

M100

Performance Standards for Antimicrobial Susceptibility Testing

This document includes updated tables for the Clinical and Laboratory Standards Institute antimicrobial susceptibility testing standards M02, M07, and M11.

A CLSI supplement for global application.

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M100, 30th ed. January 2020 Replaces M100, 29th ed.

Performance Standards for Antimicrobial Susceptibility Testing

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Abstract

The data in the tables are valid only if the methodologies in CLSI documents M02,1 M07,2 and M113 are followed. These standards contain information about disk diffusion (M021) and dilution (M072 and M113) test procedures for aerobic and anaerobic bacteria. Clinicians depend heavily on information from the microbiology laboratory for treating their seriously ill patients. The clinical importance of antimicrobial susceptibility test results demands that these tests be performed under optimal conditions and that laboratories have the capability to provide results for the newest antimicrobial agents. The tables presented in M100 represent the most current information for drug selection, interpretation, and quality control using the procedures standardized in M02,1 M07,2 and M11.3 Users should replace previously published tables with these new tables. Changes in the tables since the previous edition appear in boldface type.

Clinical and Laboratory Standards Institute (CLSI). Performance Standards for Antimicrobial Susceptibility Testing. 30th ed. CLSI supplement M100 (ISBN 978-1-68440-066-9 [Print]; ISBN 978-1-68440-067-6 [Electronic]). Clinical and Laboratory Standards Institute, 950 West Valley Road, Suite 2500, Wayne, Pennsylvania 19087 USA, 2020.

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Suggested Citation

CLSI. *Performance Standards for Antimicrobial Susceptibility Testing*. 30th ed. CLSI supplement M100. Wayne, PA: Clinical and Laboratory Standards Institute; 2020.

Previous Editions:

December 1986, December 1987, December 1991, December 1992, December 1994, December 1995, January 1997, January 1998, January 1999, January 2000, January 2001, January 2002, January 2003, January 2004, January 2005, January 2006, January 2007, January 2008, January 2010, June 2010, January 2011, January 2012, January 2013, January 2014, January 2015, January 2016, January 2017, January 2018, January 2019

ISBN 978-1-68440-066-9 (Print) ISBN 978-1-68440-067-6 (Electronic) ISSN 1558-6502 (Print) ISSN 2162-2914 (Electronic)

Volume 40, Number 1

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Contents

Abstract	i
Committee Membership	iii
Overview of Changes	xiv
Summary of CLSI Processes for Establishing Breakpoints and Quality Control Ranges	.xxvi
CLSI Reference Methods vs Commercial Methods and CLSI vs US Food and Drug Administration Breakpoints	xxvii
CLSI Breakpoint Additions/Revisions Since 2010	xxix
CLSI Epidemiological Cutoff Value Additions/Revisions Since 2015	xxxi
CLSI Archived Resources.	xxxii
Subcommittee on Antimicrobial Susceptibility Testing Mission Statement	xxxii
Instructions for Use of Tables	1
References	16
Table 1A. Suggested Groupings of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That Should Be Considered for Testing and Reporting on Nonfastidious Organisms by Microbiology Laboratories in the United States	18
Table 1B. Suggested Groupings of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That Should Be Considered for Testing and Reporting on Fastidious Organisms by Microbiology Laboratories in the United States	24
Table 1C. Suggested Groupings of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That Should Be Considered for Testing and Reporting on Anaerobic Organisms by Microbiology Laboratories in the United States	30
Table 2A. Zone Diameter and MIC Breakpoints for Enterobacterales.	32
Table 2B-1. Zone Diameter and MIC Breakpoints for <i>Pseudomonas aeruginosa</i>	42

Table 2B-2. Zone Diameter and MIC Breakpoints for <i>Acinetobacter</i> spp.	46
Table 2B-3. Zone Diameter and MIC Breakpoints for <i>Burkholderia cepacia</i> complex	50
Table 2B-4. Zone Diameter and MIC Breakpoints for Stenotrophomonas maltophilia	52
Table 2B-5. MIC Breakpoints for Other Non-Enterobacterales	54
Table 2C. Zone Diameter and MIC Breakpoints for Staphylococcus spp.	58
Table 2D. Zone Diameter and MIC Breakpoints for <i>Enterococcus</i> spp.	68
Table 2E. Zone Diameter and MIC Breakpoints for Haemophilus influenzae and Haemophilus parainfluenzae	74
Table 2F. Zone Diameter and MIC Breakpoints for Neisseria gonorrhoeae	78
Table 2G. Zone Diameter and MIC Breakpoints for Streptococcus pneumoniae	82
Table 2H-1. Zone Diameter and MIC Breakpoints for <i>Streptococcus</i> spp. β-Hemolytic Group	88
Table 2H-2. Zone Diameter and MIC Breakpoints for <i>Streptococcus</i> spp. Viridans Group	92
Table 2I. Zone Diameter and MIC Breakpoints for Neisseria meningitidis	96
Table 2J. MIC Breakpoints for Anaerobes	100
Table 3A. Tests for Extended-Spectrum β-Lactamases in <i>Klebsiella pneumoniae, Klebsiella oxytoca, Escherichia coli</i> , and <i>Proteus mirabilis</i>	104
Introduction to Tables 3B and 3C. Tests for Carbapenemases in Enterobacterales and <i>Pseudomonas aeruginosa</i>	108
Table 3B. CarbaNP Test for Suspected Carbapenemase Production in Enterobacterales and <i>Pseudomonas aeruginosa</i>	110
Table 3B-1. Modifications of Table 3B When Using MIC Breakpoints for Carbapenems Described in M100-S20 (January 2010)	114

Table 3C. Modified Carbapenem Inactivation Methods for Suspected Carbapenemase Production in Enterobacterales and <i>Pseudomonas aeruginosa</i>	118
Table 3C-1. Modifications of Table 3C When Using MIC Breakpoints for Carbapenems Described in M100-S20 (January 2010)	130
Table 3D. Tests for Colistin Resistance for Enterobacterales and <i>Pseudomonas aeruginosa</i>	132
Table 3E. Test for Detection of β-Lactamase Production in <i>Staphylococcus</i> spp.	138
Table 3F. Test for Detecting Methicillin (Oxacillin) Resistance in Staphylococcus spp	142
Table 3G. Vancomycin Agar Screen for Staphylococcus aureus and Enterococcus spp.	146
Table 3H. Test for Detecting Inducible Clindamycin Resistance in <i>Staphylococcus</i> spp., <i>Streptococcus pneumoniae</i> , and <i>Streptococcus</i> spp. β-Hemolytic Group	148
Table 3I. Test for Detecting High-Level Mupirocin Resistance in Staphylococcus aureus	152
Table 3J. Test for Detecting High-Level Aminoglycoside Resistance in <i>Enterococcus</i> spp. (Includes Disk Diffusion)	154
Table 4A-1. Disk Diffusion QC Ranges for Nonfastidious Organisms and Antimicrobial Agents Excluding β-Lactam Combination Agents	156
Table 4A-2. Disk Diffusion QC Ranges for Nonfastidious Organisms and β-Lactam Combination Agents	160
Table 4B. Disk Diffusion QC Ranges for Fastidious Organisms	164
Table 4C. Disk Diffusion Reference Guide to QC Frequency	168
Table 4D. Disk Diffusion Troubleshooting Guide	170
Table 5A-1. MIC QC Ranges for Nonfastidious Organisms and Antimicrobial Agents Excluding β-Lactam Combination Agents	174
Table 5A-2. MIC QC Ranges for Nonfastidious Organisms and β-Lactam Combination Agents	180

Table 5B. MIC QC Ranges for Fastidious Organisms (Broth Dilution Methods)	184
Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method)	188
Table 5D. MIC QC Ranges for Anaerobes (Agar Dilution Method)	190
Table 5E. MIC QC Ranges for Anaerobes (Broth Microdilution Method)	192
Table 5F. MIC Reference Guide to QC Frequency	194
Table 5G. MIC Troubleshooting Guide	196
Table 6A. Solvents and Diluents for Preparing Stock Solutions of Antimicrobial Agents	200
Table 6B. Preparing Stock Solutions for Antimicrobial Agents Provided With Activity Expressed as Units	206
Table 6C. Preparing Solutions and Media Containing Combinations of Antimicrobial Agents	208
Table 7. Preparing Dilutions of Antimicrobial Agents to Be Used in Agar Dilution Susceptibility Tests	212
Table 8A. Preparing Dilutions of Antimicrobial Agents to Be Used in Broth Dilution Susceptibility Tests	214
Table 8B. Preparing Dilutions of Water-Insoluble Antimicrobial Agents to Be Used in Broth Dilution Susceptibility Tests	216
Appendix A. Suggestions for Confirming Antimicrobial Susceptibility Test Results and Organism Identification for Agents Approved by the US Food and Drug Administration for Clinical Use	218
Appendix B. Intrinsic Resistance	226
Appendix C. QC Strains for Antimicrobial Susceptibility Tests	234
Appendix D. Anaerobe Cumulative Antibiogram	240
Appendix E. Dosage Regimens Used to Establish Susceptible or Susceptible-Dose Dependent Breakpoints	246

Table of Contents

Appendix F. Susceptible-Dose Dependent Interpretive Category	250
Appendix G. Epidemiological Cutoff Values	254
Appendix H. Using Molecular Assays for Resistance Detection	260
Appendix I. Cefiderocol Broth Preparation and Reading Broth Microdilution Minimal Inhibitory Concentration End Points	274
Glossary I (Part 1). β-Lactams: Class and Subclass Designations and Generic Names	278
Glossary I (Part 2). Non–β-Lactams: Class and Subclass Designations and Generic Names	280
Glossary II. Antimicrobial Agent Abbreviation(s), Route(s) of Administration, and Drug Class	284
Glossary III. List of Identical Abbreviations Used for More Than One Antimicrobial Agent in US Diagnostic Products	290
The Quality Management System Approach	292
Related CLSI Reference Materials	293

M100, 30th ec

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Overview of Changes

M100, 30th ed. replaces the previous edition of the supplement, M100, 29th ed., published in 2019. The major changes in M100, 30th ed., are listed below. Other minor or editorial changes were made to the general formatting and to some of the table footnotes and comments. Changes to the tables since the previous edition appear in boldface type. The following are additions or changes unless otherwise noted as a "deletion."

Users of M100, 30th ed. should note recent and new formatting changes to Tables 2, including:

• Intermediate ranges denoted with a "^" for the applicable antimicrobial agents in the drug groups in Tables 2 are based on the known ability of these agents to concentrate in the urine; some agents may also have the potential to concentrate at other anatomical sites (ie, epithelial lining).

M100 is updated and reviewed annually as new data and new agents become available. Use of outdated documents is strongly discouraged.

Section/Table	Change(s)
General	
Throughout the document	Replaced:
	• "Coagulase-negative staphylococci (CoNS)" with "other <i>Staphylococcus</i> spp."
	The term "infection control" with "infection prevention"
	Clarified:
	Methicillin and oxacillin terminology for <i>Staphylococcus</i> spp.
	Updated:
	• Genera formerly included in the family <i>Enterobacteriaceae</i> reorganized to an order (Enterobacterales) containing seven families: <i>Budviciaceae</i> , <i>Enterobacteriaceae</i> , <i>Erwiniaceae</i> , <i>Hafniaceae</i> , <i>Morganellaceae</i> , <i>Pectobacteriaceae</i> , <i>Yersiniaceae</i> ⁴
	Nomenclature for Salmonella Typhi to Salmonella enterica ser. Typhi
	Nomenclature for Salmonella Paratyphi to Salmonella enterica ser. Paratyphi

Overview	of Changes	(Continued)
Over view	or Changes	(Continucu)

Section/Table	Change(s)
General (Continued)	
CLSI Breakpoint	Added:
Additions/Revisions Since 2010	• Cefiderocol disk diffusion breakpoints for Enterobacterales (p. xxix), <i>Pseudomonas aeruginosa</i> (p. xxx), <i>Acinetobacter</i> spp. (p. xxx), and <i>Stenotrophomonas maltophilia</i> (p. xxx)
	• Colistin (p. xxix) and polymyxin B (p. xxx) minimal inhibitory concentration (MIC) breakpoints for Enterobacterales
	• Daptomycin MIC breakpoints for <i>Enterococcus faecium</i> only (originally included in the March 2019 re-released version of M100, 29th ed.) (p. xxxi)
	Revised:
	• Colistin and polymyxin B MIC breakpoints for <i>P. aeruginosa</i> and <i>Acinetobacter</i> spp. (p. xxx)
	• Daptomycin MIC breakpoints for <i>Enterococcus</i> spp. other than <i>E. faecium</i> (originally included in the March 2019
	re-released version of M100, 29th ed.) (p. xxxi)
	Reinstated:
	Norfloxacin breakpoints deleted from M100, 29th ed. (pp. xxix–xxxi)
CLSI Epidemiological Cutoff	Deleted:
Value Additions/Revisions Since 2015	Colistin epidemiological cutoff value (ECV) (Enterobacterales; now assigned a breakpoint in Table 2A)
CLSI Archived Resources	Added:
	• Link to the archived table for QC ranges eliminated from M100 since 2010 (p. xxxii)
	• Link to the archived table for ECVs eliminated from M100 since 2010 (p. xxxii)
Instructions for Use of Table	S
II. Breakpoint and	Revised:
Interpretive Category	Breakpoint examples for the nonsusceptible interpretive category (p. 4)
Definitions	Susceptible-dose dependent (SDD) category definition (p. 4)
	Intermediate category definition (p. 5)

Section/Table	Change(s)
Instructions for Use of Tables	s (Continued)
VIII. Routine, Supplemental,	Supplemental Tests (Optional) table
Screening, Surrogate Agent,	Added:
and Equivalent Agent Testing	Colistin agar test (p. 11)
to Determine Susceptibility	Colistin broth disk elution (p. 11)
and Resistance to	Supplemental Tests (Required and Optional) tables
Antimicrobial Agents	Revised:
	• References to appropriate Tables 3 (pp. 10–11)
	Screening Tests table
	Revised:
	References to appropriate Tables 3 (p. 12)
	Surrogate Agent Tests table
	Clarified:
	• Cefoxitin test description for specific <i>Staphylococcus</i> spp. (p. 12)
	Revised:
	Reference to appropriate Table 3 (p. 12) Francisco of Francisco (p. 12)
	Examples of Equivalent Agent Tests table Added:
	Colistin and polymyxin B for Enterobacterales, <i>P. aeruginosa</i> , and <i>Acinetobacter baumannii</i> complex (p. 13)
X. Abbreviations and	Added:
Acronyms	• CAT (colistin agar test) (p. 14)
Tier ony ms	CBDE (colistin broth disk elution) (p. 14)
	ICR (inducible clindamycin resistance) (p. 14)
	MH-F agar (Mueller-Hinton fastidious agar) (p. 14)
	Revised:
	MRS (methicillin [oxacillin]-resistant staphylococci) (p. 15)
	MRSA (methicillin [oxacillin]-resistant Staphylococcus aureus) (p. 15)
	 NAD (β-nicotinamide adenine dinucleotide) (p. 15)
	Deleted:
	CoNS (coagulase-negative staphylococci)
	KPC (Klebsiella pneumoniae carbapenemase)
	 NDM (New Delhi metallo-β-lactamase)

Overview	of Changes	(Continued)
		(

Section/Table	Change(s)
	gs of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That
	esting and Reporting by Microbiology Laboratories in the United States
All Tables 1	Reformatted:
	Tables to clarify criteria for inclusion in each group
	Replaced:
	Test/Report Group column and descriptions for Groups A, B, C, and U with expanded descriptions
	(as listed in the Instructions for Use of Tables, Section IC)
Table 1A. Nonfastidious	Added:
Organisms	• Footnote for Group B directing users to the Instructions for Use of Tables for examples of when a Group B agent
	might be reported (p. 18)
	Clarified for Staphylococcus spp.:
	Oxacillin footnote regarding testing methods (p. 18)
	Relocated from Group B to Group A (in the same box as trimethoprim-sulfamethoxazole) for S. maltophilia:
	• Levofloxacin (p. 19)
	Minocycline (p. 19)
Table 1B. Fastidious	Added:
Organisms	• Footnote for Group B directing users to the Instructions for Use of Tables for examples of when a Group B agent
	might be reported (p. 24)
	Clarified for Streptococcus spp.:
	Footnote regarding inducible clindamycin resistance (ICR) reporting (p. 24)
Tables 2. Zone Diameter and	
General	Added:
	• Reference for the M02 Disk Diffusion Reading Guide to appropriate tables
Table 2A. Enterobacterales	Added:
	• Salmonella enterica ser. Typhi routine QC strain recommendations for azithromycin (p. 32)
	• General comment explaining the use of the "^," with intermediate breakpoints for appropriate antimicrobial agents
	(p. 32)
	• Cefiderocol testing requirements (p. 32), reference (p. 32), and investigational disk diffusion breakpoints (p. 36)
	• Colistin and polymyxin B MIC breakpoints, warning, reporting comments, and reference (p. 38)
	• I^ designation for β-lactams (p. 33), aminoglycosides (p. 38), and fluoroquinolones (pp. 39–40)
	Clarified:
	• Ceftazidime-avibactam reporting comment (p. 33)
	• Cefazolin (surrogate test for oral cephalosporins and uncomplicated urinary tract infections) reporting comment
	(p. 36)
	Reinstated: Norfloxacin disk diffusion and MIC breakpoints and reporting comment deleted from M100, 29th ed. (p. 39)
	Normonachi disk diffusion and ivite ofeakpoints and reporting comment defeded from ivitou, 29th ed. (p. 39)

Section/Table	Change(s)
Tables 2. (Continued)	8 (/
Table 2B-1. Pseudomonas	Added:
aeruginosa	• General comment explaining the use of the "^," with intermediate breakpoints for appropriate antimicrobial agents
	(p. 42)
	• Cefiderocol testing requirements (p. 42), reference (p. 42), and investigational disk diffusion breakpoints (p. 43)
	• I^ designation for β-lactams (p. 43), aminoglycosides (p. 45), and fluoroquinolones (p. 45)
	Revised:
	Colistin and polymyxin B MIC breakpoints, warning, reporting comments, and reference (p. 44)
	Reinstated:
	Norfloxacin disk diffusion and MIC breakpoints and reporting comment deleted from M100, 29th ed. (p. 45)
Table 2B-2. Acinetobacter spp.	Added:
	• Cefiderocol testing requirements (p. 46), and reference (p. 46), and investigational disk diffusion breakpoints (p. 47)
	Revised:
	Colistin and polymyxin B MIC breakpoints, warning, reporting comments, and reference (p. 48)
Table 2B-4. Stenotrophomonas	Added:
maltophilia	• Cefiderocol testing requirements (p. 52), reference (p. 52), and investigational disk diffusion breakpoints (p. 53)
	Revised:
T. I. 4D 7 0 I	Test/report group for minocycline and levofloxacin from B to A (p. 53)
Table 2B-5. Other	Clarified:
Non-Enterobacterales	• General comment regarding the species designated as non-Enterobacterales (p. 54)
	Reinstated:
Table 2C Combalance	Norfloxacin MIC breakpoints and reporting comment deleted from M100, 29th ed. (p. 55) Added:
Table 2C. Staphylococcus spp.	 Recommendation for selecting QC strains for routine QC of β-lactam combination agents (p. 58)
	Clarified:
	 Oxacillin reporting for other Staphylococcus spp. with MICs 0.5–2 μg/mL (p. 62)
	 Oxacimin reporting for other staphytococcus spp. with wires 0.3–2 μg/miz (p. 02) ICR reporting comment (p. 65)
	Revised:
	Methods for Detection of Methicillin (Oxacillin)-Resistant <i>Staphylococcus</i> spp. table in general
	comment (5) to include incubation times for detecting methicillin (oxacillin) resistance (p. 59)
	Reinstated:
	Norfloxacin disk diffusion and MIC breakpoints and reporting comment deleted from M100, 29th ed. (p. 64)

Overview	of Changes	(Continued)
Over view	or Changes	(Continucu)

Section/Table	Change(s)
Tables 2. (Continued)	3 1,
Table 2D. Enterococcus spp.	 Added: Recommendation for selecting QC strains for routine QC of β-lactam combination agents (p. 68) General comment explaining the use of the "^," with intermediate breakpoints for appropriate antimicrobial agents (p. 68) Daptomycin MIC breakpoints (SDD and resistant only) and dosage regimen for <i>E. faecium</i> only (originally included in the March 2019 re-released version of M100, 29th ed.) (p. 70) Daptomycin intermediate MIC breakpoint and dosage regimen for <i>Enterococcus</i> spp. other than <i>E. faecium</i> (originally included in the March 2019 re-released version of M100, 29th ed.) (p. 70) I^ designation for fluoroquinolones and oxazolidinones (p. 71) Revised: Daptomycin susceptible MIC breakpoint for <i>Enterococcus</i> spp. other than <i>E. faecium</i> (originally included in the March 2019 re-released version of M100, 29th ed.) (p. 70) Reinstated:
	Norfloxacin disk diffusion and MIC breakpoints and reporting comment deleted from M100, 29th ed. (p. 71)
Table 2G. Streptococcus pneumoniae	 Added: Mueller-Hinton fastidious agar (MH-F agar) as an alternative for disk diffusion testing (p. 82) Reporting comment for oral cefuroxime (p. 84) Clarified: ICR reporting comment (p. 86)
Table 2H-1. <i>Streptococcus</i> spp. β-Hemolytic Group	Clarified: Erythromycin reporting comment (p. 90) ICR reporting comment (p. 91)
	nce Testing (NOTE: Tables following 3C were renumbered to accommodate addition of the new Table 3D.)
Table 3A. Tests for Extended- Spectrum β-Lactamases in Klebsiella pneumoniae, Klebsiella oxytoca, Escherichia coli, and Proteus mirabilis	 Revised: Aztreonam disk diffusion QC range for Klebsiella pneumoniae ATCC® 700603 for the extended-spectrum β-lactamase (ESBL) test (p. 106)
Table 3B. CarbaNP Test for Suspected Carbapenemase	Added:
Production in Enterobacterales and Pseudomonas aeruginosa	 New references (p. 110) Revised: NOTE 1 regarding ability to detect OXA-48-like producers (p. 113)

Overview of Changes (Co	
Section/Table	Change(s)
Tables 3. (Continued)	
Table 3C. Modified	Added:
Carbapenem Inactivation	New references (p. 118)
Methods for Suspected	
Carbapenemase Production in	
Enterobacterales and	
Pseudomonas aeruginosa	
Table 3D. Tests for	Added:
Colistin Resistance for	• Colistin broth disk elution (CBDE) procedure (pp. 132–134), QC recommendations (p. 134), and associated figures
Enterobacterales and	(p. 136)
Pseudomonas aeruginosa	• Colistin agar test (CAT) procedure (pp. 132–134), QC recommendations (p. 134), and associated figures (p. 137)
(new table)	
Table 3F. Detecting	Added:
Methicillin (Oxacillin)	• Options and respective procedures for detecting mecA-mediated resistance using cefoxitin or oxacillin with
Resistance in Staphylococcus	disk diffusion, broth microdilution, or agar dilution methods (pp. 142–143)
spp. (formerly Table 3E)	• QC strains recommended for when various methods are used for detecting methicillin (oxacillin) resistance (p. 144)
Table 3H. Test for Detecting	Clarified:
Inducible Clindamycin	Organism groups to be tested (pp. 148–149)
Resistance in Staphylococcus	• ICR reporting comments (p. 149)
spp., Streptococcus	
pneumoniae, and Streptococcus	
spp. β-Hemolytic Group	
(formerly Table 3G)	

Section/Table	Change(s)					
Tables 4. Disk Diffusion QC I	Ranges and Associated Tables		3 ()			
Table 4A-1. Disk Diffusion	Added:					
QC Ranges for Nonfastidious	• Sulopenem disk diffusion QC ranges for <i>Escherichia coli</i> ATCC® 25922 (p. 158)					
Organisms and Antimicrobial	Tedizolid disk diffusion QC ranges	for <i>E. faecalis</i> A	TCC® 29212 as a sup	plemental QC str	ain (p. 158)	
Agents Excluding β-Lactam	Revised:					
Combination Agents	Ciprofloxacin disk diffusion QC range for E. coli ATCC® 25922 (p. 157)					
	• Tedizolid disk content and QC ranges for S. aureus ATCC® 25923 (p. 158)					
	Reinstated:					
	Norfloxacin QC ranges for all QC st	 Norfloxacin QC ranges for all QC strains deleted from M100, 29th ed. (p. 158) 				
Table 4A-2. Disk Diffusion	Added:					
QC Ranges for Nonfastidious	Guidance on reading cefipime QC re	esults for E. coli	NCTC 13353 and A.	baumannii NCTO	C 13304 (p. 160)	
Organisms and β-Lactam	• Guidance on reading meropenem QC results for all QC organisms (p. 160)					
Combination Agents	Disk diffusion QC ranges for:					
	Antimicrobial Agent					
			Cefepime-	Cefepime-	Sulbactam-	
	QC Strain	Cefepime	enmetazobactam	taniborbactam	durlobactam	
	E. coli ATCC® 25922		X	X	X	
	P. aeruginosa ATCC® 27853		X	X		
	E. coli ATCC® 35218	X	X	X		
	K. pneumoniae ATCC® 700603		X	X		
	E. coli NCTC 13353		X	X		
	K. pneumoniae ATCC BAA-1705 TM			X		
	A. baumannii NCTC 13304				X	
Table 4B. Disk Diffusion	Added:					
QC Ranges for Fastidious	MH-F agar for S. pneumoniae only to the disk diffusion testing conditions table located in the column for					
Organisms	streptococci and Neisseria meningitidis (p. 166)					
	Revised:					
 Tedizolid disk content and QC ranges for Streptococcus pneumoniae ATCC® 49619 				CC® 49619 (p. 16	5)	
	Reinstated:					
	Norfloxacin QC ranges for all QC s	trains deleted fro	m M100, 29th ed. (p	. 165)		
Table 4D. Disk Diffusion Troubleshooting Guide						

