤ 12					
Tobramycin	10 µg	≥15	13-14	≤ 12					
Amikacin	30 µg	≥17	15–16	≤14					
TETRACYCLINES									
Minocycline	30 µg	≥16	13–15	≤12					
FLUOROQUINOLONES									
Ciprofloxacin	5 µg	≥21	16–20	≤15					
FOLATE PATHWAY INH	FOLATE PATHWAY INHIBITORS								
Trimethoprim-	1.25/ 23.75	≥16	11–15	≤10					
sulfamethoxazole	μg								



Staphylococcus aureus									
Antimicrobial Agent	Disk Content	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm			Comments				
		S	Ι	R					
PENICILLINASE-LABIL	E PENICILL	INS							
Penicillin	10 units	≥29	-	≤ 28	(a) Penicillin should be used to test the susceptibility of all staphylococci to all penicillinase- labile penicillins. Penicillin- resistant strains of staphylococci produce β -lactamase. Perform test(s) to detect β -lactamase production on staphylococci for which the penicillin MICs are ≤ 0.12 µg/mL or zone diameters ≥ 29 mm before reporting the isolate as penicillin susceptible. Rare isolates of staphylococci that contain geness for β -lactamase production may appear negative by β -lactamase tests. Consequently, for serious infections requiring penicillin therapy, laboratories should perform MIC tests and β -lactamase testing on all subsequent isolates from the same patient. PCR testing of the isolate for the <i>blaZ</i> β -lactamase gene may be considered. See Table 3F, Pages 186-187 . (b) For methicillin (oxacillin)- resistant staphylococci report penicillin as resistant or do not report.				



Staphylococcus a	Staphylococcus aureus (continued)										
PENICILLINASE-STA	BLE PENIC	ILLINS									
Oxacillin (Oxacillin disk testing is not reliable for <i>S. aureus</i> and <i>S. lugdunensis.</i>)	30 µg Cefoxitin (surrogate test for oxacillin)	≥ 22 (cefoxitin)	-	≤21 (cefoxitin)	 (a) Cefoxitin is tested as a surrogate for oxacillin for some species of <i>Staphylococcus</i>. Isolates that test resistant by cefoxitin or oxacillin, when using the appropriate test method for the species, should be reported as methicillin (oxacillin) resistant. If testing only cefoxitin, report as methicillin (oxacillin) susceptible or resistant based on the cefoxitin result. Isolates that test either <i>mecA</i> negative or PBP2a negative or cefoxitin susceptible should be reported as methicillin (oxacillin) susceptible. (b) For isolates of <i>S. aureus</i> that do not grow well on CAMHB* 						
					do not grow well on CAMHB* or unsupplemented MHA (eg, small-colony variants), testing on other media (eg, BMHA) does not reliably detect <i>mecA</i> - mediated resistance. Testing for PBP2a using induced growth (ie, growth taken from the zone margin surrounding a cefoxitin disk on either BMHA or a blood agar plate after 24 hours incubation in 5% CO ₂) or <i>mecA</i> should be done. *Cation Adgusted Mueller Hinton Agar						



Staphylococcus aur	reus (cont	tinued	l)				
GLYCOPEPTIDES							
GLYCOPEPTIDES Vancomycin	-	-	-	-	(a) For <i>S</i> susceptible vancomycin course of pr (b) MIC te to determin isolates vancomycin differentiate susceptible from <i>N</i> isolates, differentiate susceptible, resistant iso spp. other which give inhibition. (c) Send an vancomycin reference la	S aureus, w isolates m n intermediat rolonged then sts should be the suscept of staphyl n. The disk t e w isolates of vancomycin-in nor does e among , -intermedi olates of <i>Stap</i> than <i>S</i> . aureus than <i>S</i> . aureus fon is ≥ 8 µ aboratory. we Categorie akpoints, µg	vancomycin- aay become te during the rapy. e performed tibility of all ococci to est does not vancomycin- <i>S. aureus</i> intermediate the test vancomycin ate, and - <i>phylococcus</i> <i>ureus</i> all of te zones of or which the tg/mL to a
Teicoplanin (Optional)					<u> </u>	4-0	≤ 10
(Investigation)	-	-	-	-	Bre	akpoints, µg	g/mL
					S		R
					≤ 8	16	≥ 32
TETRACYCLINES	•		•				
Doxycycline	30 µg	≥16	13-15	≤ 12			
MACROLIDES	I						
Erythromycin	15 μg	≥23	14-22	≤13	Not routine isolated fro	ly reported o m the urinary	n organisms y tract.
FLUOROQUINOLONES	ſ	I			1		
Ciprofloxacin <u>or</u> levofloxacin	5 μg 5 μg	≥ 21 ≥ 19	16–20 16-18	≤15 ≤15	Staphylocod resistance c with quinol that are in become res days after	ccus spp. m luring prolon lones. Theref nitially susce istant within initiation	ay develop aged therapy ore, isolates eptible may three to four of therapy.
					varranted.	repeat isola	ttes may be