Section/Table	Change(s)						
Tables 5. MIC QC Ranges an	d Associated Tables						
Table 5A-1. MIC QC Ranges	Added:						
for Nonfastidious Organisms	• MIC QC ranges for:						
and Antimicrobial Agents	Antimicrobial Agent						
Excluding β-Lactam	QC Strain Exebacase Ozenoxacin Zoliflodacin						
Combination Agents	E. coli ATCC® 25922						
	S. aureus ATCC® 29213	X	X	X			
	E. faecalis ATCC® 29212	X	X	X			
	Footnote regarding exebacase QC	Footnote regarding exebacase QC ranges (p. 175)					
	Revised:						
	• Eravacycline QC range for <i>E. col</i>	• Eravacycline QC range for <i>E. coli</i> ATCC® 25922 (p. 175)					
	Reinstated:						
	Norfloxacin QC ranges deleted from M100, 29th ed. (p. 176)						
	Deleted:						
	• Plazomicin QC range for E. faecalis ATCC® 29212						
Table 5A-2. MIC QC Ranges	Added:						
for Nonfastidious Organisms	MIC QC ranges for:						
and β-Lactam Combination				crobial Agent		G III	
Agents	QC Strain	Cefepime- enmetazobactam	Cefepime- taniborbactam	Durlobactam	Sulbactam	Sulbactam- durlobactam	
	E. coli ATCC® 25922	X	X	X	X	uuriobactaiii	
	P. aeruginosa ATCC® 27853	X	X	71	71		
	E. coli ATCC® 35218	X	X				
	K. pneumoniae ATCC® 700603	X	X		X		
	E. coli NCTC 13353	X	X				
	K. pneumoniae ATCC® BAA-1705		X				
	K. pneumoniae ATCC® BAA-2814 TM A. baumannii NCTC 13304			X	X	X	
				Λ		Λ	
	Revised:	-lt 1 V	A TCC® I	DAA 2014TM (101)		
	MIC QC range for imipenem-rele	ebactam and K. pne	umoniae ATCC® I	3AA-2814 ^{1M} (p	. 181)		

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ΙΝΔΙΝΉΔΙΝ	of Changes	/ L'antiniiad \
	or Changes	(Continued)

Tables 5. (Continued) Table 5B. MIC QC Ranges for Fastidious Organisms (Broth Dilution Methods) Added:						Overview of Changes (Co.
Table 5B. MIC QC Ranges for Fastidious Organisms (Broth Dilution Methods) Added: MIC QC ranges for: MIC QC strain Cozenoxacin Antimicrobial Agent Ozenoxacin Antimicrobial Agent Ozenoxacin Autimicrobial Agent X S. pneumoniae ATCC® 49247 S. pneumoniae ATCC® 49619 Reinstated: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added:				Change(s)		Section/Table
Fastidious Organisms (Broth Dilution Methods) • MIC QC ranges for: QC Strain Ozenoxacin Zoliflodacin Haemophilus influenzae ATCC® 49247 X S. pneumoniae ATCC® 49619 X X Reinstated: • Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added:						Tables 5. (Continued)
Dilution Methods) QC Strain QC Strain Haemophilus influenzae ATCC® 49247 S. pneumoniae ATCC® 49619 Reinstated: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added: Antimicrobial Agent Ozenoxacin X X X X Added: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Added: Zoliflodacin MIC QC ranges for N. gonorrhoeae ATCC® 49226 (p. 188)					Added:	Table 5B. MIC QC Ranges for
QC Strain Ozenoxacin Zoliflodacin Haemophilus influenzae ATCC® 49247 X S. pneumoniae ATCC® 49619 X X Reinstated: • Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Ozenoxacin Haemophilus influenzae ATCC® 49247 X X					• MIC QC ranges for:	
Haemophilus influenzae ATCC® 49247 S. pneumoniae ATCC® 49619 X Reinstated: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added: Haemophilus influenzae ATCC® 49247 X X X X X X X X X X X X X X X X X X X					Dilution Methods)	
S. pneumoniae ATCC® 49619 Reinstated: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added: X X X X X X X X X X X X Added: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Added: **Coliflodacin MIC QC ranges for N. gonorrhoeae ATCC® 49226 (p. 188) **Added:** **Added:* **Added			Zoliflodacin	Ozenoxacin		
Reinstated: Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added:						
 Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Norfloxacin QC ranges deleted from M100, 29th ed. (p. 185) Added: Zoliflodacin MIC QC ranges for N. gonorrhoeae ATCC® 49226 (p. 188) 						
Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added: Added: Zoliflodacin MIC QC ranges for N. gonorrhoeae ATCC® 49226 (p. 188) Added:						
Neisseria gonorrhoeae (Agar Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added: A Zoliflodacin MIC QC ranges for N. gonorrhoeae ATCC® 49226 (p. 188) Added:				00, 29th ed. (p. 185)	Norfloxacin QC ranges deleted from M	
Dilution Method) Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added:					Added:	Table 5C. MIC QC Ranges for
Tables 6. Preparing Antimicrobial Agent Stock Solutions Table 6A. Solvents and Added:	• Zoliflodacin MIC QC ranges for N. gonorrhoeae ATCC® 49226 (p. 188)			Neisseria gonorrhoeae (Agar		
Table 6A. Solvents and Added:				,		
	Tables 6. Preparing Antimicrobial Agent Stock Solutions					
Dilyonts for Propaging Stock Front and appending configuring the appropriate solvents and dilyonts for activities Listt	Added:			Table 6A. Solvents and		
Footnote regarding confirming the appropriate solvents and different for antimicrobial agents with the	Footnote regarding confirming the appropriate solvents and diluents for antimicrobial agents with the			Diluents for Preparing Stock		
Solutions of Antimicrobial manufacturer (p. 200)	. ,			75 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5		
Agents • Solvent and diluent information for:					• Solvent and diluent information for:	Agents
- Durlobactam					Durlobactam	
- Enmetazobactam					Enmetazobactam	
- Exebacase	- Exebacase					
- Ozenoxacin	- Ozenoxacin					
- Taniborbactam	– Taniborbactam					
- Zoliflodacin						
Reinstated:						
 Norfloxacin solvent and diluent information deleted from M100, 29th ed. 			th ed.	tion deleted from M100, 29	• Norfloxacin solvent and diluent informa	
Table 6C. Preparing Solutions Added preparation instructions for:					Added preparation instructions for:	Table 6C. Preparing Solutions
and Media Containing • Cefepime-enmetazobactam						
Combinations of • Cefepime-taniborbactam	•			Combinations of		
Antimicrobial Agents • Sulbactam-durlobactam					Sulbactam-durlobactam	Antimicrobial Agents

Section/Table	Change(s)		
Appendixes			
Appendix A. Suggestions for Confirming Antimicrobial Susceptibility Test Results and Organism Identification for Agents Approved by the US Food and Drug Administration for Clinical Use (entire table revised)	 Added: Column for antimicrobial class or subclass Clarifying footnotes (pp. 218–219, 221) Newer agents that have US Food and Drug Administration approval (eg, ceftazidime-avibactam, ceftolozane-tazobactam, meropenem-vaborbactam, plazomicin) New ECVs Footnote regarding variations in vancomycin MICs for <i>S. aureus</i> (p. 221) Revised: Title changed from "Suggestions for Confirming Resistant, Intermediate, or Nonsusceptible Antimicrobial Susceptibility Test Results and Organism Identification" to "Suggestions for Confirming Antimicrobial Susceptibility Test Results and Organism Identification for Agents Approved by the US Food and Drug Administration for Clinical Use" Category action step definitions (p. 218) Order of the antimicrobial agents to be more consistent with Tables 2 Categories for: Colistin (Enterobacterales, <i>Acinetobacter baumannii</i> complex, <i>P. aeruginosa</i>) (pp. 218–219) Any carbapenem (<i>A. baumannii</i> complex) (p. 219) Trimethoprim-sulfamethoxazole (<i>S. maltophilia</i>) (p. 219) Vancomycin (<i>S. aureus</i>) (p. 221) Grouped classes of antimicrobial agents together and added categories for: Azithromycin (<i>Salmonella</i> and <i>Shigella</i>; <i>N. gonorrhoeae</i>) (pp. 219, 220) Ceftazidime-avibactam (Enterobacterales) (p. 218) 		
Annual P. D. Lacini	 Ceftolozane-tazobactam (<i>P. aeruginosa</i>) (p. 219) Meropenem-vaborbactam (Enterobacterales) (p. 218) Plazomicin (select Enterobacterales) (p. 218) 		
Appendix B. Intrinsic Resistance	 Added: Clostridioides spp. to section B5, Anaerobic Gram-Positive Bacilli (p. 232) 		
Appendix C. QC Strains for Antimicrobial Susceptibility Tests	Clostratolaes spp. to section B3, Anaerobic Gram-Positive Bachii (p. 232) Added:		

Section/Table	Change(s)					
Appendixes (Continued)	Appendixes (Continued)					
Appendix E. Dosage Regimens	Added:					
Used to Establish Susceptible	Daptomycin SDD MIC breakpoint and dosage regimen for <i>E. faecium</i> only (p. 248)					
or Susceptible-Dose	• Daptomycin dosage regimen for <i>Enterococcus</i> spp. other than <i>E. faecium</i> (p. 248)					
Dependent Breakpoints	• Colistin and/or polymyxin B dosage and treatment regimen reference for Enterobacterales, <i>P. aeruginosa</i> ,					
	and Acinetobacter spp. (pp. 246–247)					
	Revised: Daptomycin susceptible MIC breakpoint for <i>Enterococcus</i> spp. other than <i>E. faecium</i> (p. 248)					
	Deleted:					
	• Meropenem-vaborbactam for <i>P. aeruginosa</i> ; no breakpoints for this antimicrobial agent and organism					
	Colistin dosage regimen for <i>P. aeruginosa</i> and <i>Acinetobacter</i> spp.					
Appendix G. Epidemiological	Revised:					
Cutoff Values	Definitions for wild-type and non-wild-type in section G1, Defining Epidemiological Cutoff Values (p. 254)					
	Deleted:					
	Colistin from Table G1 (ECVs for Enterobacterales) in section G2, Epidemiological Cutoff Value Tables; now assigned a breakpoint in Table 2A					
Appendix I. Cefiderocol Broth	Added:					
Preparation and Reading	• Instructions for preparing zinc stock solution and iron-depleted cation-adjusted Mueller-Hinton broth (p. 274–275)					
Broth Microdilution Minimal	Instructions for reading results and determining end points for broth microdilution MIC tests (p. 275)					
Inhibitory Concentration End	Example photographs showing nontrailing and trailing MIC end points (p. 276)					
Points (new appendix)						

Section/Table	Change(s)				
Glossaries	Glossaries				
I (Part 1). β-Lactams: Class	Added:				
and Subclass Designations	Cefepime-enmetazobactam as a β-lactam combination agent				
and Generic Names	Cefepime-taniborbactam as a β-lactam combination agent				
	Sulbactam-durlobactam as a β-lactam combination agent				
I (Part 2): Non–β-Lactams:	Added:				
Class and Subclass	Exebacase as an antistaphylococcal lysin				
Designations and Generic	Ozenoxacin as a fluoroquinolone				
Names	Zoliflodacin as a spiropyrimidinetrione				
	Moved:				
	Ramoplanin to its own row as a lipoglycodepsipeptide				
	Reinstated:				
	Norfloxacin as a fluoroquinolone deleted from M100, 29th ed.				
II. Antimicrobial Agent	Added:				
Abbreviation(s), Route(s) of	Cefepime-enmetazobactam as a β-lactam combination agent				
Administration, and Drug	Cefepime-taniborbactam as a β-lactam combination agent				
Class	Exebacase as an antistaphylococcal lysin				
	Ozenoxacin as a fluoroquinolone				
	Sulbactam-durlobactam as a β-lactam combination agent				
	Zoliflodacin as a spiropyrimidinetrione				
	Reinstated:				
All distance ATCC® A distance T	Norfloxacin as a fluoroquinolone deleted from M100, 29th ed.				

NOTE: The content of this document is supported by the CLSI consensus process and does not necessarily reflect the views of any single individual or organization.

Abbreviation: ATCC®, American Type Culture Collection.

a ATCC® is a registered trademark of the American Type Culture Collection.

Summary of CLSI Processes for Establishing Breakpoints and Quality Control Ranges

The Clinical and Laboratory Standards Institute (CLSI) is an international, voluntary, not-for-profit, interdisciplinary, standards-developing, and educational organization accredited by the American National Standards Institute that develops and promotes the use of consensus-developed standards and guidelines within the health care community. These consensus standards and guidelines are developed in an open and consensus-seeking forum to cover critical areas of diagnostic testing and patient health care. CLSI is open to anyone or any organization that has an interest in diagnostic testing and patient care. Information about CLSI can be found at www.clsi.org.

The CLSI Subcommittee on Antimicrobial Susceptibility Testing reviews data from a variety of sources and studies (eg, *in vitro*, pharmacokinetics-pharmacodynamics, and clinical studies) to establish antimicrobial susceptibility test methods, breakpoints, and QC parameters. The details of the data necessary to establish breakpoints, QC parameters, and how the data are presented for evaluation are described in CLSI document M23.⁵

Over time, a microorganism's susceptibility to an antimicrobial agent may decrease, resulting in a lack of clinical efficacy and/or safety. In addition, microbiological methods and QC parameters may be refined to ensure more accurate and better performance of susceptibility test methods. Because of these types of changes, CLSI continually monitors and updates information in its documents. Although CLSI standards and guidelines are developed using the most current information available at the time, the field of science and medicine is always changing; therefore, standards and guidelines should be used in conjunction with clinical judgment, current knowledge, and clinically relevant laboratory test results to guide patient treatment.

Additional information, updates, and changes in this document are found in the meeting summary minutes of the Subcommittee on Antimicrobial Susceptibility Testing at https://clsi.org/meetings/ast-file-resources/.

CLSI Reference Methods vs Commercial Methods and CLSI vs US Food and Drug Administration Breakpoints

It is important for users of M02,¹ M07,² and M100 to recognize that the standard methods described in CLSI documents are reference methods. These methods may be used for routine antimicrobial susceptibility testing of patient isolates, for evaluating commercial devices that will be used in medical laboratories, or by drug or device manufacturers for testing new agents or systems. Results generated by reference methods, such as those included in CLSI documents, may be used by regulatory authorities to evaluate the performance of commercial susceptibility testing devices as part of the approval process. Clearance by a regulatory authority indicates the commercial susceptibility testing device provides susceptibility results that are substantially equivalent to results generated using reference methods for the organisms and antimicrobial agents described in the device manufacturer's approved package insert.

CLSI breakpoints may differ from those approved by various regulatory authorities for many reasons, including use of different databases, differences in data interpretation, differences in doses used in different parts of the world, and public health policies. Differences also exist because CLSI proactively evaluates the need for changing breakpoints. The reasons why breakpoints may change and the manner in which CLSI evaluates data and determines breakpoints are outlined in CLSI document M23.⁵

Following a decision by CLSI to change an existing breakpoint, regulatory authorities may also review data to determine how changing breakpoints may affect the safety and effectiveness of the antimicrobial agent for the approved indications. If the regulatory authority changes breakpoints, commercial device manufacturers may have to conduct a clinical trial, submit the data to the regulatory authority, and await review and approval. For these reasons, a delay of one or more years may be needed if a breakpoint and interpretive category change is to be implemented by a device manufacturer. In the United States, it is acceptable for laboratories that use US Food and Drug Administration (FDA)—cleared susceptibility testing devices to use existing FDA breakpoints. Either FDA or CLSI susceptibility breakpoints are acceptable to laboratory accrediting organizations in the United States. Policies in other countries may vary. Each laboratory should check with the manufacturer of its antimicrobial susceptibility test system for additional information on the breakpoints and interpretive categories used in its system's software.

Following discussions with appropriate stakeholders (eg, infectious diseases and pharmacy practitioners, the pharmacy and therapeutics and infection **prevention** committees of the medical staff, and the antimicrobial stewardship team), newly approved or revised breakpoints may be implemented by laboratories. Following verification, CLSI disk diffusion test breakpoints may be implemented as soon as they are published in M100. If a device includes antimicrobial test concentrations sufficient to allow interpretation of susceptibility and resistance to an agent using the CLSI breakpoints, a laboratory could choose to, after appropriate verification, interpret and report results using CLSI breakpoints.

CLSI Breakpoint Additions/Revisions Since 2010

	Date of Addition/Revision*	
Antimicrobial Agent	(M100 edition)	Comments
Enterobacterales		
Azithromycin – <i>S.</i> enterica ser. Typhi only	January 2015 (M100-S25)	
Aztreonam	January 2010 (M100-S20)	
Cefazolin	January 2010 (M100-S20)	Breakpoints were revised twice since 2010.
	January 2011 (M100-S21)	
	January 2014 (M100-S24)	Breakpoints were added to predict results for cefazolin when
	January 2016 (M100S, 26th ed.)	cefazolin is used for therapy of uncomplicated UTIs.
Cefepime	January 2014 (M100-S24)	
Cefiderocol	January 2019 (M100, 29th ed.)	NPBP
	January 2020 (M100, 30th ed.)	Disk diffusion breakpoints were added.
Cefotaxime	January 2010 (M100-S20)	
Ceftaroline	January 2013 (M100-S23)	NPBP
Ceftazidime	January 2010 (M100-S20)	
Ceftazidime-avibactam	January 2018 (M100, 28th ed.)	NPBP
Ceftizoxime	January 2010 (M100-S20)	
Ceftolozane-tazobactam	January 2016 (M100S, 26th ed.)	NPBP
	January 2018 (M100, 28th ed.)	Disk diffusion breakpoints were added.
Ceftriaxone	January 2010 (M100-S20)	
Ciprofloxacin	January 2019 (M100, 29th ed.)	Disk diffusion and MIC breakpoints were revised.
Ciprofloxacin – Salmonella spp.	January 2012 (M100-S22)	Anatomical site–specific breakpoint recommendations were
(including S. enterica ser. Typhi)		removed in 2013.
Colistin	January 2020 (M100, 30th ed.)	NPBP, previously assigned an ECV
Doripenem	June 2010 (M100-S20-U)	NPBP
Ertapenem	June 2010 (M100-S20-U)	Breakpoints were revised twice since 2010.
	January 2012 (M100-S22)	
Imipenem	June 2010 (M100-S20-U)	
Levofloxacin	January 2019 (M100, 29th ed.)	Disk diffusion and MIC breakpoints were revised.
Levofloxacin – Salmonella spp.	January 2013 (M100-S23)	
(including S. enterica ser. Typhi)		
Meropenem	June 2010 (M100-S20-U)	
Meropenem-vaborbactam	January 2019 (M100, 29th ed.)	NPBP
Norfloxacin	January 2020 (M100, 30th ed.)	Reinstated breakpoints deleted from M100, 29th ed.

CLSI Breakpoint Additions/Revisions Since 2010 (Continued)

	Date of Addition/Revision*	
Antimicrobial Agent	(M100 edition)	Comments
Enterobacterales (Continued)		
Ofloxacin – Salmonella spp.	June 2013 (M100-S23)	
(including <i>S.</i> enterica ser. Typhi)		
Pefloxacin – Salmonella spp.	January 2015 (M100-S25)	Surrogate test for ciprofloxacin was added.
(including S. enterica ser. Typhi)		
Polymyxin B	January 2020 (M100, 30th ed.)	NPBP
Pseudomonas aeruginosa		
Cefiderocol	January 2019 (M100, 29th ed.)	NPBP
	January 2020 (M100, 30th ed.)	Disk diffusion breakpoints were added.
Ceftazidime-avibactam	January 2018 (M100, 28th ed.)	NPBP
Ciprofloxacin	January 2019 (M100, 29th ed.)	Disk diffusion and MIC breakpoints were revised.
Colistin	January 2017 (M100, 27th ed.)	MIC breakpoints were revised.
	January 2020 (M100, 30th ed.)	MIC breakpoints were revised.
Doripenem	January 2012 (M100-S22)	
Imipenem	January 2012 (M100-S22)	
Levofloxacin	January 2019 (M100, 29th ed.)	Disk diffusion and MIC breakpoints were revised.
Meropenem	January 2012 (M100-S22)	-
Norfloxacin	January 2020 (M100, 30th ed.)	Reinstated breakpoints deleted from M100, 29th ed.
Piperacillin	January 2012 (M100-S22)	
Piperacillin-tazobactam	January 2012 (M100-S22)	
Polymyxin B	January 2020 (M100, 30th ed.)	MIC breakpoints were revised.
Ticarcillin	January 2012 (M100-S22)	•
Ticarcillin-clavulanate	January 2012 (M100-S22)	
Acinetobacter spp.		
Cefiderocol	January 2019 (M100, 29th ed.)	NPBP
	January 2020 (M100, 30th ed.)	Disk diffusion breakpoints were added.
Colistin	January 2020 (M100, 30th ed.)	MIC breakpoints were revised.
Doripenem	January 2014 (M100-S24)	•
Imipenem	January 2014 (M100-S24)	
Meropenem	January 2014 (M100-S24)	
Polymyxin B	January 2020 (M100, 30th ed.)	MIC breakpoints were revised.
Stenotrophomonas maltophilia		
Cefiderocol	January 2019 (M100, 29th ed.)	NPBP
	January 2020 (M100, 30th ed.)	Disk diffusion breakpoints were added.

CLSI Breakpoint Additions/Revisions Since 2010 (Continued)

CLSI Breakpoint Additions/Revisions Si	Date of Addition/Revision*	
Antimicrobial Agent	(M100 edition)	Comments
Other Non-Enterobacterales		
Norfloxacin	January 2020 (M100, 30th ed.)	Reinstated breakpoints deleted from M100, 29th ed.
Staphylococcus spp.		
Ceftaroline	January 2013 (M100-S23)	NPBP
	January 2019 (M100, 29th ed.)	Disk diffusion and MIC breakpoints were revised to include an
		SDD interpretive category.
Dalbavancin	January 2018 (M100, 28th ed.)	NPBP
Norfloxacin	January 2020 (M100, 30th ed.)	Reinstated breakpoints deleted from M100, 29th ed.
Oritavancin	January 2016 (M100S, 26th ed.)	NPBP
Tedizolid	January 2016 (M100S, 26th ed.)	NPBP
Telavancin	January 2016 (M100S, 26th ed.)	NPBP
Enterococcus spp.		
Dalbavancin	January 2018 (M100, 28th ed.)	NPBP
Daptomycin	January 2019 (M100, 29th ed.)	MIC breakpoints for E. faecium only were added.
	January 2020 (M100, 30th ed.)	MIC breakpoints for Enterococcus spp. other than E. faecium
		were revised.
Norfloxacin	January 2020 (M100, 30th ed.)	Reinstated breakpoints deleted from M100, 29th ed.
Oritavancin	January 2016 (M100S, 26th ed.)	NPBP
Tedizolid	January 2016 (M100S, 26th ed.)	NPBP
Telavancin	January 2016 (M100S, 26th ed.)	NPBP
Haemophilus influenzae and Haemophilus p		
Ceftaroline	January 2013 (M100-S23)	NPBP
Neisseria gonorrhoeae		
Azithromycin	January 2019 (M100, 29th ed.)	NPBP, previously assigned an ECV
Streptococcus pneumoniae		
Ceftaroline	January 2013 (M100-S23)	NPBP
Doxycycline	January 2013 (M100-S23)	NPBP
Tetracycline	January 2013 (M100-S23)	
Streptococcus spp. β-Hemolytic Group		
Ceftaroline	January 2013 (M100-S23)	NPBP
Dalbavancin	January 2018 (M100, 28th ed.)	NPBP
Oritavancin	January 2016 (M100S, 26th ed.)	NPBP
Telavancin	January 2016 (M100S, 26th ed.)	NPBP

CLSI Breakpoint Additions/Revisions Since 2010 (Continued)

	Date of Addition/Revision*			
Antimicrobial Agent	(M100 edition)	Comments		
Streptococcus spp. Viridans Group				
Ceftolozane-tazobactam	January 2016 (M100S, 26th ed.)	NPBP		
Dalbavancin	January 2018 (M100, 28th ed.)	NPBP		
Oritavancin	January 2016 (M100S, 26th ed.)	NPBP		
Tedizolid	January 2016 (M100S, 26th ed.)	NPBP		
Telavancin	January 2016 (M100S, 26th ed.)	NPBP		
Anaerobes				
Piperacillin-tazobactam	January 2017 (M100, 28th ed.)	MIC breakpoints were revised.		

^{*}Previous breakpoints can be found in the edition of M100 that precedes the document listed here, eg, previous breakpoints for aztreonam are listed in M100-S19 (January 2009). Abbreviations: ECV, epidemiological cutoff value; MIC, minimal inhibitory concentration; NPBP, no previous breakpoint existed; SDD, susceptible-dose dependent; UTI, urinary tract infection.

CLSI Epidemiological Cutoff Value Additions/Revisions Since 2015

Antimicrobial Agent	Date of Addition/Revision (M100 edition)	Comments
Enterobacterales		
Azithromycin	January 2016 (M100S, 26th ed.)	For use with Shigella flexneri and Shigella sonnei.
Anaerobes		
Vancomycin	January 2015 (M100-S25)	For use with Cutibacterium (formerly Propionibacterium) acnes.

CLSI Archived Resources

Resource	Web Address for Archived Table
Breakpoints that have been eliminated from M100 since 2010 have been relocated to the CLSI website.	https://clsi.org/media/2654/_m100_archived_drugs_table_2019.pdf
Methods that have been eliminated from M100 have been relocated to the CLSI website.	https://clsi.org/media/1899/_m100_archived_methods_table.pdf
QC ranges that have been eliminated from M100 since 2010 have been relocated to the CLSI website.	https://clsi.org/media/3202/_m100_archived_qc_table.pdf
ECVs that have been replaced by breakpoints have been relocated to the CLSI website.	https://clsi.org/media/3466/_m100_archived_ecvs_table.pdf

Abbreviations: ECV, epidemiological cutoff value; QC, quality control.

Subcommittee on Antimicrobial Susceptibility Testing Mission Statement

The Subcommittee on Antimicrobial Susceptibility Testing is composed of representatives from the professions, government, and industry, including microbiology laboratories, government agencies, health care providers and educators, and pharmaceutical and diagnostic microbiology industries. Using the CLSI voluntary consensus process, the subcommittee develops standards that promote accurate antimicrobial susceptibility testing and appropriate reporting. The mission of the Subcommittee on Antimicrobial Susceptibility Testing is to:

- Develop standard reference methods for antimicrobial susceptibility tests.
- Provide quality control parameters for standard test methods.
- Establish breakpoints and interpretive categories for the results of standard antimicrobial susceptibility tests and provide epidemiological cutoff values when breakpoints are not available.
- Provide suggestions for testing and reporting strategies that are clinically relevant and cost-effective.
- Continually refine standards and optimize detection of emerging resistance mechanisms through development of new or revised methods, breakpoints, and quality control parameters.
- Educate users through multimedia communication of standards and guidelines.
- Foster a dialogue with users of these methods and those who apply them.

The ultimate purpose of the subcommittee's mission is to provide useful information to enable laboratories to assist the clinician in the selection of appropriate antimicrobial therapy for patient care. The standards and guidelines are meant to be comprehensive and to include all antimicrobial agents for which the data meet established CLSI guidelines. The values that guide this mission are quality, accuracy, fairness, timeliness, teamwork, consensus, and trust.

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Instructions for Use of Tables

These instructions apply to:

- Tables 1A and 1B: suggested groupings of antimicrobial agents that should be considered for testing and reporting by microbiology laboratories. These guidelines are based on antimicrobial agents approved by the US Food and Drug Administration (FDA) for clinical use in the United States. In other countries, placement of antimicrobial agents in Tables 1A and 1B should be based on available drugs approved for clinical use by relevant regulatory organizations.
- Tables 2A through 2I: tables for each organism group that contain:
 - Recommended testing conditions
 - Routine QC recommendations (also see Chapter 4 in M02¹ and M07²)
 - General comments for testing the organism group and specific comments for testing particular agent/organism combinations
 - Suggested agents that should be considered for routine testing and reporting by medical microbiology laboratories, as specified in Tables 1A and 1B (test/report groups A, B, C, U)
 - Additional drugs that are appropriate for the respective organism group but would generally not warrant routine testing by a medical microbiology laboratory in the United States (test/report group O for "other"; test/report group Inv. for "investigational" [not yet FDA approved])
 - Zone diameter and minimal inhibitory concentration (MIC) breakpoints
- Tables 1C and 2J: tables containing specific recommendations for testing and reporting results on anaerobes and some of the information listed in the bullets above
- Tables 3A to 3J: tables describing tests to detect particular resistance types in specific organisms or organism groups

I. Selecting Antimicrobial Agents for Testing and Reporting

A. Selecting the most appropriate antimicrobial agents to test and report is a decision best made by each laboratory in consultation with the infectious diseases and pharmacy practitioners, the pharmacy and therapeutics and infection **prevention** committees of the medical staff, and the antimicrobial stewardship team. The recommendations for each organism group include agents of proven efficacy that show acceptable *in vitro* test performance. Considerations in the assignment of agents to specific test/report groups include clinical efficacy, prevalence of resistance, minimizing emergence of resistance, cost, FDA clinical indications for use, and current consensus recommendations for first-choice and alternative drugs. Tests on selected agents may be useful for infection **prevention** purposes.

M100, 30th ed

- B. Drugs listed together in a single box are agents for which interpretive categories (susceptible, intermediate, or resistant) and clinical efficacy are similar. Within each box, an "or" between agents indicates agents for which cross-resistance and cross-susceptibility are nearly complete. Results from one agent connected by an "or" can be used to predict results for the other agent (ie, equivalent agents). For example, Enterobacterales susceptible to cefotaxime can be considered susceptible to ceftriaxone. The results obtained from testing cefotaxime could be reported along with a comment that the isolate is also susceptible to ceftriaxone. For drugs connected with an "or," combined major and very major errors are fewer than 3%, and minor errors are fewer than 10%, based on a large population of bacteria tested (see CLSI document M23⁵ for description of error types). In addition, to qualify for an "or," at least 100 strains with resistance to the agents in question must be tested, and a result of "resistant" must be obtained with all agents for at least 95% of the strains. "Or" is also used for comparable agents when tested against organisms for which "susceptible-only" breakpoints are provided (eg, cefotaxime or ceftriaxone with *H. influenzae*). When no "or" connects agents within a box, testing of one agent cannot be used to predict results for another, owing either to discrepancies or insufficient data.
- C. Test/Report Groups
- 1. **Group A antimicrobial agents,** as listed in Tables 1A, 1B, and 1C, are considered appropriate for inclusion in a routine, primary testing panel, as well as for routine reporting of results for the specific organism groups.
- 2. **Group B** includes antimicrobial agents that may warrant primary testing, but they may be reported only selectively, such as when the organism is resistant to agents of the same antimicrobial class, as in group A. Other indications for reporting the result might include a selected specimen source (eg, a third-generation cephalosporin for enteric bacilli from cerebrospinal fluid (CSF) or trimethoprim-sulfamethoxazole for urinary tract isolates); a polymicrobial infection; infections involving multiple sites; cases of patient allergy, intolerance, or failure to respond to an antimicrobial agent in group A; or for infection **prevention**.
- 3. **Group** C includes alternative or supplemental antimicrobial agents that may necessitate testing in those institutions that harbor endemic or epidemic strains resistant to several of the primary drugs (especially in the same class, eg, β-lactams); for treatment of patients allergic to primary drugs; for treatment of unusual organisms (eg, chloramphenicol for extraintestinal isolates of *Salmonella* spp.); or for reporting to infection **prevention** as an epidemiological aid.
- 4. **Group U ("urine")** includes certain antimicrobial agents (eg, nitrofurantoin and certain quinolones) that are used only or primarily for treating UTIs. These agents should not be routinely reported against pathogens recovered from other infection sites. An exception to this rule is for **Enterobacterales** in Table 1A, in which cefazolin is listed as a surrogate agent for oral cephalosporins. Other antimicrobial agents with broader indications may be included in group U for specific urinary pathogens (eg, *Enterococcus* and ciprofloxacin).
- 5. **Group O ("other")** includes antimicrobial agents that have a clinical indication for the organism group but are generally not candidates for routine testing and reporting in the United States.

6. **Group Inv.** ("investigational") includes antimicrobial agents that are investigational for the organism group and have not yet been approved by the FDA for use in the United States.

D. Selective Reporting

Each laboratory should decide which agents in the tables to report routinely (group A) and which might be reported only selectively (from group B), in consultation with the infectious diseases and pharmacy practitioners, the pharmacy and therapeutics and infection **prevention** committees of the health care institution, and the antimicrobial stewardship team. Selective reporting should improve the clinical relevance of test reports and help minimize the selection of multiresistant, health care—associated strains by overusing broad-spectrum antimicrobial agents. Results for group B antimicrobial agents tested, but not reported routinely, should be available on request, or they may be reported for selected specimen types. Unexpected resistance, when confirmed, should be reported (eg, resistance to a secondary agent but susceptibility to a primary agent, such as a *P. aeruginosa* isolate resistant to amikacin but susceptible to tobramycin; as such, both drugs should be reported). In addition, each laboratory should develop a protocol to cover isolates that are confirmed as resistant to all agents on its routine test panels. This protocol should include options for testing additional agents in-house or sending the isolate to a referral laboratory.

II. Breakpoint and Interpretive Category Definitions

- A. **Breakpoint** minimal inhibitory concentration (MIC) or zone diameter value used to categorize an organism as susceptible, susceptible dose dependent, intermediate, resistant, or nonsusceptible; **NOTE 1:** MIC or zone diameter values generated by a susceptibility test can be interpreted based on established breakpoints; **NOTE 2:** Because breakpoints are based on pharmacologically and clinically rich datasets using *in vitro* and *in vivo* data, they are considered robust predictors of likely clinical outcome; **NOTE 3:** Also known as "clinical breakpoint"; **NOTE 4:** See **interpretive category.**
- B. **Interpretive category** category derived from microbiological characteristics, pharmacokinetic-pharmacodynamic parameters, and clinical outcome data, when available; **NOTE 1:** MIC or zone diameter values generated by a susceptibility test can be interpreted based on established breakpoints; **NOTE 2:** See **breakpoint.**

EXAMPLE:

Interpretive	Brea	kpoints
Category	MIC, μg/mL	Zone Diameter, mm
Susceptible	≤4	≥20
Susceptible-dose	8–16	15–19
dependent		
Intermediate	8–16	15–19
Resistant	≥32	≤14
Nonsusceptible	>1	<17

MIC or zone diameter value breakpoints and interpretive categories are established per CLSI document M23⁵ for categories of susceptible, intermediate, and resistant (and susceptible-dose dependent and nonsusceptible, when appropriate).

- **susceptible (S)** a category defined by a breakpoint that implies that isolates with an MIC at or below or a zone diameter at or above the susceptible breakpoint are inhibited by the usually achievable concentrations of antimicrobial agent when the dosage recommended to treat the site of infection is used, resulting in likely clinical efficacy.
- susceptible-dose dependent (SDD) a category defined by a breakpoint that implies that susceptibility of an isolate depends on the dosage regimen that is used in the patient. To achieve levels that are likely to be clinically effective against isolates for which the susceptibility testing results (either MICs or zone diameters) are in the SDD category, it is necessary to use a dosage regimen (ie, higher doses, more frequent doses, or both) that results in higher drug exposure than that achieved with the dose that was used to establish the susceptible breakpoint. Consideration should be given to the maximum, literature-supported dosage regimen, because higher exposure gives the highest probability of adequate coverage of an SDD isolate. Appendix E lists the doses used when establishing SDD categories. The drug label should be consulted for recommended doses and adjustment for organ function; NOTE: The SDD category may be assigned when doses well above those used to calculate the susceptible breakpoint are supported by the literature, widely used clinically, and/or approved and for which sufficient data to justify the designation exist and have been reviewed. This category also includes a buffer zone for inherent variability in test methods, which should prevent small, uncontrolled, technical factors from causing major discrepancies in interpretations, especially for drugs with narrow pharmacotoxicity margins. See Appendix F for additional information.

- intermediate (I) a category defined by a breakpoint that includes isolates with MICs or zone diameters within the intermediate range that approach usually attainable blood and tissue levels and/or for which response rates may be lower than for susceptible isolates; NOTE: The intermediate category implies clinical efficacy in anatomical sites where the drugs are physiologically concentrated. An I with a "^" in Tables 2 indicates agents that have the potential to concentrate at an anatomical site. The I category also includes a buffer zone for inherent variability in test methods, which should prevent small, uncontrolled, technical factors from causing major discrepancies in interpretations, especially for drugs with narrow pharmacotoxicity margins.
- resistant (R) a category defined by a breakpoint that implies that isolates with an MIC at or above or a zone diameter at or below the resistant breakpoint are not inhibited by the usually achievable concentrations of the agent with normal dosage schedules and/or that demonstrate MICs or zone diameters that fall in the range in which specific microbial resistance mechanisms are likely, and clinical efficacy of the agent against the isolate has not been reliably shown in treatment studies.
- nonsusceptible (NS) a category used for isolates for which only a susceptible breakpoint is designated because of the absence or rare occurrence of resistant strains. Isolates for which the antimicrobial agent MICs are above or the zone diameters are below the value indicated for the susceptible breakpoint should be reported as nonsusceptible; NOTE 1: An isolate that is interpreted as nonsusceptible does not necessarily mean that the isolate has a resistance mechanism. It is possible that isolates with MICs above the susceptible breakpoint that lack resistance mechanisms may be encountered within the wild-type distribution after the time the susceptible-only breakpoint was set; NOTE 2: The term "nonsusceptible" should not be used when the text is describing an organism/drug category with intermediate and resistant interpretive categories. Isolates that are in the categories of "intermediate" or "resistant" could be called "not susceptible" rather than "nonsusceptible."

C. Example of Breakpoints and Interpretive Categories as Used in Table 2

Antimicrobial	Disk	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm		_	retive Cat IC Break µg/mL	_	
Agent	Content	S	I*	R	S	I*	R
X	30 μg	≥20	15–19	≤14	≤4	8–16	≥32
Y	_	_	_	_	≤1	2	≥4
Z	10 μg	≥16	_	_	≤1	_	_

^{*} Or SDD, if appropriate.

Abbreviations: Î, intermediate; R, resistant; S, susceptible; SDD, susceptible-dose dependent.

For antimicrobial agent X with breakpoints in the table above, the susceptible breakpoint is $\leq 4~\mu g/mL$ or $\geq 20~mm$ and the resistant breakpoint is $\geq 32~\mu g/mL$ or $\leq 14~mm$. For some antimicrobial agents (eg, antimicrobial agent Y), only MIC breakpoints may be available. For these agents, the disk diffusion zone diameters do not correlate with MIC values or data have not been evaluated as described in CLSI document M23. Technical issues may also preclude the use of the disk diffusion method for some agents. For some antimicrobial agents (eg, antimicrobial agent Z) only a "susceptible" category exists. For these agents, the absence or rare occurrence of resistant strains precludes defining any results categories other than "susceptible." For strains yielding results suggestive of a "nonsusceptible" category, organism identification and antimicrobial susceptibility test results should be confirmed (see Appendix A). In examples Y and Z, a dash mark (–) indicates a disk is not available or that breakpoints are not applicable.

III. Reporting Results

A. Organisms Included in Table 2

The MIC values determined as described in M07² may be reported directly to clinicians for patient care purposes. However, it is essential that an interpretive category result (S, **SDD**, I, R, **or NS**) also be provided routinely to facilitate understanding of the MIC report by clinicians. Zone diameter measurements without an interpretive category should not be reported. Recommended interpretive categories for various MIC and zone diameter values are included in tables for each organism group and are based on the evaluation of data as described in CLSI document M23.⁵

Laboratories should only report results for agents listed in Table 2 specific to the organism being tested. It is not appropriate to apply disk diffusion or MIC breakpoints borrowed from a table in which the organism is not listed. There may be rare cases for which an agent may be appropriate for an isolate but for which there are no CLSI breakpoints (eg, tigecycline). In these cases, the FDA prescribing information document for the agent should be consulted.

For more information on reporting epidemiological cutoff values in the medical laboratory, see Appendix G.

B. Organisms Excluded From Table 2

For some organism groups excluded from Tables 2A through 2J, CLSI document M45⁶ provides suggestions for standardized methods for AST, including information about drug selection, interpretation, and QC. The organism groups covered in that guideline are *Abiotrophia* and *Granulicatella* spp. (formerly known as nutritionally deficient or nutritionally variant streptococci); *Aerococcus* spp.; *Aeromonas* spp.; *Bacillus* spp. (not *Bacillus anthracis*); *Campylobacter jejuni/coli; Corynebacterium* spp. (including *Corynebacterium diphtheriae*); *Erysipelothrix rhusiopathiae; Gemella* spp.; the HACEK group: *Aggregatibacter* spp. (formerly *Haemophilus aphrophilus, Haemophilus paraphrophilus, Haemophilus actinomycetemcomitans*), *Cardiobacterium* spp., *Eikenella corrodens*, and *Kingella* spp.; *Helicobacter pylori; Lactobacillus* spp.; *Lactococcus* spp.; *Leuconostoc* spp.; *Listeria monocytogenes; Micrococcus* spp.; *Moraxella catarrhalis; Pasteurella* spp.; *Pediococcus* spp.; *Rothia mucilaginosa;* potential agents of bioterrorism; and *Vibrio* spp., including *Vibrio cholerae*.

For organisms other than those in the groups mentioned above, studies are not yet adequate to develop reproducible, definitive standards to interpret results. These organisms may need different media or different incubation atmospheres, or they may show marked strain-to-strain variation in growth rate. For these microorganisms, consultation with an infectious diseases specialist is recommended for guidance in determining the need for susceptibility testing and in results interpretation. Published reports in the medical literature and current consensus recommendations for therapy of uncommon microorganisms may preclude the need for testing. If necessary, a dilution method usually is the most appropriate testing method, and this may necessitate submitting the organism to a referral laboratory. Physicians should be informed of the limitations of results and advised to interpret results with caution.

C. Cumulative Antibiograms

Policies regarding the generation of cumulative antibiograms should be developed together with the infectious diseases service, infection **prevention** personnel, the pharmacy and therapeutics committee, and the antimicrobial stewardship team. See CLSI document M39⁷ for detailed instructions on generating cumulative antibiograms.

D. MIC Reporting Concentrations

When serial twofold dilution MICs are being prepared and tested, the actual dilution scheme is, eg:

16, 8, 4, 2, 1, 0.5, 0.25, 0.125, 0.0625, 0.03125 μg/mL, etc. (see Table 7 for additional dilutions).

For convenience only, not because these are the actual concentrations tested, it was decided to use the following values in Tables 7, 8A, and 8B: 16, 8, 4, 2, 1, 0.5, 0.25, 0.12, 0.06, 0.03 µg/mL, etc.

The values that appear in the tables are equivalent to the actual values tested, eg, $0.12~\mu g/mL = 0.125~\mu g/mL$, and laboratories should report an MIC of $\leq 0.125~\mu g/mL$ as $\leq 0.12~\mu g/mL$.

IV. Therapy-Related Comments

Some comments in the tables relate to therapy concerns. These are denoted with an **Rx** symbol. It may be appropriate to include some of these comments (or modifications thereof) on the patient report. An example would be inclusion of a comment when rifampin is being reported stating that "Rifampin should not be used alone for antimicrobial therapy." Antimicrobial dosage regimens often vary widely among practitioners and institutions. In some cases, the MIC breakpoints rely on pharmacokinetic-pharmacodynamic (PK-PD) data, using specific human dosage regimens. In cases in which specific dosage regimens are important for properly applying breakpoints, the dosage regimen is listed. These dosage regimen comments are not generally intended for use on individual patient reports.

V. Confirmation of Patient Results

Multiple test parameters are monitored by following the QC recommendations described in M100. However, acceptable results derived from testing QC strains do not guarantee accurate results when testing patient isolates. It is important to review all the results obtained from all drugs tested on a patient's isolate before reporting the results. This review should include but not be limited to ensuring that 1) the AST results are consistent with the identification of the isolate; 2) the results from individual agents within a specific drug class follow the established hierarchy of activity rules (eg, in general, third-generation cephems are more active than first- or second-generation cephems against **Enterobacterales**); and 3) the isolate is susceptible to those agents for which resistance has not been documented (eg, vancomycin and *Streptococcus* spp.) and for which only "susceptible" breakpoints exist in M100.

Unusual or inconsistent results should be confirmed by rechecking various testing parameters detailed in Appendix A. Each laboratory must develop its own policies for confirming unusual or inconsistent antimicrobial susceptibility test results. The list provided in Appendix A emphasizes results that are most likely to affect patient care.

VI. Development of Resistance and Testing of Repeat Isolates

Isolates that are initially susceptible may become intermediate or resistant after therapy is initiated. Therefore, subsequent isolates of the same species from a similar anatomical site should be tested to detect resistance that may have developed. Development of resistance can occur within as little as three to four days and has been noted most frequently in *Enterobacter* (including *Klebsiella* [formerly *Enterobacter*] *aerogenes*), *Citrobacter*, and *Serratia* spp. with third-generation cephalosporins, in *P. aeruginosa* with all antimicrobial agents, and in staphylococci with fluoroquinolones. For *S. aureus*, vancomycin-susceptible isolates may become vancomycin intermediate during the course of prolonged therapy.

In certain circumstances, the decision to perform susceptibility tests on subsequent isolates necessitates knowledge of the specific situation and the severity of the patient's condition (eg, an isolate of *E. cloacae* from a blood culture on a premature infant or methicillin (oxacillin)-resistant *S. aureus* [MRSA] from a patient with prolonged bacteremia). Laboratory guidelines on when to perform susceptibility testing on repeat isolates should be determined after consultation with the medical staff.

VII. Warning

Some of the comments in the tables relate to dangerously misleading results that can occur when certain antimicrobial agents are tested and reported as susceptible against specific organisms. These are denoted with the word "Warning."

Location	Organism	Antimicrobial Agents		
"Warning": The f	following antimicrobial agent/organism	combinations may appear active in vitro but are not effective clinically and		
must not be reporte	ed as susceptible.			
Table 2A	Salmonella spp., Shigella spp.	1st- and 2nd-generation cephalosporins, cephamycins, and aminoglycosides		
Table 2D	Enterococcus spp.	Aminoglycosides (except for high-level resistance testing), cephalosporins, clindamycin, and trimethoprim-sulfamethoxazole		
"Warning": The following antimicrobial agents that are included in this document should not be routinely reported for bacteria isolated from CSF. These antimicrobial agents are not the drugs of choice and may not be effective for treating CSF infections caused by these organisms (ie, the bacteria included in Tables 2A through 2J):				
Tables 2A through 2J	Bacteria isolated from CSF	Agents administered by oral route only, 1st- and 2nd-generation cephalosporins and cephamycins, clindamycin, macrolides, tetracyclines, and fluoroquinolones		

Abbreviation: CSF, cerebrospinal fluid.

VIII. Routine, Supplemental, Screening, Surrogate Agent, and Equivalent Agent Testing to Determine Susceptibility and Resistance to Antimicrobial Agents

The testing categories are defined as follows:

- Routine test: disk diffusion or broth or agar dilution MIC tests for routine clinical testing
- Supplemental (not routine) test: test that detects susceptibility or resistance to a drug or drug class by method other than routine disk diffusion or broth or agar dilution MIC and does not need additional tests to confirm susceptibility or resistance
 - Some supplemental tests identify a specific resistance mechanism and may be required or optional for reporting specific clinical results.
- Screening test: test that provides presumptive results; additional testing typically only needed for a specific result (eg, only if screen is positive)

M100, 30th ed

- Surrogate agent test: test performed with an agent that replaces a test performed with the antimicrobial agent of interest and is used when the agent of interest cannot be tested due to availability or performance issues (eg, surrogate agent performs better than the agent of interest)
- Equivalent agent test: test performed with an agent that predicts results of closely related agents of the same class and increases efficiency by limiting testing of multiple closely related agents. Equivalent agents are identified by:
 - Listing equivalent agents with an "or" in Tables 1 and 2. "Or" indicates cross-susceptibility and cross-resistance is nearly complete (very major error + major error < 3%; minor error < 10%) and only one agent needs to be tested.
 - Listing agents that are equivalent and results that can be deduced by testing the equivalent agent in a comment (see Tables 1 and 2).

The following tables include tests that fall into the supplemental, screening, surrogate agent, and equivalent agent test categories. The tables for supplemental, screening, and surrogate agent tests are comprehensive. The table for equivalent agent tests includes several examples, and many other equivalent agent tests are described throughout Tables 1 and 2.

Supplemental Tests (Required)

Supplemental Test	Organisms	Test Description	Required for:	Table Location
Inducible clindamycin resistance	 Staphylococcus spp. S. pneumoniae Streptococcus spp. β-hemolytic group 	Broth microdilution or disk diffusion with clindamycin and erythromycin tested together	Isolates that test erythromycin resistant and clindamycin susceptible or intermediate before reporting the isolate as clindamycin susceptible	3Н
β-lactamase	Staphylococcus spp.	Chromogenic cephalosporin (all staphylococci), penicillin disk diffusion zoneedge test (S. aureus only)	Isolates that test penicillin susceptible before reporting the isolate as penicillin susceptible	3E

Supplemental Tests (Optional)

Supplemental Test	Organisms	Test Description	Optional for:	Table Location
ESBL	 E. coli K. pneumoniae Klebsiella oxytoca Proteus mirabilis 	Broth microdilution or disk diffusion clavulanate inhibition test for ESBLs	Isolates demonstrating reduced susceptibility to cephalosporins Results that indicate presence or absence of ESBLs	3A
CarbaNP	EnterobacteralesP. aeruginosa	Colorimetric assay for detecting carbapenem hydrolysis	Isolates demonstrating reduced susceptibility to carbapenems Results that indicate presence or absence of certain carbapenemases	3B, 3B-1
mCIM with or without eCIM	 mCIM only: Enterobacterales and P. aeruginosa mCIM with eCIM: Enterobacterales only 	Disk diffusion for detecting carbapenem hydrolysis (inactivation) eCIM add-on enables differentiation of metallo-β-lactamases from serine carbapenemases in Enterobacterales isolates that are positive for mCIM	Isolates demonstrating reduced susceptibility to carbapenems Results that indicate presence or absence of certain carbapenemases	3C
Colistin agar test	• Enterobacterales • P. aeruginosa	Modified agar dilution	Determining the colistin MIC	3D
Colistin broth disk elution	Enterobacterales P. aeruginosa	Tube dilution using colistin disks as antimicrobial agent source	Determining the colistin MIC	3D
Oxacillin salt agar	• S. aureus	Agar dilution; MHA with 4% NaCl and 6 µg/mL oxacillin	Detecting MRSA; see cefoxitin surrogate agent tests, which are preferred	3F

Abbreviations: eCIM, EDTA-modified carbapenem inactivation method; ESBL, extended-spectrum β-lactamase; mCIM, modified carbapenem inactivation method; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; MRSA, methicillin (oxacillin)-resistant Staphylococcus aureus.