Staphylococcus aureus (continued)									
NITROFURANTOINS									
Nitrofurantoin	300 µg	≥17	15-16	≤14	Report only on organisms isolated from the urinary tract.				
FOLATE PATHWAY INH	IBITORS	•	•	•					
Trimethoprim-	1.25/23.75	≥16	11-15	≤ 10					
sulfamethoxazole	μg								
LINCOSAMIDES									
Clindamycin	2 μg	≥21	15-20	≤ 14	 (a) Not routinely reported on organisms isolated from the urinary tract. (b) For isolates that test erythromycin resistant and clindamycin susceptible or intermediate, testing for ICR* by disk diffusion using the D-zone test or by broth microdilution is required befor reporting clindamycin (See Table 3I, Pages 196-198). (c) D-zone test: 15-µg erythromycin and 2-µg clindamycin disks spaced 15–26 mm apart. Report isolates with ICR as "clindamycin resistant" (See Table 3I, Pages 196-198). *ICR: Inducible clindamycin resistance 				
ANSAMYCINS		1							
Rifampin	5 µg	≥ 20	17-19	≤16	<i>Rx</i> : should not be used alone for antimicrobial therapy.				



Enterococcus spp.					
Antimicrobial Agent	Disk Content	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm		egories neter s, mm R	Comments
		5	1	N	
PENICILLINS					
Ampicillin	10 μg	≥17	-	≤ 16	The results of ampicillin susceptibility tests should be used to predict the activity of amoxicillin. Ampicillin results may be used to predict susceptibility to amoxicillin- clavulanate, ampicillin-sulbactam, and piperacillin-tazobactam among non– β -lactamase producing enterococci. Ampicillin susceptibility can be used to predict imipenem susceptibility, providing the species is confirmed to be <i>E. faecalis</i> .
GLYCOPEPTIDES			1	•	
Vancomycin	30 µg	≥ 17	15-16	≤ 14	When testing vancomycin against enterococci, plates should be held a full 24 hours for accurate detection of resistance. Zones should be examined using transmitted light; the presence of a haze or any growth within the zone of inhibition indicates resistance. Organisms with intermediate zones should be tested by an MIC method as described in M07. For isolates for which the vancomycin MICs are 8 to 16 µg/mL, perform biochemical tests for identification as listed under the "Vancomycin MIC \geq 8 µg/mL" test found in Table 3H, Pages 194-195 .
FLUOROQUINOLONES			16.004	. 1 5	
Ciprofloxacin (Urine)	5 μg	≥ 21	16-20^	≤15	Report only on organisms isolated from the urinary tract.
levofloxacin (Urine)	5 µg	≥ 21	14-16^	≤ 13	
NITROFURANTOINS Nitrofurantoin (Urine)	300 µg	≥17	15-16	≤14	Report only on organisms isolated from the urinary tract.
OXAZOLIDINONES					-
Linezolid	30 µg	≥23	21-22	≤ 20	



Test for Gentamicin High-Level Aminoglycoside Resistance in							
Enterococcus spp.	1	1					
Antimicrobial Agent	Disk	Interp	retive Categori	ies and	Comments		
	Content		Zone Diameter	•			
			Breakpoints,				
		n	earest whole m	m			
		S	Inconclusive	R			
Gentamicin	120 µg	≥ 10	7-9	= 6	If disk diffusion result is		
					inconclusive: perform an agar		
					dilution or broth dilution MIC		
					test to confirm (See Table 3K,		
					Pages 202-204).		