Screening Tests

			When to Perform		
Screening Test	Organisms	Test Description	Confirmatory Test	Confirmatory Test	Table Location
Vancomycin agar screen	• S. aureus • Enterococcus spp.	Agar dilution; BHI with 6 μg/mL vancomycin	If screen positive	Vancomycin MIC	3 G
HLAR by disk diffusion	• Enterococcus spp.	Disk diffusion with gentamicin and streptomycin	If screen inconclusive	Broth microdilution, agar dilution MIC	3J

Abbreviations: BHI, brain heart infusion; HLAR, high-level aminoglycoside resistance; MIC, minimal inhibitory concentration.

Surrogate Agent Tests

Surrogate				Table
Agent	Organisms	Test Description	Results	Location
Cefazolin	 E. coli Klebsiella pneumoniae P. mirabilis 	Broth microdilution or disk diffusion	When used for therapy of uncomplicated UTIs, predicts results for the following oral antimicrobial agents: cefaclor, cefdinir, cefpodoxime, cefprozil, cefuroxime, cephalexin, and loracarbef Cefazolin as a surrogate may overcall resistance to cefdinir, cefpodoxime, and cefuroxime. If cefazolin tests resistant, test these drugs individually if needed for therapy.	1A, 2A
Cefoxitin	 S. aureus S. lugdunensis S. epidermidis Other Staphylococcus spp. (excluding S. pseudintermedius and S. schleiferi) 	Broth microdilution: S. aureus S. lugdunensis Disk diffusion: S. aureus S. lugdunensis Other Staphylococcus spp., excluding S. pseudintermedius and S. schleiferi	Predicts results for mecA-mediated methicillin (oxacillin) resistance.	1A, 2C, 3F
Oxacillin	S. pneumoniae	Disk diffusion	Predicts penicillin susceptibility if oxacillin zone is ≥20 mm. If oxacillin zone is ≤19 mm, penicillin MIC must be done.	1B, 2G
Pefloxacin	• Salmonella spp.	Disk diffusion	Predicts reduced susceptibility to ciprofloxacin	2A

Abbreviations: MIC, minimal inhibitory concentration; PBP2a, penicillin-binding protein 2a; UTI, urinary tract infection.

Examples of Equivalent Agent Tests

Agents	Organisms	Identified by	Table Location
Cefotaxime or ceftriaxone	Enterobacterales	"Or"	1A and 2A
Colistin or polymyxin B	Enterobacterales, Pseudomonas aeruginosa, Acinetobacter baumannii complex	"Or"	2A, 2B-1, and 2B-2
Azithromycin or clarithromycin or erythromycin	Staphylococcus spp.	"Or"	1A and 2C
Penicillin-susceptible staphylococci are susceptible to other β-lactam agents with established clinical efficacy for staphylococcal infections (including both penicillinase-labile and penicillinase-stable agents; see Glossary I). Penicillinresistant staphylococci are resistant to penicillinase-labile penicillins.	Staphylococcus spp.	Note listed	1A and 2C
The results of ampicillin susceptibility tests should be used to predict the activity of amoxicillin.	Haemophilus spp.	Note listed	1B and 2E
The results of ampicillin susceptibility tests should be used to predict the activity of amoxicillin.	Anaerobes	Note listed	2Ј

IX. Quality Control and Verification

Recommendations for QC are included in various tables and appendixes. Acceptable ranges for QC strains are provided in Tables 4A-1 through 4B for disk diffusion and Tables 5A-1 through 5E for MIC testing. Guidance for QC frequency and modifications of AST systems is found in Table 4C for disk diffusion and Table 5F for MIC testing. Guidance for troubleshooting out-of-range results is included in Table 4D for disk diffusion and Table 5G for MIC testing. Additional information is available in Appendix C (eg, QC organism characteristics, QC testing recommendations).

Implementing any new diagnostic test requires verification.⁸ Each laboratory that introduces a new AST system or adds a new antimicrobial agent to an existing AST system must verify or establish that, before reporting patient test results, the system meets performance specifications for that system. Verification generally involves testing patient isolates with the new AST system and comparing results to those obtained with an established reference method or a system that has been previously verified. Testing patient isolates may be done concurrently with the two systems. Alternatively, organisms with known MICs or zone sizes may be used for the verification. Guidance on verification studies is not included in this document. Other publications describe AST system verification (eg, CLSI document M52⁹ and Patel J, et al.¹⁰).

X. Abbreviations and Acronyms

AST antimicrobial susceptibility testing $ATCC^{\otimes_a}$ American Type Culture Collection

BHI brain heart infusion

BLNAR β-lactamase negative, ampicillin-resistant

BMHA blood Mueller-Hinton agar BSC biological safety cabinet

BSL-2 biosafety level 2 BSL-3 biosafety level 3

CAMHB cation-adjusted Mueller-Hinton broth

CAT colistin agar test

CBDE colistin broth disk elution CFU colony-forming unit(s)

CMRNG chromosomally mediated penicillin-resistant Neisseria gonorrhoeae

CSF cerebrospinal fluid DMSO dimethyl sulfoxide

ECV epidemiological cutoff value

eCIM EDTA-modified carbapenem inactivation method

EDTA ethylenediaminetetraacetic acid
ESBL extended-spectrum β-lactamase
FDA US Food and Drug Administration
HLAR high-level aminoglycoside resistance

HTM Haemophilus test medium

I intermediate

ICR inducible clindamycin resistance

IM intramuscular
ID identification
LHB lysed horse blood

mCIM modified carbapenem inactivation method

MHA Mueller-Hinton agar

MH-F agar Mueller-Hinton fastidious agar

 $^{^{\}text{a}}$ $\text{ATCC}^{\text{\circledR}}$ is a registered trademark of the American Type Culture Collection.

MHB	Mueller-Hinton broth
MIC	minimal inhibitory concentration

MRS methicillin (oxacillin)-resistant staphylococci

MRSA methicillin (oxacillin)-resistant Staphylococcus aureus

NAD β-nicotinamide adenine dinucleotide
NCTC National Collection of Type Cultures
NPBP no previous breakpoint existed

NS nonsusceptible NWT non-wild-type

PBP2a penicillin-binding protein 2a PCR polymerase chain reaction

PK-PD pharmacokinetic-pharmacodynamic

pH negative logarithm of hydrogen ion concentration

QC quality control R resistant S susceptible

SDD susceptible-dose dependent

TSA tryptic soy agar
TSB trypticase soy broth
UTI urinary tract infection

WT wild-type

References

- ¹ CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ³ CLSI. Methods for Antimicrobial Susceptibility Testing of Anaerobic Bacteria. 9th ed. CLSI standard M11. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ⁴ Adeolu M, Alnajar S, Naushad S, Gupta RS. Genome-based phylogeny and taxonomy of the 'Enterobacteriales': proposal for Enterobacterales ord. nov. divided into the families *Enterobacteriaceae, Erwiniaceae* fam. nov., *Pectobacteriaceae* fam. nov., *Yersiniaceae* fam. nov., *Hafniaceae* fam. nov., *Morganellaceae* fam. nov., and *Budviciaceae* fam. nov. *Int J Syst Evol Microbiol.* 2016;66(12):5575-5599.
- 5 CLSI. Development of In Vitro Susceptibility Testing Criteria and Quality Control Parameters. 5th ed. CLSI guideline M23. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 6 CLSI. Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently Isolated or Fastidious Bacteria. 3rd ed. CLSI guideline M45. Wayne, PA: Clinical and Laboratory Standards Institute; 2016.
- 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.
- 8 Centers for Medicare & Medicaid Services, US Department of Health and Human Services. Part 493—Laboratory Requirements; Standard: Establishment and verification of performance specifications (Codified at 42 CFR §493.1253). Office of the Federal Register; published annually.
- 9 CLSI. Verification of Commercial Microbial Identification and Antimicrobial Susceptibility Testing Systems. 1st ed. CLSI guideline M52. Wayne, PA: Clinical and Laboratory Standards Institute; 2015.
- Patel J, Sharp S, Novak-Weekley S. Verification of antimicrobial susceptibility testing methods: a practical approach. Clin Microbiol Newslett. 2013;35(13):103-109.

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M100, 30th ed.

Table 1A. Suggested Groupings of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That Should Be Considered for Testing and Reporting on Nonfastidious Organisms by Microbiology Laboratories in the United States

Group A: Includes antimicrobia		olon in a routino, primary tooting pain	or, ao tron ao for foatino reporting of
results for the specific organism Enterobacterales	m group. Pseudomonas aeruginosa	Staphylococcus spp.	Enterococcus spp. ⁿ
Ampicillin ^d	Ceftazidime	Azithromycin ^b or	Ampicillin ^o
Cefazoline	Gentamicin	clarithromycin ^b or	Penicillin
0014201111	Tobramycin	erythromycin ^b	1 Grilomin
Gentamicin ^d	Piperacillin-tazobactam	Clindamycin ^b	
Tobramycin ^d	'	Oxacillin ^{j,l,*,†,§}	
		Cefoxitin ^{j,l,†}	
		(surrogate test for oxacillin)	
		Penicillin ^j	
		Trimethoprim-sulfamethoxazole	
Group B: Includes antimicrobia	I agents that may warrant primary testing		such as when the organism is resista
to agents of the same antimicro		,	, cach ac 1111011 and or g ament is 1001014
Amikacin ^d	Amikacin	Ceftaroline ⁱ	Daptomycin ^{k,*}
Amoxicillin-clavulanate	Aztreonam	Daptomycin ^{k,*}	Linezolid
Ampicillin-sulbactam			Tedizolid ^q
Ceftazidime-avibactam	Cefepime	Linezolid	Vancomycin
Ceftolozane-tazobactam	Ceftazidime-avibactam	Tedizolid ⁱ	
Meropenem-vaborbactam	Ceftolozane-tazobactam		
Piperacillin-tazobactam			
Cefuroxime	Ciprofloxacin	Doxycycline	
	Levofloxacin	Minocycline ^b	
Cefepime	Doripenem	Tetracycline ^a	
Cefotetan	Imipenem	Vancomycin*	
Cefoxitin	Meropenem	·	
Cefotaxime ^{d,e} or			
Ceftriaxone ^{d,e}			
Ciprofloxacin ^d		Rifampin ^h	
Levofloxacind			
Doripenem			
Ertapenem			
Imipenem			
Meropenem			
Trimethoprim-sulfamethoxazole ^d			

Table 1A. (Continued)

Group C: Includes alternative or supplemental antimicrobial agents that may require testing in institutions that harbor endemic or epidemic strains for treatment of patients allergic to primary drugs, for treatment of unusual organisms, or for reporting to resistant to several of the primary drugs, for treatment of patients allergic to primary drugs, for treatment of unusual organisms, or for reporting to infection prevention as an epidemiological aid.

Enterobacterales	Pseudomonas aeruginosa	Staphylococcus spp.	Enterococcus spp.n
Aztreonam		Chloramphenicol ^b	Gentamicin (high-level
Ceftazidime			resistance testing only)
		Ciprofloxacin or	Streptomycin (high-level
Ceftaroline		levofloxacin	resistance testing only)
Chloramphenicol ^{b,d}			
Tetracycline ^a		Moxifloxacin	
,			Dalbavancin ^{s,*}
		Gentamicin ^m	Oritavancin ^{s,*}
		Dalbavancin ^{i,*}	Telavancin ^{s,*}
		Oritavancin ^{i,*}	
		Telavancin ^{i,*}	
Group U: Includes antimicrobial agents	that are used only or primarily for tro	eating UTIs.	
Cefazolin		Nitrofurantoin	Ciprofloxacin
(surrogate test for uncomplicated UTI) [‡]			Levofloxacin
Fosfomycin ^f		Sulfisoxazole	
Nitrofurantoin		Trimethoprim	Fosfomycin ^r
Sulfisoxazole		·	Nitrofurantoin
Trimethoprim			Tetracycline ^a
Group A: Includes antimicrobial agents	considered appropriate for inclusior	n in a routine, primary testing panel	, as well as for routine reporting of
results for the specific organism group.			
A simusta ha a tamana	Develop a laborito de acesta de acesta la co	04	Other New Entereder structure (*

results for the specific organism grou	0.		
Acinetobacter spp.	Burkholderia cepacia complex	Stenotrophomonas maltophilia	Other Non-Enterobacterales ^{g,*}
Ampicillin-sulbactam	Levofloxacin*	Levofloxacin	Ceftazidime
Ceftazidime	Meropenem	Minocycline	Gentamicin
Ciprofloxacin	Trimethoprim-sulfamethoxazole	Trimethoprim-sulfamethoxazole	Tobramycin
Levofloxacin			
Doripenem			
Imipenem			
Meropenem			
Gentamicin			
Tobramycin			

Table 1A. (Continued)			
Group B: Includes antimicrobial ag	ents that may warrant primary testing	but may be reported only selectively, s	uch as when the organism is resistant
to agents of the same antimicrobia	l class in Group A.c		
Amikacin	Ceftazidime	Ceftazidime*	Amikacin
Piperacillin-tazobactam	Minocycline		Aztreonam
Cefepime			Cefepime
Cefotaxime			Ciprofloxacin
Ceftriaxone			Levofloxacin
Doxycycline			Imipenem
Minocycline			Meropenem
Trimethoprim-sulfamethoxazole			Piperacillin-tazobactam
			Trimethoprim-sulfamethoxazole
Group C: Includes alternative or su	ipplemental antimicrobial agents that	may require testing in institutions that h	narbor endemic or epidemic strains
	• •	ic to primary drugs, for treatment of unu	sual organisms, or for reporting to
infection prevention as an epidemi			
	Chloramphenicol ^{b,*}	Chloramphenicol ^{b,*}	Cefotaxime
			Ceftriaxone
			Chloramphenicol ^b
Group U: Includes antimicrobial ag	ents that are used only or primarily fo	or treating UTIs.	
Tetracycline ^a			Sulfisoxazole
			Tetracycline ^a

Abbreviations: MIC, minimal inhibitory concentration; UTI, urinary tract infection.

^{*} MIC testing only; disk diffusion test is unreliable.

[†] See oxacillin and cefoxitin comments in Table 2C for using cefoxitin as a surrogate for oxacillin.

[‡] See cefazolin comments in Table 2A for using cefazolin as a surrogate for oral cephalosporins and for reporting cefazolin when used for therapy in uncomplicated UTIs.

[§] For S. aureus, S. lugdunensis, and other Staphylococcus spp. (excluding S. epidermidis, S. pseudintermedius, and S. schleiferi), only MIC testing, not disk diffusion testing, is acceptable; see exceptions in Table 2C.

Table 1A. (Continued)

"Warning": The following antimicrobial agents that are included in this document should not be routinely reported for bacteria isolated from CSF. These antimicrobial agents are not the drugs of choice and may not be effective for treating CSF infections caused by these organisms (ie, the bacteria included in Tables 2A through 2J):

- Agents administered by oral route only
- 1st- and 2nd-generation cephalosporins and cephamycins
- Clindamycin
- Macrolides
- Tetracyclines
- Fluoroquinolones

Footnotes

<u>General</u>

- a. 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.
- b. Not routinely reported on organisms isolated from the urinary tract.
- c. Section I, C.2. in the Instructions for Use of Tables lists additional examples of when a Group B agent might be reported.

Enterobacterales

d. **WARNING:** For *Salmonella* spp. and *Shigella* spp., aminoglycosides, first- and second-generation cephalosporins, and cephamycins may appear active *in vitro*, but are not effective clinically and should not be reported as susceptible.

Routine susceptibility testing is not indicated for nontyphoidal *Salmonella* spp. isolated from intestinal sources. In contrast, susceptibility testing is indicated for all *Shigella* isolates.

When fecal isolates of *Salmonella* and *Shigella* spp. are tested, only ampicillin, a fluoroquinolone, and trimethoprim-sulfamethoxazole should be reported routinely. In addition, for extraintestinal isolates of *Salmonella* spp., a third-generation cephalosporin should be tested and reported, and if requested, chloramphenicol may be tested and reported. Susceptibility testing is indicated for typhoidal *Salmonella* (*S.* enterica ser. Typhi and *Salmonella* enterica ser. Paratyphi A–C) isolated from extraintestinal and intestinal sources.

- e. Cefotaxime or ceftriaxone should be tested and reported on isolates from CSF in place of cefazolin.
- f. For testing and reporting of *E. coli* urinary tract isolates only.

Table 1A. (Continued)

Other Non-Enterobacterales

g. Other non-Enterobacterales include *Pseudomonas spp.* and other nonfastidious, glucose-nonfermenting, gram-negative bacilli but exclude *P. aeruginosa, Acinetobacter* spp., *B. cepacia* complex, and *S. maltophilia*. Refer to each respective organism column for suggested antimicrobial agents to test and report.

Recommendations for testing and reporting of *Aeromonas hydrophila* complex, *Burkholderia mallei*, *Burkholderia pseudomallei*, and *Vibrio* spp. (including *V. cholerae*) are found in CLSI document M45.¹

Staphylococcus spp.

- h. Rx: Rifampin should not be used alone for antimicrobial therapy.
- i. For S. aureus only, including methicillin (oxacillin)-resistant S. aureus (MRSA).
- j. Penicillin-susceptible staphylococci are also susceptible to other β-lactam agents with established clinical efficacy for staphylococcal infections. Penicillin-resistant staphylococci are resistant to penicillinase-labile penicillins. **Methicillin** (oxacillin)-resistant staphylococci are resistant to all currently available β-lactam antimicrobial agents, with the exception of **ceftaroline**. Thus, susceptibility or resistance to a wide array of β-lactam antimicrobial agents may be deduced from testing only penicillin and either cefoxitin or oxacillin. Routine testing of other β-lactam agents, except **ceftaroline**, is not advised.
- k. Daptomycin should not be reported for isolates from the respiratory tract.
- I. If a penicillinase-stable penicillin is tested, oxacillin is the preferred agent, and results can be applied to the other penicillinase-stable penicillins (refer to Glossary I). Detection of **methicillin** (oxacillin) resistance in staphylococci is achieved by using specific methods as described in Tables 2C and **3F**.
- m. For staphylococci that test susceptible, gentamicin is used only in combination with other active agents that test susceptible.

Enterococcus spp.

- n. **Warning:** For *Enterococcus* spp., cephalosporins, aminoglycosides (except for high-level resistance testing), clindamycin, and trimethoprim-sulfamethoxazole may appear active *in vitro*, but are not effective clinically and should not be reported as susceptible.
- o. 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 *Enterococcus faecalis*.

Table 1A. (Continued)

- p. Enterococci susceptible to penicillin are predictably susceptible to ampicillin, amoxicillin, ampicillin-sulbactam, amoxicillin-clavulanate, and piperacillin-tazobactam for non–β-lactamase-producing enterococci. However, enterococci susceptible to ampicillin cannot be assumed to be susceptible to penicillin. If penicillin results are needed, testing of penicillin is required. *Rx:* Combination therapy with ampicillin, penicillin, or vancomycin (for susceptible strains) plus an aminoglycoside is usually indicated for serious enterococcal infections, such as endocarditis, unless high-level resistance to both gentamicin and streptomycin is documented; such combinations are predicted to result in synergistic killing of the *Enterococcus*. For strains with low-level penicillin or ampicillin resistance when combination therapy with a β-lactam is being considered, see additional testing and reporting information in Table 3J.²
- q. For testing and reporting of *E. faecalis* only.
- r. For testing and reporting of *E. faecalis* urinary tract isolates only.
- s. For testing and reporting of vancomycin-susceptible *E. faecalis* only.

References for Table 1A

- ¹ CLSI. *Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently Isolated or Fastidious Bacteria*. 3rd ed. CLSI guideline M45. Wayne, PA: Clinical and Laboratory Standards Institute; 2016.
- Murray BE, Arias CA, Nannini EC. Glycopeptides (vancomycin and teicoplanin), streptogramins (quinupristin-dalfopristin), lipopeptides (daptomycin), and lipoglycopeptides (telavancin). In: Bennett JE, Dolin R, Blaser MJ. *Mandell, Douglas, and Bennett's Principles and Practice of Infectious Diseases*. 8th ed. Philadelphia, PA: Elsevier Saunders; 2015:377-400.

Meropenem^e

M100, 30th ed

Table 1B. Suggested Groupings of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That Should Be Considered for Testing and Reporting on Fastidious Organisms by Microbiology Laboratories in the United States

Group A: Includes antimicrobial agents considered appropriate for inclusion in a routine, primary testing panel, as well as for routine reporting of results for the specific organism group. Haemophilus influenzaee and Haemophilus Neisseria Streptococcus Streptococcus spp. Streptococcus spp. parainfluenzae gonorrhoeae pneumoniaek **β-Hemolytic Group**^q Viridans Group^q Ampicillin^{n,*} Ampicilline,g Azithromycin*,† Erythromycina,c Clindamycin^{c,p} Penicillin^{n,*} Ceftriaxone[†] Cefixime[†] Ciprofloxacin† Penicillin^I Erythromycin^{a,c,p} Tetracycline^{b,†} (oxacillin disk) Penicillino,† or Trimethoprimampicillino,† sulfamethoxazole Group B: Includes antimicrobial agents that may warrant primary testing but may be reported only selectively, such as when the organism is resistant to agents of the same antimicrobial class, as in Group A.d Ampicillin-sulbactam Cefepime* Cefepime or Cefepime Cefotaxime1,* cefotaxime or Cefotaxime Cefotaximee or Ceftriaxone^{I,*} ceftriaxone Ceftriaxone ceftazidimee or Clindamycin^c Vancomycin Vancomycin ceftriaxone^e Ciprofloxacin or Doxycycline levofloxacin or Levofloxacin^k moxifloxacin Moxifloxacin^k

Meropenem^{l,*}
Tetracycline^b
Vancomycin^l

Table 1B. (Continued)

Group C: Includes alternative or supplemental antimicrobial agents that may require testing in institutions that harbor endemic or epidemic strains resistant to several of the primary drugs, for treatment of patients allergic to primary drugs, for treatment of unusual organisms, or for reporting to infection prevention as an epidemiological aid.

Haemophilus influenzae ^e and Haemophilus parainfluenzae	Neisseria gonorrhoeae ^j	Streptococcus pneumoniae ^k	<i>Streptococcus</i> spp. β-Hemolytic Group ^q	<i>Streptococcus</i> spp. Viridans Group ^q
Azithromycin ^f Clarithromycin ^f		Amoxicillin* Amoxicillin-clavulanate*	Ceftaroline	Ceftolozane-tazobactam
Aztreonam		Cefuroxime*	Chloramphenicol ^c	Chloramphenicol ^c
Amoxicillin-clavulanatef		Ceftaroline	Daptomycin ^{r,*}	Clindamycin ^c
Cefaclor ^f Cefprozil ^f		Chloramphenicol ^c	Levofloxacin	Erythromycin ^{a,c}
Cefdinir ^f or cefixime ^f or cefpodoxime ^f		Ertapenem* Imipenem*	Linezolid Tedizolid ^s Dalbavancin ^{u,*}	Linezolid Tedizolid ^t Dalbavancin ^{u,*}
Ceftaroline ^h		Linezolid Rifampin ^m	Oritavancin [*] Telavancin [*]	Oritavancin* Telavancin*
Cefuroxime ^f		, <u>.</u>		
Chloramphenicol ^c				
Ertapenem or imipenem				
Rifampin ⁱ				
Tetracycline ^b				
Trimethoprim-sulfamethoxazole				

Abbreviations: CSF, cerebrospinal fluid; MIC, minimal inhibitory concentration.

^{*} MIC testing only; disk diffusion test is unreliable.

[†] Routine testing is not necessary (see footnotes j and o).

M100, 30th ed

Table 1B. (Continued)

"Warning": The following antimicrobial agents that are included in this document should not be routinely reported for bacteria isolated from CSF. These antimicrobial agents are not the drugs of choice and may not be effective for treating CSF infections caused by these organisms (ie, the bacteria included in Tables 2A through 2J):

- Agents administered by oral route only
- 1st- and 2nd-generation cephalosporins and cephamycins
- Clindamycin
- Macrolides
- Tetracyclines
- Fluoroquinolones

Footnotes

General

- a. Susceptibility and resistance to azithromycin, clarithromycin, and dirithromycin can be predicted by testing erythromycin.
- b. Organisms that are susceptible to tetracycline are also considered susceptible to doxycycline and minocycline.
- c. Not routinely reported for organisms isolated from the urinary tract.
- d. Section I, C.2. in the Instructions for Use of Tables lists additional examples of when a Group B agent might be reported.

Haemophilus spp.

- e. For isolates of *H. influenzae* from CSF, only results of testing with ampicillin, any of the third-generation cephalosporins listed, and meropenem are appropriate to report.
- f. Amoxicillin-clavulanate, azithromycin, cefaclor, cefdinir, cefixime, cefpodoxime, cefprozil, cefuroxime, and clarithromycin are used as empiric therapy for respiratory tract infections due to *Haemophilus* spp. The results of susceptibility tests with these antimicrobial agents are often not necessary for managing individual patients.
- g. The results of ampicillin susceptibility tests should be used to predict the activity of amoxicillin. The majority of *H. influenzae* isolates that are resistant to ampicillin and amoxicillin produce a TEM-type β-lactamase. In most cases, a direct β-lactamase test can provide a rapid means of detecting ampicillin and amoxicillin resistance.
- h. For H. influenzae only.

Table 1B. (Continued)

i. May be appropriate only for prophylaxis of case contacts. Refer to Table 2E.

Neisseria gonorrhoeae

j. Culture and susceptibility testing of *N. gonorrhoeae* should be considered in cases of treatment failure. Antimicrobial agents recommended for testing include, at a minimum, the agents listed in group A. The most current guidelines for treatment and testing are available from the Centers for Disease Control and Prevention at https://www.cdc.gov/std/gonorrhea/stdfact-gonorrhea.htm.

Streptococcus pneumoniae

- k. S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin or moxifloxacin cannot be assumed to be susceptible to levofloxacin.
- I. Penicillin and cefotaxime, ceftriaxone, or meropenem should be tested by a reliable MIC method (such as that described in M07¹) and reported routinely with CSF isolates of *S. pneumoniae*. 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 ≤ 19 mm, penicillin, cefotaxime, ceftriaxone, or meropenem MICs should be determined.
- m. *Rx:* Rifampin should not be used alone for antimicrobial therapy.

Streptococcus spp.

- n. *Rx:* Penicillin- or ampicillin-intermediate isolates may necessitate combined therapy with an aminoglycoside for bactericidal action.
- o. Penicillin and ampicillin are drugs of choice for treating β-hemolytic streptococcal infections. Susceptibility testing of penicillins and other β-lactams approved by the US Food and Drug Administration for treating β-hemolytic streptococcal infections does not need to be performed routinely, because nonsusceptible isolates (ie, penicillin MICs > 0.12 and ampicillin MICs > 0.25 μg/mL) are extremely rare in any β-hemolytic streptococci and have not been reported for *Streptococcus pyogenes*. If testing is performed, any β-hemolytic streptococcal isolate found to be nonsusceptible should be re-identified, retested, and, if confirmed, submitted to a public health laboratory (see Appendix A for additional instructions).

M100, 30th ed

Table 1B. (Continued)

- p. *Rx:* Recommendations for intrapartum prophylaxis for group B streptococci are penicillin or ampicillin. Although cefazolin is recommended for penicillin-allergic women at low risk for anaphylaxis, those at high risk for anaphylaxis may receive clindamycin. Group B streptococci are susceptible to ampicillin, penicillin, and cefazolin, but may be resistant to erythromycin and clindamycin. When group B *Streptococcus* is isolated from a pregnant woman with severe penicillin allergy (high risk for anaphylaxis), erythromycin and clindamycin (including inducible clindamycin resistance [ICR]) should be tested, and only clindamycin should be reported. Erythromycin, even when tested for determination of ICR, should not be reported. See Table 3H.
- q. For this table, the β-hemolytic group includes the large colony–forming pyogenic strains of streptococci with group A (*S. pyogenes*), C, or G antigens and strains with group B (*S. agalactiae*) antigen. Small colony–forming β-hemolytic strains with group A, C, F, or G antigens (*Streptococcus anginosus* group, previously termed "*Streptococcus milleri*") are considered part of the viridans group, and breakpoints for the viridans group should be used.
- Daptomycin should not be reported for isolates from the respiratory tract.
- s. For reporting against *S. pyogenes* and *Streptococcus agalactiae* only.
- t. For reporting against S. anginosus group (includes S. anginosus, Streptococcus intermedius, and Streptococcus constellatus) only.
- u. For reporting against S. pyogenes, S. agalactiae, Streptococcus dysgalactiae, and S. anginosus group.

NOTE 1: For information about the selection of appropriate antimicrobial agents; explanation of test/report groups A, B, C, and U; and explanation of the listing of agents within boxes, including the meaning of "or" between agents, refer to the Instructions for Use of Tables that precede Table 1A.

NOTE 2: Information in boldface type is new or modified since the previous edition.

Reference for Table 1B

¹ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

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M100, 30th ed

Table 1C. Suggested Groupings of Antimicrobial Agents Approved by the US Food and Drug Administration for Clinical Use That Should Be Considered for Testing and Reporting on Anaerobic Organisms by Microbiology Laboratories in the United States

Group A: Includes antimicrobial agents considered to be appro of results for the specific organism group.	priate for inclusion in a routine, primary testing panel, as well as for routine reporting
Gram-Negative Anaerobes	Gram-Positive Anaerobes ^a
Amoxicillin-clavulanate	Ampicillin ^b
Ampicillin-sulbactam	Penicillin ^b
Piperacillin-tazobactam	Amoxicillin-clavulanate
	Ampicillin-sulbactam
	Piperacillin-tazobactam
Clindamycin	Clindamycin
Doripenem	Doripenem
Ertapenem	Ertapenem
Imipenem	Imipenem
Meropenem	Meropenem
Metronidazole	Metronidazole
	ents that may require testing in institutions that harbor endemic or epidemic strains
infection prevention as an epidemiological aid.	nts allergic to primary drugs, for treatment of unusual organisms, or for reporting to
infection prevention as an epidemiological aid. Penicillin ^b Ampicillin ^b Cefotetan	Cefotetan
infection prevention as an epidemiological aid. Penicillin ^b Ampicillin ^b	
infection prevention as an epidemiological aid. Penicillin ^b Ampicillin ^b Cefotetan Cefoxitin Ceftizoxime	Cefotetan Cefoxitin Ceftizoxime
infection prevention as an epidemiological aid. Penicillin ^b Ampicillin ^b Cefotetan Cefoxitin	Cefotetan Cefoxitin
infection prevention as an epidemiological aid. Penicillin ^b Ampicillin ^b Cefotetan Cefoxitin Ceftizoxime Ceftriaxone Chloramphenicol	Cefotetan Cefoxitin Ceftizoxime
infection prevention as an epidemiological aid. Penicillin ^b Ampicillin ^b Cefotetan Cefoxitin Ceftizoxime Ceftriaxone	Cefotetan Cefoxitin Ceftizoxime

Footnotes

- a. Many non-spore-forming, gram-positive anaerobic rods are resistant to metronidazole (see Appendix D).
- b. If β -lactamase positive, report as resistant to penicillin and ampicillin. Be aware that β -lactamase-negative isolates may be resistant to penicillin and ampicillin by other mechanisms.

For Use With M11

Table 1C. (Continued)

NOTE 1: For information about the selection of appropriate antimicrobial agents; explanation of test/report groups A and C; and explanation of the listing of agents within boxes, refer to the Instructions for Use of Tables that precede Table 1A.

NOTE 2: Most anaerobic infections are polymicrobial, including both β -lactamase-positive and β -lactamase-negative strains. Testing may not be necessary for isolates associated with polymicrobial anaerobic infections. However, if susceptibility testing is requested, only the organism most likely to be resistant (eg, *Bacteroides* spp. and *Parabacteroides* spp.) should be tested and results reported (see Appendix D).

NOTE 3: Specific *Clostridium* spp. (eg, *Clostridium septicum, Clostridium sordellii*) may be the singular cause of infection and are typically susceptible to penicillin and ampicillin. Penicillin and clindamycin resistance have been reported in *Clostridium perfringens*. Agents in group A of Table 1C should be tested and reported for *Clostridium* spp.

NOTE 4: Information in boldface type is new or modified since the previous edition.

Table 2A. Zone Diameter and MIC Breakpoints for Enterobacterales

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB; iron-depleted CAMHB for

cefiderocol (see Appendix I)1

Agar dilution: MHA

Inoculum: Broth culture method or colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35°C±2°C; ambient air

Disk diffusion: 16–18 hours Dilution methods: 16–20 hours **Routine QC Recommendations** (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Escherichia coli ATCC®a 25922

Pseudomonas aeruginosa ATCC® 27853 (for carbapenems)

Staphylococcus aureus ATCC® 25923 (for Salmonella enterica ser. Typhi azithromycin disk diffusion testing only; see Table 4A-1)

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

Refer to Tables 3A, 3B, and 3C for additional testing, reporting, and QC for Enterobacterales.

General Comments

- (1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02,² Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide³). Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. Strains of Proteus spp. may swarm into areas of inhibited growth around certain antimicrobial agents. With Proteus spp., ignore the thin veil of swarming growth in an otherwise obvious zone of growth inhibition. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.
- (2) When fecal isolates of Salmonella and Shigella spp. are tested, only ampicillin, a fluoroquinolone, and trimethoprim-sulfamethoxazole should be reported routinely. In addition, for extraintestinal isolates of Salmonella spp., a 3rd-generation 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. In contrast, susceptibility testing is indicated for all Shigella isolates.
- (3) The dosage regimens shown in the comments column below are those needed to achieve plasma drug exposures (in adults with normal renal and hepatic functions) on which breakpoints were based. When implementing new breakpoints, it is strongly recommended that laboratories share this information with infectious diseases practitioners, pharmacists, pharmacy and therapeutics committees, infection **prevention** committees, and the antimicrobial stewardship team.
- (4) Intermediate ranges denoted with a "^" for the applicable antimicrobial agents in the drug groups in Tables 2 are based on the known ability of these agents to concentrate in the urine; some agents may also have the potential to concentrate at other anatomical sites (eg, epithelial lining).

NOTE: Information in boldface type is new or modified since the previous edition.

Table 2A. Enterobacterales (Continued)

Test/Report Antimicrobial		Disk	Zone	Diamete	ategories er Breakpo whole mm	ints,	l:		ve Categories Breakpoints, µg/mL	and	
Group Agent	Content	S	SDD	ı	R	S	SDD	I	R	Comments	
PENICILLINS											
Α	Ampicillin	10 μg	≥17	-	14–16 ^	≤13	≤8	-	16 ^	≥32	(5) Results of ampicillin testing can be used to predict results for amoxicillin. See general comment (2).
0	Piperacillin	100 μg	≥21	. –	18–20 ^	≤17	≤16	-	32-64^	≥128	
0	Mecillinam	10 μg	≥15	_	12–14^	≤11	≤8	-	16 ^	≥32	(6) For testing and reporting of <i>E. coli</i> urinary tract isolates only.
β-LACTAM C	OMBINATION AGENTS										
В	Amoxicillin-clavulanate	20/10 µg	≥18	-	14–17^	≤13	≤8/4	- 1	16/8 ^	≥32/16	
В	Ampicillin-sulbactam	10/10 µg	≥15	-	12-14^	≤11	≤8/4	- :	16/8 ^	≥32/16	
В	Ceftolozane- tazobactam	30/10 µg	≥21	-	18–20 ^	≤17	≤2/4	-	4/4^	≥8/4	(7) Breakpoints are based on a dosage regimen of 1.5 g administered every 8 h.
В	Ceftazidime-avibactam	30/20 μg	≥21	-	-	≤20 :	≤8/4	- :	-	≥16/4	(8) Breakpoints are based on a dosage regimen of 2.5 g (2 g ceftazidime+0.5 g avibactam) every 8 h administered over 2 h.
				-							(9) Confirmatory MIC testing is indicated for isolates with zones of 20–22 mm to avoid reporting falsesusceptible or false-resistant results.
В	Meropenem- vaborbactam	20/10 µg	≥18	_	15–17^	: ≤14 :	≤4/8	-	8/8^	≥16/8	(10) Breakpoints are based on a dosage regimen of 4 g (2 g meropenem + 2 g vaborbactam) every 8 h administered over 3 h.
В	Piperacillin-tazobactam	100/10 µg	≥21	_	18–20 ^	≤17	≤ 16/4	-	32/4-64/4^	≥128/4	
0	Ticarcillin-clavulanate	75/10 µg	≥20	<u> </u>	15–19 ^	≤14	≤ 16/2	-	32/2-64/2^	≥128/2	

Table 2A. Enterobacterales (Continued)

					ategories er Breakpo		Int	erpretive (MIC Bre	Categories akpoints,	and		
Test/Report	Antimicrobial	Disk		nearest	whole mm	1		μg	/mL			
Group	Agent	Content	S	S SDD I R				SDD	ı	R	Comments	
CEDUENIC (D	CERUFAC (DARENTERAL) (Including confederacing L.H. III) and IV. Places refer to Classery L.											

CEPHEMS (PARENTERAL) (Including cephalosporins I, II, III, and IV. Please refer to Glossary I.)

- (11) WARNING: For Salmonella spp. and Shigella spp., 1st- and 2nd-generation cephalosporins and cephamycins may appear active in vitro but are not effective clinically and should not be reported as susceptible.
- (12) Following evaluation of PK-PD properties, limited clinical data, and MIC distributions, revised breakpoints for cephalosporins (cefazolin, cefotaxime, ceftazidime, ceftizoxime, and ceftriaxone) and aztreonam were first published in January 2010 (M100-S20) and are listed in this table. Cefuroxime (parenteral) was also evaluated; however, no change in breakpoints was necessary for the dosage indicated below. When using the current breakpoints, routine ESBL testing is no longer necessary before reporting results (ie, it is no longer necessary to edit results for cephalosporins, aztreonam, or penicillins from susceptible to resistant). However, ESBL testing may still be useful for epidemiological or infection prevention purposes. For laboratories that have not implemented the current breakpoints, ESBL testing should be performed as described in Table 3A.

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* spp., or *Proteus* spp., ESBL testing should be performed (see Table 3A). If isolates test ESBL positive, the results for moxalactam, cefonicid, cefamandole, and cefoperazone should be reported as resistant.

(13) Enterobacter, Klebsiella (formerly Enterobacter) aerogenes, Citrobacter, and Serratia may develop resistance during prolonged therapy with 3rd-generation cephalosporins as a result of derepression of AmpC β-lactamase. Therefore, isolates that are initially susceptible may become resistant within 3 to 4 days after initiation of therapy. Testing repeat isolates may be warranted.

A	Cefazolin	30 µg	≥23	_	20–22	≤19	≤2	-	4	≥8	(14) Breakpoints when cefazolin is used for therapy of infections other than uncomplicated UTIs due to <i>E. coli, K. pneumoniae,</i> and <i>P. mirabilis.</i> Breakpoints are based on a dosage regimen of 2 g administered every 8 h. See comment (12).
U	Cefazolin	30 µg	≥15	-	_	≤14	≤16	_	_	≥32	(15) Breakpoints when cefazolin is used for therapy of uncomplicated UTIs due to <i>E. coli, K. pneumoniae</i> , and <i>P. mirabilis</i> . Breakpoints are based on a dosage regimen of 1 g administered every 12 h. See additional information in CEPHEMS (ORAL).
С	Ceftaroline	30 µg	≥23	_	20–22^	≤19	≤0.5	-	1^	≥2	(16) Breakpoints are based on a dosage regimen of 600 mg administered every 12 h.

Table 2A. Enterobacterales (Continued)

Test/Report	Antimicrobial	Disk		Diamete	Categories er Breakpo whole mm	ints,	Interpretive Categories and MIC Breakpoints, µg/mL		and		
Group Agent	Content	S	SDD	I	R	S	SDD	ı	R	Comments	
CEPHEMS (P	ARENTERAL) (Including of	cephalospor	ins I, II, I	II, and IV	. Please re	fer to G	lossary I.)	(Continue	ed)		
В	Cefepime	30 µg	≥25	19– 24	-	≤18	≤2	4–8	_	≥16	(17) The breakpoint for susceptible is based on a dosage regimen of 1 g administered every 12 h. The breakpoint for SDD is based on dosage regimens that result in higher cefepime exposure, either higher doses or more frequent doses or both, up to approved maximum dosage regimens. See Appendix E for more information about breakpoints and dosage regimens. Also see the definition of SDD in the Instructions for Use of Tables section.
B B	Cefotaxime or ceftriaxone	30 µg 30 µg	≥26 ≥23	-	23–25 ^ 20–22 ^	≤22 ≤19	≤1 ≤1	-	2^ 2^	≥4 ≥4	(18) 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. See comment (12).
В	Cefotetan	30 μg	≥16	-	13–15^	≤12	≤16	-	32^	≥64	, ,
В	Cefoxitin	30 μg	≥18	-	15–17 ^	≤14	≤8	-	16^	≥32	(19) Breakpoints are based on a dosage regimen of at least 8 g per day (eg, 2 g administered every 6 h).
В	Cefuroxime (parenteral)	30 μg	≥18	-	15–17 ^	≤14	≤8	-	16^	≥32	(20) Breakpoints are based on a dosage regimen of 1.5 g administered every 8 h. See comment (12).
С	Ceftazidime	30 μg	≥21	-	18–20 ^	≤17	≤4	-	8^	≥16	(21) Breakpoints are based on a dosage regimen of 1 g administered every 8 h. See comment (12).
0	Cefamandole	30 μg	≥18	-	15–17 ^	≤14	≤8	-	16 ^	≥32	See comment (12).
0	Cefmetazole	30 μg	≥16	-	13–15^	≤12	≤16	-	32^	≥64	(22) Insufficient new data exist to reevaluate breakpoints listed here.
0	Cefonicid	30 μg	≥18	-	15–17^	≤14	≤8	<u> </u>	16^	≥32	See comment (12).
0	Cefoperazone	75 μg	≥21	_	16–20	≤15	≤16	 -	32	≥64	See comment (12).
0	Ceftizoxime	30 μg	≥25	_	22–24^	≤21	≤1	-	2^	≥4	(23) Breakpoints are based on a dosage regimen of 1 g administered every 12 h. See comment (12).
0	Moxalactam	30 μg	≥23	-	15–22^	≤14	≤8	-	16–32 ^	≥64	See comment (12).

M100, 30th ed

Interpretive Categories and Interpretive Categories and Zone Diameter Breakpoints, MIC Breakpoints, Test/Report Antimicrobial Disk nearest whole mm μg/mL Group Agent Content S SDD R S SDD R Comments CEPHEMS (PARENTERAL) (Including cephalosporins I, II, III, and IV. Please refer to Glossary I.) (Continued) Cefiderocol 30 μg ≥16 12-15^ ≤4 ≥16 (24) Breakpoints are based on a dosage Inv. ≤11 regimen of 2 g every 8 h administered over 3 h. **CEPHEMS (ORAL)** Cefuroxime 30 μg ≥23 15-22^ ≤4 8-16^ ≥32 See comment (25). ≤14 U Cefazolin 30 μg ≥15 ≤14 ≤16 ≥32 (25) Breakpoints are for cefazolin when (surrogate test for used as a surrogate test to predict oral cephalosporins results for the oral agents cefaclor, and uncomplicated cefdinir, cefpodoxime, cefprozil, cefuroxime, cephalexin, and loracarbef UTIs) when used for therapy of uncomplicated UTIs due to E. coli, K. pneumoniae, and P. mirabilis. Cefazolin as a surrogate may overcall resistance to cefdinir, cefpodoxime, and cefuroxime. If cefazolin tests resistant, test these drugs individually if needed for therapy. 0 Loracarbef 30 μg 15-17^ ≤14 ≤8 16^ ≥32 (26) Do not test Citrobacter, Providencia, ≥18 or Enterobacter spp. with cefdinir or loracarbef by disk diffusion because falsesusceptible results have been reported. See comment (25). 0 Cefaclor 15-17^ 16^ See comment (25). 30 μα ≥18 ≤8 ≥32 ≤14 0 Cefdinir See comments (25) and (26). 5 μg ≥20 17-19^ ≤16 ≤1 ≥4 0 Cefixime ≥19 16-18^ ≤15 ≤1 2^ \geq 4 (27) Do not test Morganella spp. with 5 μg cefixime, cefpodoxime, or cefetamet by disk diffusion. 0 18-20**^** See comments (25) and (27). Cefpodoxime 10 μg ≥21 ≤17 ≤2 ≥8

Test/Report	Antimicrobial	Disk		Diamete	ategories a er Breakpoir whole mm		Inte	MIC Bre	Categories akpoints, /mL		
Group	Agent	Content	S	SDD	1	R	S	SDD	ı	R	Comments
CEPHEMS (OF	RAL) (Continued)										
0	Cefprozil	30 μg	≥18	-	15–17^	≤14	≤8	-	16^	≥32	(28) Do not test <i>Providencia</i> spp. with cefprozil by disk diffusion because false-susceptible results have been reported. See comment (25).
Inv.	Cefetamet	10 μg	≥18	-	15–17 ^	≤14	≤4	-	8^	≥16	See comment (27).
Inv.	Ceftibuten	30 μg	≥21	-	18–20 ^	≤17	≤8	-	16 ^	≥32	(29) For testing and reporting of urinary tract isolates only.
MONOBACTA	MS										
С	Aztreonam	30 µg	≥21	-	18–20 ^	≤17	≤4	-	8^	≥16	(30) Breakpoints are based on a dosage regimen of 1 g administered every 8 h. See comment (12).

CARBAPENEMS

(31) Following evaluation of PK-PD properties, limited clinical data, and MIC distributions that include recently described carbapenemase-producing strains, revised breakpoints for carbapenems were first published in June 2010 (M100-S20-U) and are listed below. Because of limited treatment options for infections caused by organisms with carbapenem MICs or zone diameters in the intermediate range, clinicians may wish to design carbapenem dosage regimens that use maximum recommended doses and possibly prolonged intravenous infusion regimens, as has been reported in the literature.⁴⁻⁷ Consultation with an infectious diseases practitioner is recommended for isolates for which the carbapenem MICs or zone diameter results from disk diffusion testing are in the intermediate or resistant ranges.

Laboratories using **Enterobacterales** MIC breakpoints for carbapenems described in M100-S20 (January 2010) should perform the CarbaNP test, mCIM, eCIM, and/or a molecular assay (refer to Tables 3B and 3C for methods) when isolates of **Enterobacterales** are suspicious for carbapenemase production based on imipenem or meropenem MICs 2–4 µg/mL or ertapenem MIC 2 µg/mL (refer to Tables 3B-1 and 3C-1 for guidance on reporting). After implementing the current breakpoints, these additional tests may not need to be performed other than for epidemiological or infection **prevention** purposes (ie, it is no longer necessary to edit results for the carbapenems to resistant if a carbapenemase producer is detected). See Appendix H, Table H3 regarding suggestions for reporting when molecular and phenotypic methods are discordant.

The following information is provided as background on carbapenemases in **Enterobacterales** that are largely responsible for MICs and zone diameters in the intermediate and resistant ranges, and thus the rationale for setting revised carbapenem breakpoints:

- The clinical effectiveness of carbapenem treatment of infections produced by isolates for which the carbapenem MIC or disk diffusion test results are within the intermediate range is uncertain due to lack of controlled clinical studies.
- Imipenem MICs for *Proteus* spp., *Providencia* spp., and *Morganella morganii* tend to be higher (eg, MICs in the intermediate or resistant range) than meropenem or doripenem MICs. These isolates may have elevated imipenem MICs by mechanisms other than production of carbapenemases.

donpene	ili Milos. Tricsc isolates il	lay have cic	vated inhip	CHCIII WIIC	23 by Incone	111131113 0	trici triari pre	Judellon	or carbaper	icitiases.	
В	Doripenem	10 µg	≥23	-	20–22	≤19	≤1	-	2	≥4	(32) Breakpoints are based on a dosage
				:		:			:	:	regimen of 500 mg administered every
				İ		•		i	1		8 h.
В	Ertapenem	10 µg	≥22	-	19–21	≤18	≤0.5	-	1	≥2	(33) Breakpoints are based on a dosage
				į	ļ	į		į	i	į	regimen of 1 g administered every 24 h.
В	Imipenem	10 µg	≥23	-	20–22	≤19	≤1	-	2	≥4	(34) Breakpoints are based on a dosage
				Ì	ļ	Ì		į	i	•	regimen of 500 mg administered every
				-	-	ŀ		į	-	•	6 h or 1 g every 8 h.

Table 2A. Enterobacterales (Continued)

	toropaotoraioo (o										
				•	ategories r Breakpoi		lı lı	•	re Categories Breakpoints,		
Test/Report	Antimicrobial	Disk		nearest v	whole mm				μg/mL		
Group	Agent	Content	S	SDD	I	R	S	SDD	I	R	Comments
CARBAPENE	MS (Continued)										
В	Meropenem	10 µg	≥23	-	20–22	≤19	≤1	-	2	≥4	(35) Breakpoints are based on a dosage regimen of 1 g administered every 8 h.
LIBOREDTINE	9										

LIPOPEPTIDES

(36) 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.