* For disk diffusion, test a maximum of 9 disks on a 150-mm plate and 4 disks on a 100-mm plate.

Streptococcus pneumoniae										
Antimicrobial Agent	Disk Content	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm		egories leter S, mm	Comments					
		S	Ι	R						
Penicillin and cefotaxime, ceftriaxone, or meropenem should be tested by a reliable MIC method (such as that described in M071) and reported routinely with <i>S. pneumoniae</i> isolated from CSF. Such isolates can also be tested against vancomycin using the MIC or disk diffusion method. With isolates from other sites, the oxacillin disk test may be used. If the oxacillin zone size is \leq 19 mm, cefotaxime, ceftriaxone, meropenem, or penicillin MICs should be determined.										
PENICILLINS	1	> 20		[
(nonmeningitis)	I μg Oxacillin	≥ 20	-	-	Isolates of pneumococci with oxacillin zone sizes of ≥ 20 mm are susceptible (MIC $\leq 0.06 \mu g/mL$) to penicillin. Penicillin and cefotaxime, ceftriaxone, or meropenem MICs should be determined for those isolates with oxacillin zone diameters of \leq 19 mm, because zones of \leq 19 mm occur with penicillin-resistant, -intermediate, or certain -susceptible strains. For isolates with oxacillin zones \leq 19 mm, do not report penicillin as resistant without performing a penicillin MIC test.					
Penicillin parenteral	-	-	-	-	Interpretive Categories and MIC					
(nonmeningitis)					Breakpoints, μg/mL					
(optional)					S I R					
					(a) Rx : Doses of intravenous penicillin of at least 2 million units every 4 hours in adults with normal renal function (12 million units per day) can be used to treat nonmeningeal pneumococcal infections due to strains with penicillin MICs $\leq 2 \ \mu g/mL$. Strains with an intermediate MIC of 4 $\ \mu g/mL$ may require penicillin doses of 18 to 24 million units per day. (b) For all isolates other than those from CSF, report interpretations for both meningitis and nonmeningitis.					



Streptococcus pneumoniae (continued)									
CEPHEMS									
Ceftriaxone (nonmeningitis)	-	-	-	-	Interpretive Categories and MIC Breakpoints, µg/mL				
_					S I R				
					≤ 1 2 ≥ 4				
					For all isolates other than those from CSF, report interpretations for both meningitis and nonmeningitis.				
TETRACYCLINES									
Doxycycline	30 µg	≥28	25-27	≤24	Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline. However, resistance to doxycycline cannot be inferred from tetracycline resistance.				
MACROLIDES									
Erythromycin	15 μg	≥21	16-20	≤15	(a) Susceptibility and resistance to azithromycin, clarithromycin, and dirithromycin can be predicted by testing erythromycin.				
					(b) Not routinely reported on organisms isolated from the urinary tract.				
FLUOROQUINOLONES									
Levofloxacin	5 µg	≥17	14-16	≤13					
FOLATE PATHWAY INI	HIBITORS		1						
Trimethoprim-	1.25/23.75	≥ 19	16-18	≤ 15					
sulfamethoxazole	μg								
Clindamycin	2 µg	> 10	16.18	< 15	(a) Not routinely reported on organisms				
	2 #5	_ 17			 (a) For formerly reported on organisms isolated from the urinary tract. (b) For isolates that test erythromycin resistant and clindamycin susceptible or intermediate, testing for ICR* by disk diffusion using the D-zone test or by broth microdilution is required befor reporting clindamycin (See Table 3I, Pages 196-198). (c) D-zone test: 15-μg erythromycin and 2-μg clindamycin disks spaced 15–26 mm apart. Report isolates with ICR as "clindamycin resistant" (See Table 3I, Pages 196-198). *ICR: Inducible clindamycin resistance 				