0	species are intrinsical Colistin or		_	_	_	_	-	-	≤2^	≥4	(38) Colistin (methanesulfonate)
Ü	polymyxin B		_	<u>-</u>	-	-	-	_	<u>≤2</u>	≥4	should be given with a loading do and maximum renally adjusted doses (see International Consens Guidelines ⁸).
											(39) Polymyxin B should be given with a loading dose and maximus recommended doses (see International Consensus Guidelines ⁸).
											(40) When colistin or polymyxin I given systemically, neither is like to be effective for pneumonia.
											(41) For colistin, broth microdilut CBDE, and CAT MIC methods are acceptable. For polymyxin B, bromicrodilution is the only approve method. Disk diffusion and gradi diffusion methods should not be performed (see Table 3D).
NOGLYC	OSIDES										
WARNIN	G: For Salmonella spp.	and <i>Shigella</i> s	pp., amin	oglycosid	es may app	ear activ	ve in vitro	but are no	ot effective cli	nically and s	should not be reported as susceptible.
Α	Gentamicin	10 μg	≥15	_	13–14^	≤12	≤4	_	8^	≥16	
Α	Tobramycin	10 μg	≥15	-	13–14 ^	≤12	≤4	_	8^	≥16	
В	Amikacin	30 μg	≥17	-	15–16 ^	≤14	≤16	_	32^	≥64	
0	Kanamycin	30 μg	≥18	-	14–17^	≤13	≤16	-	32^	≥64	
0	Netilmicin	30 μg	≥15	_	13–14^	≤12	≤8	-	16 ^	≥32	
0	Streptomycin	10 ug	≥15		12-14^	≤11	_	_	_	_	

Test/Report	Antimicrobial	Disk		Diamet	Categories er Breakpo whole mm	oints,	ı		Categories eakpoints, g/mL		
Group	Agent	Content	S	SDD	I	R	S	SDD	1	R	Comments
MACROLIDES											
Inv.	Azithromycin	15 μg	≥13	-	-	≤12	≤16	_	-	≥32	(43) S. enterica ser. Typhi only: breakpoints are based on MIC distribution data and limited clinical data For S. flexneri and S. sonnei, see Appendix G, Table G1.
TETRACYCLI	INES										
	s that are susceptible to ay be susceptible to dox				susceptible	to doxycy	cline an	d minocyclir	ne. Howeve	r, some org	anisms that are intermediate or resistant to
С	Tetracycline	30 μg	≥15	_	12–14	≤11	≤4	_	8	≥16	
0	Doxycycline	30 μg	≥14	_	11–13	≤10	≤4	_	8	≥16	
0	Minocycline	30 μg	≥16	-	13–15	≤12	≤4	_	8	≥16	
QUINOLONES	AND FLUOROQUINO	LONES for E	nterobact	erales ex	cept Salm	onella sp	p. (Plea	se refer to (Glossary I.		
В	Ciprofloxacin	5 µg	≥26	-	22–25^	≤21	≤	_	0.5^	≥1	(45) Breakpoints for ciprofloxacin are
В	Levofloxacin	5 µg	≥21		17–20 ^	≤16	0.25	-	1^	≥2	based on a dosage regimen of
							≤ 0.5				400 mg IV or 500 mg orally administered every 12 h.
											(46) Breakpoints for levofloxacin are based on a dosage regimen of 750 mg administered every 24 h.
0	Cinoxacin	100 µg	≥19	-	15–18 ^	≤14	≤16	_	32^	≥64	See comment (29).
0	Enoxacin	10 µg	≥ 18	_	15–17^	≤ 14	≤2	_	4^	≥8	See comment (29).
0	Gatifloxacin	5 µg	≥ 18	_	15–17^	≤ 14	≤2	_	4^	≥8	
0	Gemifloxacin	5 μg	≥ 20	-	16–19	≤ 15	≤ 0.25	_	0.5	≥1	(47) For testing and reporting of K. pneumoniae only.
0	Grepafloxacin	5 µg	≥18	-	15–17	≤14	≤1	_	2	≥4	
	Lomefloxacin	10 µg	≥22	-	19–21^	≤18	≤2	_	4^	≥8	
0											
0	Nalidixic acid	30 μg 10 μg	≥19 ≥17	_	14–18 13–16	≤13 ≤12	≤16 ≤4	_	_	≥32 ≥16	See comment (29). See comment (29).

QUINOLONES AND FLUOROQUINOLONES for Salmonella spp. (Please refer to Glossary I.)

5 ua

5 µg

≥16

≥19

(48) For testing and reporting of Salmonella spp. (including S. enterica ser. Typhi and S. enterica ser. Paratyphi A-C). Routine susceptibility testing is not indicated for nontyphoidal Salmonella spp. isolated from intestinal sources.

≤12

≤2

4^

≥8

≥8

(49) The preferred test for assessing fluoroquinolone 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. If a ciprofloxacin, levofloxacin, or ofloxacin MIC or ciprofloxacin disk diffusion test cannot be done, pefloxacin disk diffusion may be used as surrogate test to predict ciprofloxacin susceptibility.

(50) No single test detects resistance resulting from all possible fluoroquinolone resistance mechanisms that have been identified in Salmonella spp.

13-15^

16-18^

0

Inv.

Ofloxacin

Fleroxacin

should not be performed.

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Table 2A. Enterobacterales (Continued)

| Interpretive Categories and Zone Diameter Breakpoints, Hill Breakpoints, Content S SDD I R S S

Test/Report	Antimicrobial	Disk			r Breakpoi vhole mm	nts,		MIC	Breakpoints, μg/mL		
Group	Agent	Content	S	SDD	1	R	S	SDD	<u> </u>	R	Comments
	S AND FLUOROQUING	LONES for S	Salmonell	a spp. (F	lease refer	r to Glo	ssary I.) (Continue	ed)	•	2 2 2
В	Ciprofloxacin	5 µg	≥31	: -	21–30^	: ≤20	≤0.06	: -	0.12-0.5 ^	≥1	(51) Isolates of Salmonella spp. that test
В	Levofloxacin	_	_	_	- -	-	≤0.12	-	0.25–1^	≥2	not susceptible to ciprofloxacin, levofloxacin, ofloxacin, or pefloxacin may be associated with clinical failure or delayed response in fluoroquinolone-treated patients with salmonellosis.
0	Ofloxacin	_	_	-	-	-	≤0.12	-	0.25-1^	≥2	
lnv.	Pefloxacin (surrogate test for ciprofloxacin)	5 μg	≥24		-	≤23	_		-	-	(52) Report results as ciprofloxacin susceptible or resistant based on the pefloxacin test result. Pefloxacin will not detect resistance in Salmonella spp. due to aac(6')-lb-cr. Pefloxacin disks are not available in the United States. See comment (50).
FOLATE PAT	HWAY ANTAGONISTS	3									
В	Trimethoprim- sulfamethoxazole	1.25/ 23.75 µg	≥16	_	11–15	≤10	≤2/38	-	-	≥4/76	See general comment (2).
U	Sulfonamides	250 or 300 μg	≥17	-	13–16	≤12	≤256	-	-	≥512	(53) Sulfisoxazole can be used to represen any of the currently available sulfonamide preparations.
U	Trimethoprim	5 µg	≥16	; –	11–15	≤10	≤8	. –	: - :	≥16	
PHENICOLS											
С	Chloramphenicol	30 µg	≥18	-	13–17	≤12	≤8	-	16	≥32	(54) Not routinely reported on isolates from the urinary tract.
FOSFOMYCII	NS										
U	Fosfomycin	200 µg	≥16	_	13–15	≤12 :	≤64	_	128	≥256	 (55) Disk diffusion and MIC breakpoints apply only to <i>E. coli</i> urinary tract isolates and should not be extrapolated to other species of Enterobacterales. (56) The 200-μg fosfomycin disk contains 50 μg of glucose-6-phosphate.
											(57) The only approved MIC method for testing is agar dilution using agar media supplemented with 25 μg/mL of glucose-6-phosphate. Broth dilution MIC testing

Table 2A. Enterobacterales (Continued)

Test/Report	Antimicrobial	Disk	Zone [Diamet	Categories er Breakpo whole mm	ints,	In		Bre	Categori akpoint /mL		
Group	Agent	Content	S	SDD	1	R	S	SDE) :	ı	R	Comments
NITROFURAN:	S											
U	Nitrofurantoin	300 µg	≥17	_	15–16	≤14	≤32	. –	i	64	≥128	

Abbreviations: ATCC[®], American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; **CAT, colistin agar test; CBDE, colistin broth disk elution;** eCIM, EDTA-modified carbapenem inactivation method; ESBL, extended-spectrum β-lactamase; I, intermediate; IV, intravenous; mCIM, modified carbapenem inactivation method; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; PK-PD, pharmacokinetic-pharmacodynamic; QC, quality control; R, resistant; S, susceptible; SDD, susceptible-dose dependent; UTI, urinary tract infection.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2A

- Hackel MA, Tsuji M, Yamono Y, Echols R, Karlowsky JA, Sahm DF. Reproducibility of broth microdilution MICs for the novel siderophore cephalosporin, cefiderocol, determined using iron-depleted cation-adjusted Mueller-Hinton broth. *Diagn Microbiol Infect Dis.* 2019;94(4):321-325.
- ² CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 3 CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ⁴ Perrott J, Mabasa VH, Ensom MH. Comparing outcomes of meropenem administration strategies based on pharmacokinetic and pharmacodynamic principles: a qualitative systematic review. *Ann Pharmacother*. 2010;44(3):557-564.
- ⁵ Cirillo I, Vaccaro N, Turner K, Solanki B, Natarajan J, Redman R. Pharmacokinetics, safety, and tolerability of doripenem after 0.5-, 1-, and 4-hour infusions in healthy volunteers. *J Clin Pharmacol*. 2009;49(7):798-806.
- Sakka SG, Glauner AK, Bulitta JB, et al. Population pharmacokinetics and pharmacodynamics of continuous versus short-term infusion of imipenem-cilastatin in critically ill patients in a randomized, controlled trial. *Antimicrob Agents Chemother*. 2007;51(9):3304-3310.
- Peleg AY, Hooper DC. Hospital-acquired infections due to gram-negative bacteria. *N Engl J Med*. 2010;362(19):1804-1813.
- Tsuji BT, Pogue JM, Zavaxcki AP, et al. International consensus guidelines for the optimal use of the polymyxins: endorsed by the American College of Clinical Pharmacy (ACCP), European Society of Clinical Microbiology and Infectious Diseases (ESCMID), Infectious Diseases Society of America (IDSA), International Society for Anti-Infective Pharmacology (ISAP), Society of Critical Care Medicine (SCCM), and Society of Infectious Diseases Pharmacists (SIDP). Pharmacotherapy. 2019;39(1):10-39.

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Table 2B-1. Zone Diameter and MIC Breakpoints for Pseudomonas aeruginosa

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB; iron-depleted CAMHB for

cefiderocol (see Appendix I)¹

Agar dilution: MHA

Inoculum: Broth culture method or colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35°C±2°C; ambient air

Disk diffusion: 16–18 hours Dilution methods: 16–20 hours **Routine QC Recommendations** (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Pseudomonas aeruginosa ATCC®a 27853

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02,² Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide³). Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth.
- (2) The susceptibility of *P. aeruginosa* isolated from patients with cystic fibrosis can be reliably determined by disk diffusion or dilution methods but may need extended incubation for up to 24 hours before reporting as susceptible.
- (3) P. aeruginosa may develop resistance during prolonged therapy with all antimicrobial agents. Therefore, isolates that are initially susceptible may become resistant within 3 to 4 days after initiation of therapy. Testing of repeat isolates may be warranted.
- (4) The dosage regimens shown in the comments column below are those necessary to achieve plasma drug exposures (in adults with normal renal and hepatic functions) on which breakpoints were derived. When implementing new breakpoints, it is strongly recommended that laboratories share this information with infectious diseases practitioners, pharmacists, pharmacy and therapeutics committees, infection **prevention** committees, and the antimicrobial stewardship team.
- (5) Intermediate ranges denoted with a "A" for the applicable antimicrobial agents in the drug groups in Tables 2 are based on the known ability of these agents to concentrate in the urine; some agents may also have the potential to concentrate at other anatomical sites (eg, epithelial lining).

Table 2B-1. Pseudomonas aeruginosa (Continued)

Test/Report	Antimicrobial	Disk	Zone Di	tive Catego ameter Brea arest whole	akpoints,		pretive Categor MIC Breakpoin µg/mL		
Group	Agent	Content	S	ı	R	s	ı	R	Comments
PENICILLINS 1									
0	Piperacillin	100 μg	≥21	15–20 ^	≤14	≤16	32–64 ^	≥128	(6) Breakpoints for piperacillin (alone or with tazobactam) are based on a piperacillin dosage regimen of at least 3 g administered every 6 h.
β-LACTAM C	OMBINATION AGENTS								
А	Piperacillin-tazobactam	100/10 μg	≥21	15–20 ^	≤14	≤16/4	32/4–64/4^	≥128/4	(7) Breakpoints for piperacillin (alone or with tazobactam) are based on a piperacillin dosage regimen of at least 3 g administered every 6 h.
В	Ceftazidime-avibactam	30/20 μg	≥21	_	≤20	≤8/4	_	≥16/4	(8) Breakpoints are based on a dosage regimen of 2.5 g (2 g ceftazidime+0.5 g avibactam) administered every 8 h over 2 h.
В	Ceftolozane-tazobactam	30/10 μg	≥21	17–20^	≤16	≤4/4	8/4^	≥ 16/4	(9) Breakpoints are based on a dosage regimen of 1.5 g administered every 8 h.
0	Ticarcillin-clavulanate	75/10 μg	≥24	16–23 ^	≤15	≤16/2	32/2–64/2 ^	≥128/2	(10) Breakpoints for ticarcillin (alone or with clavulanate) are based on a ticarcillin dosage regimen of at least 3 g administered every 6 h.
CEPHEMS (P.	ARENTERAL) (Including ce	phalosporins	I, II, III, an	d IV. Please	refer to G	lossary I.)			
Α	Ceftazidime	30 µg	≥18	15 – 17 ^	≤14	≤8	16^	≥32	(11) Breakpoints are based on a dosage regimen of 1 g administered every 6 h or 2 g administered every 8 h.
В	Cefepime	30 μg	≥18	15 – 17 ^	≤14	≤8	16^	≥32	(12) Breakpoints are based on a dosage regimen of 1 g administered every 8 h or 2 g administered every 12 h.
Inv.	Cefiderocol	30 µg	≥18	13–17^	≤12	≤4	8^	≥16	(13) Breakpoints are based on a dosage regimen of 2 g every 8 h administered over 3 h.
MONOBACTA	AMS						,		•
В	Aztreonam	30 μg	≥22	16–21 ^	≤15	≤8	16 ^	≥32	(14) Breakpoints are based on a dosage regimen of 1 g administered every 6 h or 2 g administered every 8 h.

Table 2B-1. Pseudomonas aeruginosa (Continued)

Test/Report	Antimicrobial	Disk	Zone D	retive Catego Diameter Bre earest whole	akpoints,		etive Cate IIC Breakpo µg/mL		
Group	Agent	Content	S	I	R	S	1	R	Comments
CARBAPENE	MS								
В	Doripenem	10 μg	≥19	16–18 ^	≤15	≤2	4^	≥8	(15) Breakpoints for doripenem are based on a dosage regimen of 500 mg administered every 8 h.
В	Imipenem	10 μg	≥19	16–18 ^	≤15	≤2	4^	≥8	(16) Breakpoints for imipenem are based on a dosage regimen of 1 g administered every 8 h or 500 mg administered every 6 h.
В	Meropenem	10 μg	≥19	16–18 ^	≤15	≤2	4^	≥8	(17) Breakpoints for meropenem are based on a dosage regimen of 1 g administered every 8 h.
LIPOPEPTIDE	S		•						

(18) 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.

0	Colistin or	_	_	: -	: -	_	: ≤2	: ≥4	(19) Colistin (methanesulfonate) should be
	polymyxin B	_	_	: –	: -	_	≤2	: >4	given with a loading dose and maximum
	' ' '			:	:		:	: -:	renally adjusted doses (see International
				:	:		:	:	Consensus Guidelines ⁴).
				:	:		:	:	Consensus Guidennes J.
				:	:			:	(00) Delements Delement to the selection with a
				:	:			:	(20) Polymyxin B should be given with a
				-	:			:	loading dose and maximum recommended
					1				doses (see International Consensus
					:				Guidelines4).
				:	:		:	:	,
				:	:		:	:	(21) When colistin or polymyxin B is given
				:	:		:	:	systemically, neither is likely to be
				:	:		:	:	
				:	:			:	effective for pneumonia.
				:	:			:	
				:				:	(22) For colistin, broth microdilution,
				-	-		-		CBDE, and CAT MIC methods are
				:					acceptable. For polymyxin B, broth
					:				microdilution is the only approved method.
				:	:		:	:	Disk diffusion and gradient diffusion
				:	:		:	:	
				:	:		:	:	methods should not be performed (see
				:	:		-	:	Table 3D).

Table 2B-1. Pseudomonas aeruginosa (Continued)

Test/Report	Antimicrobial	Disk	Zone D	retive Catego Diameter Bre earest whole	akpoints,		etive Catego IC Breakpoi µg/mL		
Group	Agent	Content	S	1	R	S	I	R	Comments
AMINOGLYC	OSIDES								
Α	Gentamicin	10 μg	≥15	13–14^	≤12	≤4	8^	≥16	
Α	Tobramycin	10 μg	≥15	13–14^	≤12	≤4	8^	≥16	
В	Amikacin	30 μg	≥17	15–16 ^	≤14	≤16	32^	≥64	
0	Netilmicin	30 μg	≥15	13–14^	≤12	≤8	16 ^	≥32	
FLUOROQUI	INOLONES								
В	Ciprofloxacin	5 μg	≥25	19–24 ^	≤18	≤ 0.5	1^	≥2	(23) Breakpoints are based on a dosage regimen of 400 mg IV administered every 8 h
В	Levofloxacin	5 μg	≥22	15–21 ^	≤14	≤ 1	2^	≥4	(24) Breakpoints are based on a dosage regimen of 750 mg administered every 24 h.
0	Lomefloxacin	10 μg	≥.22	19–21 ^	≤18	≤ 2	4^	≥8	(25) For testing and reporting of urinary tract isolates only.
0	Norfloxacin	10 μg	≥17	13–16	≤12	≤ 4	8	≥16	See comment (25).
0	Ofloxacin	5 μg	≥16	13–15 ^	≤12	≤ 2	4^	≥8	
0	Gatifloxacin	5 μg	≥18	15–17 ^	≤14	≤ 2	4^	≥8	

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; **CAT**, **colistin agar test**; **CBDE**, **colistin broth disk elution**; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; **PK-PD**, **pharmacokinetic-pharmacodynamic**; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2B-1

- Hackel MA, Tsuji M, Yamono Y, Echols R, Karlowsky JA, Sahm DF. Reproducibility of broth microdilution MICs for the novel siderophore cephalosporin, cefiderocol, determined using iron-depleted cation-adjusted Mueller-Hinton broth. *Diagn Microbiol Infect Dis.* 2019;94(4):321-325.
- ² CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ³ CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- Tsuji BT, Pogue JM, Zavaxcki AP, et al. International consensus guidelines for the optimal use of the polymyxins: endorsed by the American College of Clinical Pharmacy (ACCP), European Society of Clinical Microbiology and Infectious Diseases (ESCMID), Infectious Diseases Society of America (IDSA), International Society for Anti-Infective *Pharmacology* (ISAP), Society of Critical Care Medicine (SCCM), and Society of Infectious Diseases Pharmacists (SIDP). *Pharmacotherapy*. 2019;39(1):10-39.

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Table 2B-2. Zone Diameter and MIC Breakpoints for Acinetobacter spp.

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB; iron-depleted CAMHB for

cefiderocol (see Appendix I)1

Agar dilution: MHA

Inoculum: Broth culture method or colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35°C±2°C; ambient air; 20–24 hours, all methods

Routine QC Recommendations (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Escherichia coli ATCC®a 25922 (for tetracyclines and trimethoprim-

sulfamethoxazole)

Pseudomonas aeruginosa ATCC® 27853

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comment

(1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02,² Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide³). Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.

Table 2B-2. Acinetobacter spp. (Continued)

14510 25 2.	Acinetobacter Spp.	(Commuca)	Interp	retive Cate	gories				
Test/Report	Antimicrobial	Disk	Е	and one Diamet Breakpoints rest whole	s,		retive Categor IIC Breakpoin µg/mL		
Group	Anumicrobiai	Content	S	1	R	s		R	Comments
PENICILLINS		Content	J	<u> </u>	<u> </u>	3	<u> </u>	<u>, K</u>	Comments
O	Piperacillin	100 μg	≥21	18–20	≤17	≤16	32–64	≥128	
β-LACTAM C	OMBINATION AGENTS								
Α	Ampicillin-sulbactam	10/10 μg	≥15	12–14	≤11	≤8/4	16/8	≥32/16	
В	Piperacillin-tazobactam	100/10 μg	≥21	18–20	≤17	≤16/4	32/4–64/4	≥128/4	
0	Ticarcillin-clavulanate	75/10 μg	≥20	15–19	≤14	≤16/2	32/2-64/2	≥128/2	
CEPHEMS (P.	ARENTERAL) (Including		, II, III, an	d IV. Pleas	e refer to	Glossary I	.)		
Α	Ceftazidime	30 μg	≥18	15–17	≤14	≤8	16	≥32	
В	Cefepime	30 μg	≥18	15–17	≤14	≤8	16	≥32	
В	Cefotaxime	30 μg	≥23	15–22	≤14	≤8	16–32	≥64	
В	Ceftriaxone	30 μg	≥21	14–20	≤13	≤8	16–32	≥64	
Inv.	Cefiderocol	30 μg	≥15	11–14	≤10	≤4	8	≥16	(2) Breakpoints are based on a dosage regimen of 2 g every 8 h administered over 3 h.
CARBAPENE	MS							•	, ,
А	Doripenem	10 μg	≥18	15–17	≦14	≤2	4	≥8	(3) Breakpoints for doripenem are based on a dosage regimen of 500 mg administered every 8 h.
Α	Imipenem	10 μg	≥22	19–21	≤18	≤2	4	≥8	(4) Breakpoints for imipenem are based on a dosage regimen of 500 mg administered every 6 h.
Α	Meropenem	10 μg	≥18	15–17	≤14	≤2	4	≥8	(5) Breakpoints for meropenem are based on a dosage regimen of 1 g administered every 8 h or 500 mg administered every 6 h.

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Test/Report	Antimicrobial	Disk	Zo	retive Cate and one Diamet reakpoints rest whole	er s,		etive Catego IC Breakpoii µg/mL		
Group	Agent	Content	S	I	R	S	ı	R	Comments
are strongly p	: Clinical and PK-PD dat preferred. Colistin and precommended. Colistin or	a demonstrate colymyxin B shot	colistin and uld be use	d polymyxi d in combi	n B have nation wi	limited clir	nical efficacy nore active a ≤2 <2	/, even if an antimicrobia	intermediate result is obtained. Alternative ager al agents. Consultation with an infectious diseas (7) Colistin (methanesulfonate) should be give with a loading dose and maximum renally
	polymyxin B						52	24	adjusted doses (see International Consensus Guidelines ⁴). (8) Polymyxin B should be given with a loading dose and maximum recommended doses (see International Consensus Guidelines ⁴). (9) When colistin or polymyxin B is given systemically, the drug is unlikely to be effective for pneumonia. (10) The only approved MIC method is broth microdilution. CBDE, CAT, disk diffusion, and gradient diffusion should not be performed. (11) Applies to A. baumannii complex only.
AMINOGLYC	OSIDES								
Α	Gentamicin	10 μg	≥15	13–14	≤12	≤4	8	≥16	
Α	Tobramycin	10 μg	≥15	13–14	≤12	≤4	8	≥16	
В	Amikacin	30 μg	≥17	15–16	≤14	≤16	32	≥64	
0	Netilmicin	_	_	-	-	≤8	16	≥32	
					tible to do ≤9 ≤12	exycycline a ≤4 ≤4	nd minocyclir	ne. However	, some organisms that are intermediate or resistant
U	Tetracycline	30 μg	≥15	12–14	<u> </u>	<u> </u>	8	≥16	
ŭ		υ μ <u>ο</u>		14-14	<u>: = 11</u>	<u>≥</u> •• '	U	: = 10	
LUOROOUII	10LUITLU				•	1		1	
	Ciprofloyacin	5	> 21	16_20	. /15	/1	?		
FLUOROQUIN A A	Ciprofloxacin Levofloxacin	5 μg 5 μg	≥21 ≥17	16–20 14–16	≤15 ≤13	≤1 ≤2	2 4	≥4 ≥8	

Table 2B-2 Acinetobacter spp. (Continued)

I GOIO ED E.	Acinetobacter app.	(Gontiniaga)							
			Z	Interpretive Categories and Zone Diameter Breakpoints,			etive Cate C Breakpo	gories and pints,	
Test/Report	Antimicrobial	Disk	nearest whole mm				μg/mL		
Group	Agent	Content	S	1	R	S	ı	R	Comments
FOLATE PAT	HWAY ANTAGONISTS								
В	Trimethoprim-	1.25/23.75 μg	≥ 16	11–1	5 ≤10	≤2/38	_	≥4/76	
	sulfamethoxazole	. •		į	į	į		į	

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; CAT, colistin agar test; CBDE, colistin broth elution test; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; PK-PD, pharmacokinetic-pharmacodynamic; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2B-2

- Hackel MA, Tsuji M, Yamono Y, Echols R, Karlowsky JA, Sahm DF. Reproducibility of broth microdilution MICs for the novel siderophore cephalosporin, cefiderocol, determined using iron-depleted cation-adjusted Mueller-Hinton broth. Diagn Microbiol Infect Dis. 2019;94(4):321-325.
- CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- Tsuji BT, Pogue JM, Zavaxcki AP, et al. International consensus guidelines for the optimal use of the polymyxins: endorsed by the American College of Clinical Pharmacy (ACCP), European Society of Clinical Microbiology and Infectious Diseases (ESCMID), Infectious Diseases Society of America (IDSA), International Society for Anti-Infective Pharmacology (ISAP), Society of Critical Care Medicine (SCCM), and Society of Infectious Diseases Pharmacists (SIDP). Pharmacotherapy. 2019;39(1):10-39.

Table 2B-3. Zone Diameter and MIC Breakpoints for Burkholderia cepacia complex

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB Agar dilution: MHA

Inoculum: Broth culture method or colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35°C±2°C; ambient air; 20–24 hours, all methods

Routine QC Recommendations (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Escherichia coli ATCC^{®a} 25922 (for chloramphenicol, minocycline, and trimethoprim-sulfamethoxazole)

Pseudomonas aeruginosa ATCC® 27853

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comment

(1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02,¹ Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide²). Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.

		5	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm Interpretive Categories and MIC Breakpoints, μg/mL						
Test/Report Group	Antimicrobial Agent	Disk Content	s	; ,	R	s	,	R	Comments
	MBINATION AGENTS	Content			, IX			<u>, ix</u>	Comments
0	Ticarcillin-clavulanate	_	_	: -	_	≤16/2	32/2-64/2	: ≥128/2	
CEPHEMS (PA	RENTERAL) (Including o	ephalosporins I,	II, III, and I	IV. Please re	fer to Glo	ossary I.)	-		
В	Ceftazidime	30 μg	≥21	18–20	≤17	≤8	16	≥32	
CARBAPENEN	ns .								
Α	Meropenem	10 μg	≥20	16–19	≤15	≤4	8	≥16	
TETRACYCLIN	IES								
В	Minocycline	30 μg	≥19	15–18	≤14	≤4	8	≥16	
FLUOROQUIN	OLONES								
Α	Levofloxacin	-	_	-	-	≤2	4	≥8	
FOLATE PATH	IWAY ANTAGONISTS								
Α	Trimethoprim- sulfamethoxazole	1.25/23.75 μg	≥16	11–15	≤10	≤2/38	-	≥4/76	
PHENICOLS									
С	Chloramphenicol	_	_	-	-	≤8	16	≥32	(2) Not routinely reported on isolates from the urinary tract.

Abbreviations: ATCC[®], American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. $ATCC^{\otimes}$ is a registered trademark of the American Type Culture Collection.

References for Table 2B-3

- ¹ CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

M100, 30th ed

Table 2B-4. Zone Diameter and MIC Breakpoints for Stenotrophomonas maltophilia

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB; iron-depleted CAMHB for

cefiderocol (see Appendix I)1

Agar dilution: MHA

Inoculum: Broth culture method or colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35°C±2°C; ambient air; 20–24 hours, all methods

Routine QC Recommendations (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Escherichia coli ATCC®a 25922 (for chloramphenicol, minocycline, and

trimethoprim-sulfamethoxazole)

Pseudomonas aeruginosa ATCC® 27853

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comment

(1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02,² Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide³). Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.

Test/Report	Antimicrobial	Disk	2	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm			etive Categorie Breakpoints µg/mL		
Group	Agent	Content	S	1	R	S	1	R	Comments
β-LACTAM CC	MBINATION AGENTS								
0	Ticarcillin-clavulanate	_	_	-	-	≤16/2	32/2-64/2	≥128/2	
CEPHEMS (PA	RENTERAL) (Including cept	nalosporins I, II	, III, and	IV. Please re	fer to G	lossary I.)			
В	Ceftazidime	_	_	-	_	≤8	16	≥32	
Inv.	Cefiderocol	30 μg	≥17	13–16	≤12	≤ 4	8	≥ 16	(2) Breakpoints are based on a dosage regimen of 2 g every 8 h administered over 3 h.
TETRACYCLI	NES								
Α	Minocycline	30 μg	≥19	15–18	≤14	≤4	8	≥16	
FLUOROQUIN	OLONES								
Α	Levofloxacin	5 μg	≥17	14–16	≤13	≤2	4	≥8	
FOLATE PATH	WAY ANTAGONISTS								
Α	Trimethoprim- sulfamethoxazole	1.25/23.75 μg	≥16	11–15	≤10	≤2/38		≥4/76	
PHENICOLS									
С	Chloramphenicol	_	-	_	_	≤8	16	≥32	(3) Not routinely reported on isolates from the urinary tract.

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2B-4

- Hackel MA, Tsuji M, Yamono Y, Echols R, Karlowsky JA, Sahm DF. Reproducibility of broth microdilution MICs for the novel siderophore cephalosporin, cefiderocol, determined using iron-depleted cation-adjusted Mueller-Hinton broth. *Diagn Microbiol Infect Dis.* 2019;94(4):321-325.
- ² CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 3 CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

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Table 2B-5. MIC Breakpoints for Other Non-Enterobacterales (Refer to General Comment 1)

Testing Conditions

Medium: Broth dilution: CAMHB Agar dilution: MHA

Inoculum: Broth culture method or colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35°C±2°C; ambient air; 16–20 hours

Routine QC Recommendations (see Table 5A-1 for acceptable QC ranges)

Escherichia coli ATCC^{®a} 25922 (for chloramphenicol, tetracyclines, sulfonamides, and trimethoprim-sulfamethoxazole)
Pseudomonas aeruginosa ATCC[®] 27853

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) Other non-Enterobacterales include *Pseudomonas* spp. and other nonfastidious, glucose-nonfermenting, gram-negative bacilli but exclude *P. aeruginosa, Acinetobacter* spp., *B. cepacia* complex, and *S. maltophilia* (refer to Tables 2B-2, 2B-3, and 2B-4, respectively). Recommendations for testing and reporting *Aeromonas hydrophila* complex, *Burkholderia mallei*, *Burkholderia pseudomallei*, and *Vibrio* spp. (including *V. cholerae*) are found in CLSI document M45.¹
- (2) For other non-Enterobacterales, the disk diffusion method has not been systematically studied. Therefore, for this organism group, disk diffusion testing is not recommended.

			Zone Dia	tive Catego ameter Brea arest whole	kpoints,	Interpretive Categories and MIC Breakpoints, µg/mL			
Test/Report Group	Antimicrobial Agent	Disk Content	s	1	R	s	1	R	Comments
PENICILLINS									
0	Piperacillin	_	_	_	_	≤16	32–64	≥128	
β-LACTAM CC	MBINATION AGENTS							•	
В	Piperacillin-tazobactam	_	_	-	_	≤16/4	32/4-64/4	≥ 128/4	
0	Ticarcillin-clavulanate	_	_	_	-	≤16/2	32/2-64/2	≥128/2	
CEPHEMS (PA	RENTERAL) (Including ceph	alosporins I,	II, III, and I	V. Please re	fer to Glos	sary I.)			
Α ,	Ceftazidime		<u> </u>	_	_	≤8	16	≥32	
В	Cefepime	_	_	_	_	≤8	16	≥32	
С	Cefotaxime	_	_	_	_	≤8	16–32	≥64	
С	Ceftriaxone	_	_	-	_	≤8	16–32	≥64	
0	Cefoperazone	_	_	_	_	≤16	32	≥64	
0	Ceftizoxime	_	_	-	-	≤8	16–32	≥64	
0	Moxalactam	_	_	-	_	≤8	16–32	≥64	
MONOBACTA	MS								
В	Aztreonam	_	_		<u> </u>	≤8	16	≥32	
CARBAPENE									
В	Imipenem	_	-	_	_	≤4	8	≥16	
В	Meropenem	_	-	<u> </u>	-	≤4	8	≥16	
AMINOGLYCC			•	-					,
Α	Gentamicin	-	_	_	_	≤4	8	≥16	
Α	Tobramycin	-	_	-	_	≤4	8	≥16	
В	Amikacin	-	_	_	<u> </u>	≤16	32	≥64	
0	Netilmicin	_	_	<u> </u>	<u> </u>	≤8	16	≥32	
tetracycline ma	that are susceptible to tetracycl y be susceptible to doxycycline	, minocycline,		1	1				organisms that are intermediate or resistant to
U	Tetracycline	_	_	<u> </u>	_	≤4	8	≥16	
0	Doxycycline	_	_	-	_	≤4	8	≥16	
0	Minocycline	_	_	<u> </u>	-	≤4	8	<u> ≥16</u>	
FLUOROQUIN		T T	1		·				T
B B	Ciprofloxacin	_	_	-	_	≤1	2	≥4	
	Levofloxacin	_	_		<u> </u>	≤2	4	≥8	
0	Gatifloxacin	_	-		_	≤2	4	≥8	
0	Lomefloxacin	_	_	-	_	≤2	4	≥8	105 1 11 11 11 11
0	Norfloxacin	-	-	-	-	≤4	8	≥16	(4) For testing and reporting of urinary tract isolates only.
Ο	Ofloxacin	_	_	-	-	≤2	4	≥8	

Table 2B-5. Other Non-Enterobacterales (Continued)

			Zone Di	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm Interpretive Catego MIC Breakpoin µg/mL					
Test/Report Group	Antimicrobial Agent	Disk Content	S	ı	R	S I R			Comments
FOLATE PATH	IWAY ANTAGONISTS								
В	Trimethoprim- sulfamethoxazole	-	_	-	-	≤2/38	-	≥4/76	
U	Sulfonamides	_	_	-	-	≤256	-	≥512	(5) Sulfisoxazole can be used to represent any of the currently available sulfonamide preparations.
PHENICOLS									<u> </u>
С	Chloramphenicol	-	_	_	_	≤8	16	≥32	(6) Not routinely reported on isolates from the urinary tract.

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

Reference for Table 2B-5

¹ CLSI. *Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently Isolated or Fastidious Bacteria.* 3rd ed. CLSI guideline M45. Wayne, PA: Clinical and Laboratory Standards Institute; 2016.

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Table 2C. Zone Diameter and MIC Breakpoints for Staphylococcus spp.

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB; CAMHB + 2% NaCl for oxacillin; CAMHB supplemented to 50 µg/mL calcium for daptomycin.

Agar dilution: MHA; MHA + 2% NaCl for oxacillin.

NOTE: Agar dilution has not been validated for daptomycin.

Inoculum: Colony suspension, equivalent to a 0.5 McFarland

standard

Incubation: 35°C±2°C; ambient air

Disk diffusion: 16–18 hours; 24 hours (for cefoxitin when testing *Staphylococcus* spp., excluding *S. aureus*, *S. lugdunensis*, *S. pseudintermedius*, and *S. schleiferi*) Dilution methods: 16–20 hours; 24 hours for oxacillin and

vancomycin

Testing at temperatures above 35°C may not detect methicillin (oxacillin)-resistant staphylococci (MRS).

Routine QC Recommendations (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Disk diffusion:

S. aureus ATCC®a 25923

Dilution methods:

S. aureus ATCC® 29213

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02, Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide²). Hold the Petri plate a few inches above a black background illuminated with reflected light, except for linezolid, which should be read with transmitted light (plate held up to light source). The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter. For linezolid, any discernible growth within the zone of inhibition is indicative of resistance to the respective agent.
- (2) For staphylococci when testing chloramphenicol, clindamycin, erythromycin, linezolid, tedizolid, and tetracycline by broth microdilution MIC, trailing growth can make end-point determination difficult. In such cases, read the MIC at the lowest concentration where the trailing begins. Tiny buttons of growth should be ignored (see M07,³ Figures 3 and 4). With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, read the end point at the concentration in which there is ≥80% reduction in growth compared with the control (see M07,³ Figure 5).
- (3) Routine testing of urine isolates of *Staphylococcus saprophyticus* is not advised, because infections respond to concentrations achieved in urine of antimicrobial agents commonly used to treat acute, uncomplicated UTIs (eg, nitrofurantoin, trimethoprim±sulfamethoxazole, or a fluoroquinolone).

Table 2C. Staphylococcus spp. (Continued)

- (4) Historically, resistance to the penicillinase-stable penicillins (see Glossary I) has been referred to as "methicillin resistance" or "oxacillin resistance." MRSA are strains of *S. aureus* that express *mecA*, *mecC*, or another mechanism of methicillin (oxacillin) resistance, such as changes in affinity of penicillin-binding proteins for oxacillin (modified *S. aureus* strains).
- (5) Most methicillin (oxacillin) resistance is mediated by mecA, encoding PBP2a (also called PBP2'). Isolates that test positive for mecA or PBP2a should be reported as methicillin (oxacillin) resistant (see Appendix H).

Detection of methicillin (oxacillin) resistance in staphylococci is achieved by using specific methods as listed in Table 2C and further described in Table 3F.

	Meth	ods for Detection of Me	thicillin (Oxacillin)-Res	sistant Staphylococcus	s spp.
Organism	Cefoxitin MIC	Cefoxitin disk diffusion	Oxacillin MIC	Oxacillin disk diffusion	Oxacillin salt agar
S. aureus	Yes (16–20 h)	Yes (16–18 h)	Yes (24 h)	No	Yes (24 h)
S. lugdunensis	Yes (16–20 h)	Yes (16–18 h)	Yes (24 h)	No	No
S. epidermidis	No	Yes (16–18 h)	Yes (24 h)	Yes (16–18 h)	No
S. pseudintermedius	No	No	Yes (24 h)	Yes (16–18 h)	No
S. schleiferi	No	No	Yes (24 h)	Yes (16–18 h)	No
Other <i>Staphylococcus</i> spp. (not listed above)	No	Yes ^a (24 h)	Yes ^a (24 h)	No	No

Abbreviations: h, hour(s); MIC, minimal inhibitory concentration; MRS, methicillin (oxacillin)-resistant staphylococci; PBP2a, penicillin-binding protein 2a.

Mechanisms of **methicillin** (oxacillin) resistance other than *mecA* are rare and include a novel *mecA* homologue, *mecC*.⁴ MICs for strains with *mecC* are typically cefoxitin resistant and oxacillin susceptible; *mecC* resistance cannot be detected by tests directed at *mecA* or PBP2a.

- (6) MRS, as defined by cefoxitin or oxacillin testing, as appropriate to the species, are considered resistant to other β-lactam agents, ie, penicillins, β-lactam combination agents, cephems (with the exception of ceftaroline), and carbapenems. This is because most cases of documented MRS infections have responded poorly to β-lactam therapy or because convincing clinical data that document clinical efficacy for those agents have not been presented.
- (7) For tests for β-lactamase production, **methicillin** (oxacillin) resistance and *mecA*-mediated **methicillin** (oxacillin) resistance using cefoxitin, reduced susceptibility to vancomycin, **ICR**, and high-level mupirocin resistance (S. *aureus* only), refer to Tables **3E**, **3F**, **3G**, **3H**, and **3I**, respectively.

^a For isolates of "other *Staphylococcus* spp." from serious infections for which the oxacillin MICs are 0.5–2 μg/mL, testing for *mecA* or PBP2a should be considered (see comment [17]). Cefoxitin disk diffusion is not currently recommended.

M100, 30th ed

Table 2C. Staphylococcus spp. (Continued	Table 2C.	Staph	ylococcus	spp.	(Continued
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	, ,	• • •	,	
				Interpretive Categories and
		Staphylococcus		Zone Diameter Breakpoints, MIC Breakpoints,
Test/Report	Antimicrobial	spp.	Disk	nearest whole mm μg/mL
Group	Agent	Indications	Content	S SDD I R S SDD I R Comments
DENIIOU LINIA	OF LABUE DENI	OIL LINIO		

- PENICILLINASE-LABILE PENICILLINS
- (8) Penicillin-susceptible staphylococci are susceptible to other β-lactam agents with established clinical efficacy for staphylococcal infections (including both penicillinase-labile and penicillinase-stable agents; see Glossary I). Penicillin-resistant staphylococci are resistant to penicillinase-labile penicillins.
- (9) Penicillin should be used to test the susceptibility of all staphylococci to penicillinase-labile penicillins (see Glossary I). Penicillin-resistant strains of staphylococci produce β -lactamase. Perform a test(s) to detect β -lactamase production on staphylococci for which the penicillin MICs are \leq 0.12 μg/mL or zone diameters \geq 29 mm before reporting the isolate as penicillin susceptible. Rare isolates of staphylococci that contain genes 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 *blaZ* β -lactamase gene maybe considered. See Tables 3D and 3E.

A Penicillin All staphylococci 10 units ≥29 - - ≤28 ≤0.12 - ≥0.25 (10) For methicillin (oxacillin)-resistant staphylococci, report penicillin as resistant or do not report.

PENICILLINASE-STABLE PENICILLINS

- (11) Cefoxitin is tested as a surrogate for oxacillin for some species of *Staphylococcus*. 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 *mecA* negative or PBP2a negative or cefoxitin susceptible should be reported as **methicillin** (oxacillin) susceptible.
- (12) Oxacillin (or cefoxitin) results can be applied to the other penicillinase-stable penicillins (cloxacillin, dicloxacillin, methicillin, and nafcillin). For agents with established clinical efficacy and considering site of infection and appropriate dosing, methicillin (oxacillin)-susceptible staphylococci can be considered susceptible to:
- β-lactam combination agents (amoxicillin-clavulanate, ampicillin-sulbactam, piperacillin-tazobactam)
- Oral cephems (cefaclor, cefdinir, cephalexin, cefpodoxime, cefprozil, cefuroxime, loracarbef)
- Parenteral cephems including cephalosporins I, II, III, and IV (cefamandole, cefazolin, cefepime, cefmetazole, cefonicid, cefoperazone, cefotaxime, ceftizoxime, ceftriaxone, cefuroxime, ceftaroline, moxalactam)
- Carbapenems (doripenem, ertapenem, imipenem, meropenem)

Methicillin (oxacillin)-resistant staphylococci are resistant to all currently available β -lactam antimicrobial agents, with the exception of ceftaroline. Thus, susceptibility or resistance to a wide array of β -lactam antimicrobial agents may be deduced from testing only penicillin and either cefoxitin or oxacillin. Testing of other β -lactam agents, except ceftaroline, is not advised. See general comments (5) and (6).

Additional explanation on the use of cefoxitin for prediction of mecA-mediated methicillin (oxacillin) resistance can be found in Subchapter 3.12 of M07³ and Subchapter 3.9 of M02.1

Table 2C. Staphylococcus spp. (Continued)

Test/		Staphylococcus		Zone D	etive Car liameter earest wh	Break	points,	Interp	retive Cat IIC Break µg/m	point	es and S,	
Group		spp. Indications	Disk Content	s	SDD	ı	R	s	SDD	ı	R	Comments
		LE PENICILLINS (Cor	ntinued)		•		-	-	•			
A	Oxacillin	S. aureus and S. lugdunensis	_	-	-	_	-	≤2 (oxacillin)	-	-	≥4 (oxacillin)	(13) Oxacillin disk testing is not reliable for <i>S. aureus</i> and <i>S. lugdunensis</i> .
			30 µg cefoxitin (surrogate test for oxacillin)	≥ 22	-	-	≤21	≤4 (cefoxitin)	-	-	≥8 (cefoxitin)	that 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.
A	Oxacillin	S. epidermidis	1 μg oxacillin	≥18 (oxacillin)	-	_	≤17 (oxacillin)	≤0.25 (oxacillin)	-	-	≥0.5 (oxacillin)	and (12). See general comments (5) and (6) and comments (8), (11), and (12).
			30 µg cefoxitin (surrogate test for oxacillin)	≥25 (cefoxitin)	-	-	≤24 (cefoxitin)	-	-	-	-	(15) Cefoxitin MIC testing is not reliable for detecting <i>mecA</i> -mediated resistance in S. <i>epidermidis</i> .
		S. pseudintermedius and S. schleiferi	1 μg oxacillin	≥18	-	_	≤ 17	≤0.25	-	_	≥0.5	(16) Neither cefoxitin MIC nor cefoxitin disk tests are reliable for detecting mecA-mediated resistance in S. pseudintermedius and S. schleiferi.
						: 						See general comments (5) and (6) and comments (8), (11), and (12).

is based on a dosage regimen of

600 mg administered every 12 h.

(19) The breakpoint for SDD is based on a dosage of 600 mg every

8 h administered over 2 h.

M100, 30th ed

Table 2C. Staphylococcus spp. (Continued)

Test/
Report Antimicrobial Group Agent Indications Content S SDD I R S SDD I R

PENICIL INASE STAPLE PENICIL INS (Continued)

including MRSA

Test/		Staphylococcus			earest v		mm		μg/n	•	,	Comments
	Antimicrobial	spp.	Disk		:				:	:	1	
Group	Agent	Indications	Content	S	SDD	<u> </u>	R	S	SDD	<u>: L</u>	R	
PENIC	PENICILLINASE-STABLE PENICILLINS (Continued)											
A	Oxacillin	Other Staphylococcus spp., excluding S. aureus S. lugdunensis S. epidermidis S. pseudintermedius S. schleiferi	30 µg cefoxitin (surrogate test for oxacillin)	≥ 25 (cefoxitin)			≤ 24 (cefoxitin)	≤0.25 (oxacillin)			≥0.5 (oxacillin)	(17) Oxacillin MIC breakpoints may overcall resistance, and some isolates for which the oxacillin MICs are 0.5–2 µg/mL may be mecA negative. Isolates from serious infections for which oxacillin MICs are 0.5–2 µg/mL may be tested for mecA or for PBP2a. Isolates that test mecA or
CERH	EMS /DADENTI											PBP2a negative should be reported as methicillin (oxacillin) susceptible. See general comments (5) and (6) and comments (8), (11), and (12).
	EMS (PARENTI		1	1 -							:	
В	Ceftaroline	S. aureus,	30 µg	≥25	20–	- 1	≤19	≤1	2–4	-	≥8	(18) The breakpoint for susceptible

24

Table 2C	Staphylococcus spp.	(Continued)

Test/Report	Antimicrobial		Disk	Inte Zon	e Diame	Categorie ter Breakp t whole mr	oints,	Inte	MIC Bre	Categorie: eakpoints, g/mL		Comments
Group	Agent		Content	S	SDD	ı	R	s	SDD	1 1	R	
isolates of	ests should be pe S. aureus from va		e isolates,	nor does	s the test							not differentiate vancomycin-susceptible and -resistant isolates of <i>Staphylococcus</i>
В	Vancomycin	Staphylococcus spp. other than S. aureus	_		_	=	-	≤2 ≤4		8–16	≥16	(21) For <i>S. aureus</i> , vancomycinsusceptible isolates may become vancomycin intermediate during the course of prolonged therapy. (22) Send any <i>S. aureus</i> for which the vancomycin is ≥8 μg/mL to a referral laboratory. See Appendix A. Also refer to Table 3F for <i>S. aureus</i> , Subchapter 3.12 in M07,³ and Subchapter 3.9 in M02.¹ See comment (19). (23) Send any <i>Staphylococcus</i> spp. other than <i>S. aureus</i> for which the vancomycin MIC is ≥32 μg/mL to a referral laboratory. See Appendix A.
LIBOCI VO	OPEPTIDES				-		-		:	:	:	See also Subchapter 3.12 in M07³ and Subchapter 3.9 in M02.1
C	Dalbavancin	S. aureus,	T _ T	_	: _		1 _	≤0.25	: _	: _		
C	Oritavancin	including MRSA		_	<u> </u>		: -	≤0.12	: -	- -	: -	
C	Telavancin		_	_			†	≤0.12 ≤0.12	†	 	;	
Inv.	Teicoplanin	All staphylococci	_	_	 	_	 	<8	 	16	≥32	
LIPOPEPT		o.apj.oooo	I									
В	Daptomycin	All staphylococci	-	-	-	-	<u> </u>	≤1	<u> </u>	-	-	(24) Daptomycin should not be reported for isolates from the respiratory tract.
AMINOGLY (25) For sta		est susceptible, gentar	micin is use	ed only ir	n combin	ation with c	ther activ	e agents t	hat test s	usceptible		
C	Gentamicin	All staphylococci	10 μg	≥15	! _	13–14	≤12	<4	1	8	≥16	

or Or
Use
With
M02
and
M07

M100, 30th ed.

		Staphylococcus spp. Indications			e [·] Diam	e Categorie leter Breakp est whole m	ooints,	Inte	MIC Bro	Categorie eakpoints g/mL		
	Antimicrobial Agent		Disk Content	s	SD		R	s	SDD	,	R	Comments
MACROL		maications	Content			<u> </u>	1 1		. 000	<u> </u>	<u> </u>	Comments
		n organisms isolated f	rom the urina	ry tract.								
Α	Azithromycin	All staphylococci	15 μg	≥18	-	14–17	≤13	≤2	. – !	4	≥8	
Α	or clarithromycin or		15 μg	≥18	 	14–17	≤13	≤2		4	≥8	
Α	erythromycin		15 μg	≥23	! !	14–22	≤13	≤0.5		1–4	≥8	
0	Dirithromycin		15 μg	≥19	-	16–18	<u>≤15</u>	≤2		4	≥8	
		ible to doxycycline, mil			: -	15–18	≤14	<u>≤4</u>		8	≥16	ms that are intermediate or resistant to
					<u> </u>	15–18	<14	<4	:	8	>16	
В	Doxycycline		30 μg	≥16	<u> </u>	13–15	≤12	≤4	- 1	8	≥16	
В	Minocycline		30 μg	≥19	. –	15–18	≤14	≤4	: - :	8	≥16	See comment (26).
(28) Stap	ation of therapy. Te Ciprofloxacin	ay develop resistance sting of repeat isolates All staphylococci			rapy w		es. Therefo	ore, isolat	tes that a	re initially s	susceptible	may become resistant within 3 to 4 days
0	or				1	16–20	≤15	≤1		2	≥4	
С	Levofloxacin		F		:				-			
	Movifloyacin		5 μg	≥19	_	16–18	≤15	≤1	-	2	≥4	
0 0	Moxifloxacin Enoxacin		5 μg 5 μg 10 μg		- - -				- - -			
			5 μg 10 μg	≥19 ≥24 ≥18		16–18 21–23	≤15 ≤20 ≤14	≤1 ≤0.5 ≤2	- - -	2 1	≥4 ≥2 ≥8	(29) For testing and reporting of urinar tract isolates only.
0	Enoxacin		5 μg 10 μg 5 μg	≥19 ≥24 ≥18 ≥23	-	16–18 21–23 15–17	≤15 ≤20	≤1 ≤0.5		2 1 4	≥4 ≥2	
0	Enoxacin Gatifloxacin		5 μg 10 μg 5 μg 5 μg	≥19 ≥24 ≥18 ≥23 ≥18	-	16–18 21–23 15–17	≤15 ≤20 ≤14 ≤19	≤1 ≤0.5 ≤2 ≤0.5	_	2 1 4	≥4 ≥2 ≥8 ≥2	
0 0	Enoxacin Gatifloxacin Grepafloxacin		5 μg 10 μg 5 μg 5 μg 10 μg	≥19 ≥24 ≥18 ≥23	-	16–18 21–23 15–17 20–22 15–17	≤15 ≤20 ≤14 ≤19 ≤14	≤1 ≤0.5 ≤2 ≤0.5 ≤1	-	2 1 4 1 2	≥4 ≥2 ≥8 ≥2 ≥4	
0 0 0	Enoxacin Gatifloxacin Grepafloxacin Lomefloxacin		5 μg 10 μg 5 μg 5 μg	≥19 ≥24 ≥18 ≥23 ≥18 ≥22	-	16–18 21–23 15–17 20–22 15–17 19–21	≤15 ≤20 ≤14 ≤19 ≤14 ≤18	≤1 ≤0.5 ≤2 ≤0.5 ≤1 ≤2	-	2 1 4 1 2 4	≥4 ≥2 ≥8 ≥2 ≥4 ≥8	tract isolates only.
0 0 0 0	Enoxacin Gatifloxacin Grepafloxacin Lomefloxacin Norfloxacin		5 μg 10 μg 5 μg 5 μg 10 μg	≥19 ≥24 ≥18 ≥23 ≥18 ≥22 ≥17	-	16–18 21–23 15–17 20–22 15–17 19–21 13–16	≤15 ≤20 ≤14 ≤19 ≤14 ≤18 ≤12	≤1 ≤0.5 ≤2 ≤0.5 ≤1 ≤2 ≤4	-	2 1 4 1 2 4 8	≥4 ≥2 ≥8 ≥2 ≥4 ≥8 ≥16	tract isolates only.
0 0 0 0 0	Enoxacin Gatifloxacin Grepafloxacin Lomefloxacin Norfloxacin Ofloxacin		5 μg 10 μg 5 μg 5 μg 10 μg 10 μg 5 μg	≥19 ≥24 ≥18 ≥23 ≥18 ≥22 ≥17 ≥18	- - - - -	16–18 21–23 15–17 20–22 15–17 19–21 13–16 15–17	≤15 ≤20 ≤14 ≤19 ≤14 ≤18 ≤12 ≤14	≤1 ≤0.5 ≤2 ≤0.5 ≤1 ≤2 ≤4 ≤1	- - - -	2 1 4 1 2 4 8 2	≥4 ≥2 ≥8 ≥2 ≥4 ≥8 ≥16 ≥4	,
0 0 0 0 0 0	Enoxacin Gatifloxacin Grepafloxacin Lomefloxacin Norfloxacin Ofloxacin Sparfloxacin		5 μg 10 μg 5 μg 5 μg 10 μg 10 μg 10 μg 5 μg 5 μg 5 μg	≥19 ≥24 ≥18 ≥23 ≥18 ≥22 ≥17 ≥18 ≥19	- - - - -	16–18 21–23 15–17 20–22 15–17 19–21 13–16 15–17	≤15 ≤20 ≤14 ≤19 ≤14 ≤18 ≤12 ≤14		- - - -	2 1 4 1 2 4 8 2	≥4 ≥2 ≥8 ≥2 ≥4 ≥8 ≥16 ≥4 ≥2	tract isolates only.

Table 2C. Staphylococcus spp. (Continued)

Test/ Report	Antimicrobial	Disk		e Diamet	Categorie ter Breakp t whole mi	oints,	Inte	rpretive C MIC Bre µg	•			
Group	Agent	spp. Indications	Content	s	SDD	1	R	S	SDD	1	R	Comments
LINCOS	AMIDES											
A	Clindamycin	All staphylococci	2 μg	≥21	-	15–20	≤14	≤0.5	-	1–2	≥4	(30) 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 before reporting clindamycin (see Table 3H, Subchapter 3.9 in M02,¹ and Subchapter 3.12 in M07³).
							:		:		:	See comment (26).
FOLATE	PATHWAY ANTAG	ONISTS										
Α	Trimethoprim- sulfamethoxazole	All staphylococci	1.25/23.75 μg	≥16	-	11–15	≤10	≤2/38	-	-	≥4/76	
U	Sulfonamides	All staphylococci	250 or 300 μg	≥17	-	13–16	≤12	≤256	-	-	≥512	(31) Sulfisoxazole can be used to represent any of the currently available sulfonamide preparations.
U	Trimethoprim	All staphylococci	5 μg	≥16	-	11–15	≤10	≤8	-	_	≥16	
PHENIC	OLS											
С	Chloramphenicol	All staphylococci	30 μg	≥18	-	13–17	≤12	≤8	-	16	≥32	See comment (26).
ANSAM	YCINS	•										
В	Rifampin	All staphylococci	5 μg	≥20	-	17–19	≤16	≤1	-	2	≥4	(32) Rx: Rifampin should not be used alone for antimicrobial therapy.
STREPT	OGRAMINS											
0	Quinupristin- dalfopristin	S. aureus	15 μg	≥19	-	16–18	≤15	≤1	-	2	≥4	(33) For reporting against methicillin (oxacillin)-susceptible S. aureus.

Table 2C. Staphylococcus spp. (Continued)

Test/ Report	Antimicrobial	Staphylococcus spp.	Disk		Diame	Categorie ter Breakp whole mi	oints,	Inte	MIC Bre	Categorie: eakpoints, g/mL		Comments
Group	Agent	Indications	Content	s	SDD	<u>.</u> .	R	S	SDD	ı	R	
OXAZOL	IDINONES											
В	Linezolid	All staphylococci	30 μg	≥21	_	_	≤20	≤ 4	_	-	≥8	(34) When testing linezolid, disk diffusion zones should be examined using transmitted light. Organisms with resistant results by disk diffusion should be confirmed using an MIC method.
В	Tedizolid	S. aureus, including MRSA	ı	_	_	_	_	≤0.5	_	1	≥2	

Abbreviations: ATCC®, American Type Culture Collection; BMHA, blood Mueller-Hinton agar; CAMHB, cation-adjusted Mueller-Hinton broth; I, intermediate; ICR, inducible clindamycin resistance; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; MRS, methicillin (oxacillin)-resistant staphylococci; MRSA, methicillin (oxacillin)-resistant S. aureus; PBP2a, penicillin-binding protein 2a; PCR, polymerase chain reaction; QC, quality control; R, resistant; S, susceptible; SDD, susceptible-dose dependent; UTI, urinary tract infection.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2C

- CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 3 CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- García-Álvarez L, Holden MT, Lindsay H, et al. Methicillin-resistant Staphylococcus aureus with a novel mecA homologue in human and bovine populations in the UK and Denmark: a descriptive study. Lancet Infect Dis. 2011;11(8):595-603.

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Table 2D. Zone Diameter and MIC Breakpoints for Enterococcus spp.

Testing Conditions

Medium: Disk diffusion: MHA

Broth dilution: CAMHB; CAMHB supplemented to

50 μg/mL calcium for daptomycin

Agar dilution: MHA; agar dilution has not been validated

for daptomycin

Inoculum: Broth culture method or colony suspension, equivalent to

a 0.5 McFarland standard

Incubation: 35°C±2°C; ambient air

Disk diffusion: 16–18 hours Dilution methods: 16–20 hours

All methods: 24 hours for vancomycin

Routine QC Recommendations (see Tables 4A-1 and 5A-1 for acceptable QC ranges)

Disk diffusion:

S. aureus ATCC®a 25923

Dilution methods:

E. faecalis ATCC® 29212

Refer to Tables 4A-2 and 5A-2 to select strains for routine QC of β -lactam combination agents.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

Refer to Tables 3F and 3I for additional testing recommendations, reporting suggestions, and QC.

General Comments

- (1) For disk diffusion, test a maximum of 12 disks on a 150-mm plate and no more than 6 disks on a 100-mm plate; disks should be placed no less than 24 mm apart, center to center (see M02, Subchapter 3.6). Each zone diameter should be clearly measurable; overlapping zones prevent accurate measurement. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide²). Hold the Petri plate a few inches above a black background illuminated with reflected light, except for vancomycin, which should be read with transmitted light (plate held up to light source). The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. Any discernible growth within the zone of inhibition indicates vancomycin resistance.
- (2) For enterococci when testing chloramphenicol, erythromycin, linezolid, tedizolid, and tetracycline by broth microdilution MIC, trailing growth can make endpoint determination difficult. In such cases, read the MIC at the lowest concentration where the trailing begins. Tiny buttons of growth should be ignored (see M07.³ Figures 3 and 4).
- (3) WARNING: For *Enterococcus* spp., aminoglycosides (except for high-level resistance testing), cephalosporins, clindamycin, and trimethoprim-sulfamethoxazole may appear active *in vitro*, but they are not effective clinically, and isolates should not be reported as susceptible.
- (4) Synergy between ampicillin, penicillin, or vancomycin and an aminoglycoside can be predicted for enterococci by using a high-level aminoglycoside (gentamicin and streptomycin) test (see Table 3J).
- (5) Intermediate ranges denoted with a "A" for the applicable antimicrobial agents in the drug groups in Tables 2 are based on the known ability of these agents to concentrate in the urine; some agents may also have the potential to concentrate at other anatomical sites (eg, epithelial lining).

Table 2D. Enterococcus spp. (Continued)

Test/Report	Antimicrobial	microbial Disk		etive Ca Zone Di Breakp earest w	amete oints	,	Int		Categori eakpoint g/mL		
Group	Agent	Content	S	1		R	S	SDD	ı	R	Comments
PENICILLINS											
A A	Penicillin Ampicillin	10 units 10 μg	≥15 ≥17			≤14 ≤16	≤8 ≤8			≥16 ≥16	(6) 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> . (7) Enterococci susceptible to penicillin are predictably susceptible to ampicillin, amoxicillin, ampicillin-sulbactam, amoxicillin-clavulanate, and piperacillintazobactam for non-β-lactamase-producing enterococci. However, enterococci susceptible to ampicillin cannot be assumed to be susceptible to penicillin. If penicillin results are needed, testing of penicillin is required. (8) <i>Rx:</i> Combination therapy with ampicillin, penicillin, or vancomycin (for susceptible strains only), plus an aminoglycoside, is usually indicated for serious enterococcal infections, such as endocarditis, unless high-level resistance to both gentamicin and streptomycin is documented; such combinations are predicted to result in synergistic killing of the <i>Enterococcus</i> . For strains with low-level penicillin or ampicillin resistance when combination therapy with a β-lactam is being considered, also see additional testing and reporting information in Table 3J. ⁴ (9) Penicillin or ampicillin resistance among enterococci due to β-lactamase production has been reported very rarely. Penicillin or ampicillin resistance due to β-lactamase production is not reliably detected with routine disk or dilution methods but is detected using a direct, nitrocefin-based β-lactamase test. Because of the rarity of β-lactamase—positive enterococci, this test does not need to be performed routinely but can be used in selected cases. A positive
				:	:			:	• • •	-	β-lactamase test predicts resistance to penicillin as
				:						:	well as amino- and ureidopenicillins (see Glossary I).

Test/Report	Antimicrobial Agent	Disk	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm			Int	MIC Bre	Categories a akpoints, //mL	and	
Group			s	1	R	S	SDD	1	R	Comments
GLYCOPEPT	IDES									·
В	Vancomycin	30 µg	≥17	15–16	≤14	≤4	-	8–16	≥32	(10) 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 M07.³ For isolates for which the vancomycin MICs are 8–16 μg/mL, perform biochemical tests for identification as listed under the "Vancomycin MIC≥ ≥ 8 μg/mL" test found in Table 3G.
LIPOGLYCOF	PERTINES						<u>:</u>	<u>:</u>	<u>:</u>	See general comment (4) and comment (8).
C	Dalbavancin	1	1			≤0.25		,		(44) For reporting against vancomy sin supportible
	Daibavancin	_	_	_	_	≥0.25	_	_	_	(11) For reporting against vancomycin-susceptible <i>E. faecalis</i> .
С	Oritavancin	_	_		_	≤0.12		<u>:</u> –	_	See comment (11).
С	Telavancin	_	_	-	_	≤0.25	<u> </u>	: -	: -	See comment (11).
Inv.	Teicoplanin	30 μg	≥14	11–13	≤10	≤8	-	16	≥32	
LIPOPEPTIDE	S									
В	Daptomycin <i>E. faecium</i> only	-	_	-	-	-	≤4	_	≥8	 (12) Daptomycin should not be reported for isolates from the respiratory tract. (13) The breakpoint for SDD is based on a dosage regimen of 8–12 mg/kg administered every 24 h and is intended for serious infections due to <i>E. faecium</i>. Consultation with an infectious diseases specialist is recommended.
В	Daptomycin Enterococcus spp. other than E. faecium	_	-	-	-	≤2	-	4	≥8	(14) The breakpoint for susceptible is based on a dosage regimen of 6 mg/kg administered every 24 h. See comment (12).
MACROLIDES	S									
0	Erythromycin	15 μg	≥23	14–22	≤13	≤0.5	<u> </u>	1–4	≥8	(15) Not routinely reported on isolates from the urinary tract.

Table 2D. Enterococcus spp. (Continued)

Test/Report	Antimicrobial	Disk	Zone Dia	tive Catego ameter Brea arest whole	akpoints,	Interp		tegories kpoints, p/mL	and MIC		
Group	Agent	Content	s	1	R	S	SDD	- 1	R	Comments	
TETRACYCLIN	NES										
	that are susceptible y be susceptible to d				susceptible	to doxyc	ycline and	minocycl	ine. Howeve	er, some organisms that are intermediate or resistant to	
Ū	Tetracycline	30 μg	≥19	15–18	≤14	≤4	_	8	≥16		
0	Doxycycline	30 μg	≥16	13–15	≤12	≤4	-	8	≥16		
0	Minocycline	30 μg	≥19	15–18	≤14	≤4	-	8	≥16		
FLUOROQUIN	OLONES										
U	Ciprofloxacin	5 μg	≥21	16–20 ^	≤15	≤1	-	2^	≥4		
U	Levofloxacin	5 μg	≥17	14–16 ^	≤13	≤2		4^	≥8		
0	Gatifloxacin	5 μg	≥18	15–17 ^	≤14	≤2	-	4^	≥8		
0	Norfloxacin	10 μg	≥17	13–16	≤12	≤4	-	8	≥16	(17) For testing and reporting of urinary tract isolates only.	
NITROFURAN	TOINS										
U	Nitrofurantoin	300 μg	≥17	15–16	≤14	≤32	-	64	≥128		
ANSAMYCINS	3										
0	Rifampin	5 μg	≥20	17–19	≤16	≤1	-	2	≥4	(18) Rx: Rifampin should not be used alone for antimicrobial therapy.	
FOSFOYCINS									•		
U	Fosfomycin	200 μg	≥16	13–15	≤12	≤64	-	128	≥256	(19) For testing and reporting of <i>E. faecalis</i> urinary tract isolates only.	
										(20) The approved MIC testing method is agar dilution. Agar media should be supplemented with 2 μg/mL of glucose-6-phosphate. Broth dilution testing should not be performed.	
				:	· · ·					(21) The 200-μg fosfomycin disk contains 50 μg glucose-6-phosphate.	
PHENICOLS											
0	Chloramphenicol	30 μg	≥18	13–17	≤12	≤8	_	16	≥32	See comment (15).	
STREPTOGRA											
0	Quinupristin- dalfopristin	15 μg	≥19	16–18	≤15	≤1		2	≥4	(22) For reporting against vancomycin-resistant Enterococcus faecium.	
OXAZOLIDINO	NES										
В	Linezolid	30 μg	≥23	21–22^	≤20	≤2	<u> </u>	4^	≥8		
В	Tedizolid		_		_	≤0.5	. – :	_	_	(23) For reporting against E. faecalis only.	

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible; SDD, susceptible-dose dependent.

Table 2D. Enterococcus spp. (Continued)

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2D

- CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 3 CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- Murray BE, Arias CA, Nannini EC. Glycopeptides (vancomycin and teicoplanin), streptogramins (quinupristin-dalfopristin), lipopeptides (daptomycin), and lipoglycopeptides (telavancin). In: Bennett JE, Dolin R, Blaser MJ. *Mandell, Douglas, and Bennett's Principles and Practice of Infectious Diseases.* 8th ed. Philadelphia, PA: Elsevier Saunders; 2015;377-400.

M100, 30th ed

Table 2E. Zone Diameter and MIC Breakpoints for Haemophilus influenzae and Haemophilus parainfluenzae

Testing Conditions

Medium: Disk diffusion: HTM
Broth dilution: HTM broth

Inoculum: Colony suspension, equivalent to a 0.5 McFarland standard

prepared using colonies from an overnight (preferably 20- to

24-hour) chocolate agar plate (see comment [2])

Incubation: 35°C±2°C

Disk diffusion: 5% CO₂; 16–18 hours Broth dilution: ambient air; 20–24 hours **Routine QC Recommendations** (see Tables 4A-1, 4B, 5A-1, and 5B for acceptable QC ranges)

H. influenzae ATCC®a 49247 H. influenzae ATCC® 49766

Use either *H. influenzae* ATCC® 49247 or *H. influenzae* ATCC® 49766 or both of these strains, based on the antimicrobial agents to be tested. Neither strain has QC ranges for all agents that might be tested against *H. influenzae* or *H. parainfluenzae*.

E. coli ATCC® 35218 (when testing amoxicillin-clavulanate)

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) Haemophilus spp., as used in this table, includes only H. influenzae and H. parainfluenzae. See CLSI document M45¹ for testing and reporting recommendations for other species of Haemophilus.
- (2) The 0.5 McFarland suspension contains approximately 1 to 4 × 10⁸ CFU/mL. Use care in preparing this suspension, because higher inoculum concentrations may lead to false-resistant results with some β-lactam antimicrobial agents, particularly when β-lactamase-producing strains of *H. influenzae* are tested.
- (3) For disk diffusion, test a maximum of 9 disks on a 150-mm plate and 4 disks on a 100-mm plate. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk. Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.
- (4) For isolates of *H. influenzae* from CSF, only results of testing with ampicillin, any of the 3rd-generation cephalosporins listed below, chloramphenicol, and meropenem are appropriate to report.
- (5) Amoxicillin-clavulanate, azithromycin, cefaclor, cefdinir, cefixime, cefpodoxime, cefprozil, cefuroxime, and clarithromycin are used as empiric therapy for respiratory tract infections due to *Haemophilus* spp. The results of susceptibility tests with these antimicrobial agents are often not necessary for management of individual patients.

Table 2E. Haemophilus influenzae and Haemophilus parainfluenzae (Continued)

(6) To make HTM: Prepare a fresh hematin stock solution by dissolving 50 mg of hematin powder in 100 mL of 0.01 mol/L NaOH with heat and stirring until the powder is thoroughly dissolved. Add 30 mL of the hematin stock solution and 5 g of yeast extract to 1 L of MHA, and autoclave. After autoclaving and cooling, add 3 mL of an NAD stock solution (50 mg NAD dissolved in 10 mL distilled water, filter sterilized) aseptically.

Test/Report	Antimicrobial	Disk	Zone Di	etive Catego ameter Brea arest whole	kpoints,	Interpreti MIC	ve Catego Breakpoi µg/mL		
Group	Agent	Content	S	ı	R	s	1	R	Comments
PENICILLINS			•						
A	Ampicillin	10 μg	≥22	19–21	≤18	≤1	2	≥4	See general comment (4). (7) The results of ampicillin susceptibility tests should be used to predict the activity of amoxicillin. The majority of isolates of <i>H. influenzae</i> that are resistant to ampicillin and amoxicillin produce a TEM-type β-lactamase. In most cases, a direct β-lactamase test can provide a rapid means of detecting resistance to ampicillin and amoxicillin. (8) Rare BLNAR strains of <i>H. influenzae</i> should be considered resistant to amoxicillinclavulanate, ampicillin-sulbactam, cefaclor, cefamandole, cefetamet, cefonicid, cefprozil, cefuroxime, loracarbef, and piperacillintazobactam, despite apparent <i>in vitro</i>
									susceptibility of some BLNAR strains to these agents.
B-L ACTAM COL	MBINATION AGENTS			<u> </u>	<u> </u>		<u>: </u>		agents.
В	Ampicillin-sulbactam	10/10 μg	≥20	-	≤19	≤2/1	1 - 1	≥4/2	See comment (8).
С	Amoxicillin-clavulanate	20/10 μg	≥20	_	≤19	≤4/2	: - :	≥8/4	See general comment (5) and comment (8).
0	Piperacillin-tazobactam	100/10 μg	≥21	_	_	≤ 1/4		≥2/4	See comment (8).
CEPHEMS (PAI	RENTERAL) (Including cep	halosporins I, II	, III, and IV	. Please refe	r to Gloss	ary I.)			, ,
B B	Cefotaxime or ceftazidime or	30 μg 30 μg	≥26 ≥26	- -	_ _	≤2 ≤2	_	_	See general comment (4).
В	ceftriaxone	30 μg	≥26	_	_	≤2	-	_	
С	Cefuroxime	30 μg	≥20	17–19	≤16	≤4	8	≥16	See general comment (5) and comment (8).
С	Ceftaroline	30 μg	≥30	-	_	≤0.5	-	_	(9) For <i>H. influenzae</i> only. (10) Breakpoints are based on a dosage regimen of 600 mg administered every 12 h.
0	Cefonicid	30 μg	≥20	17–19	≤16	≤4	8	≥16	See comment (8).

Table 2E. Haemophilus influenzae and Haemophilus parainfluenzae (Continued)

Test/Report	Zone Diameter Breakpoints, MIC nearest whole mm					Interpretiv MIC I	/e Catego Breakpoi μg/mL		
Group	Agent	Content	S	1	R	s	1	R	Comments
CEPHEMS (PA	RENTERAL) (Including ce	phalosporins I, I	, III, and IV	. Please refe	er to Gloss	ary I.) (Contir	nued)		
0	Cefamandole	-	_	-	_	≤4	8	≥16	See comment (8).
0	Cefepime	30 μg	≥26	-	_	≤2	- :	_	
0	Ceftizoxime	30 μg	≥26	-	-	≤2	- :	_	See general comment (4).
CEPHEMS (OR	AL)								
С	Cefaclor	30 μg	≥20	17–19	≤16	≤8	16	≥32	See general comment (5) and comment (8).
С	Cefprozil	30 μg	≥18	15–17	≤14	≤8	16	≥32	
С	Cefdinir or	5 μg	≥20	! –	_	≤1	-	_	See general comment (5).
С	cefixime or	5 μg	≥21	-	-	≤1	. – !	_	
С	cefpodoxime	10 μg	≥21	-	-	≤2	- :	_	
С	Cefuroxime	30 μg	≥20	17–19	≤16	≤4	8	≥16	See general comment (5) and comment (8).
0	Loracarbef	30 μg	≥19	16–18	≤15	≤8	16	≥32	See general comment (5) and comment (8).
0	Ceftibuten	30 μg	≥28	 	i –	≤2	_	_	, , , , , ,
Inv.	Cefetamet	10 μg	≥18	15–17	≤14	≤4	8	≥16	See comment (8).
MONOBACTAN	MS	1 10				•			. ,
С	Aztreonam	30 μg	≥26	-	-	≤2	- 1	_	
CARBAPENEM	IS								
В	Meropenem	10 μg	≥20	-	_	≤0.5	- 1	_	See general comment (4).
С	Ertapenem or	10 μg	≥19	: -	-	≤0.5	: - :	_	
С	imipenem	10 μg	≥16	-	<u> </u>	≤4	. – !	_	
0	Doripenem	10 μg	≥16	<u> </u>	_	≤1	- 1	_	
MACROLIDES					,				
С	Azithromycin	15 μg	≥12	-	_	≤4	- :	_	See general comment (5).
С	Clarithromycin	15 μg	≥13	11–12	≤10	≤8	16	≥32	
nferred from tet	that are susceptible to tetra tracycline resistance.								stance to doxycycline and minocycline cannot
С	Tetracycline	30 μg	≥29	26–28	≤25	≤2	4	≥8	
FLUOROQUING		1				1	,		
В	Ciprofloxacin or	5 μg	≥21	-	-	≤1	-	-	
В	levofloxacin or	5 μg	≥17	_	_	≤2	-	-	
В	moxifloxacin	5 μg	≥18	-	_	≤1	-		
0	Gemifloxacin	5 μg	≥18	_	-	≤0.12	- :	-	
0	Gatifloxacin	5 μg	≥18	<u> </u>	_	≤1	<u> </u>	_	
0	Grepafloxacin	5 μg	≥24	-	_	≤0.5	- :	_	
0	Lomefloxacin	10 μg	≥22	_	_	≤2	- :	_	
0	Ofloxacin	5 μg	≥16	<u> </u>	_	≤2	- !	_	
0	Sparfloxacin		_	: -	: -	≤0.25	: - :		

Table 2E. Haemophilus influenzae and Haemophilus parainfluenzae (Continued)

Test/Report	Antimicrobial	Disk	Zone Dia	tive Catego ameter Brea arest whole	kpoints,		re Categories Breakpoints, µg/mL	and MIC	
Group	Agent	Content S I R S I				R	Comments		
FLUOROQUINO	DLONES (Continued)								
0	Trovafloxacin	10 μg	≥22	-	-	≤1	_	-	
lnv.	Fleroxacin	5 μg	≥19	-	_	≤2	-	-	
FOLATE PATH	WAY ANTAGONISTS								
С	Trimethoprim- sulfamethoxazole	1.25/23.75 μg	≥16	11–15	≤10	≤0.5/9.5	1/19–2/38	≥4/76	
PHENICOLS									
С	Chloramphenicol	30 μg	≥29	26–28	≤25	≤2	4	≥8	See general comment (4).
									(12) Not routinely reported on isolates from the urinary tract.
ANSAMYCINS									
С	Rifampin	5 μg	≥20	17–19	≤16	≤1	2	≥4	(13) May be appropriate only for prophylaxis of case contacts. These breakpoints do not apply to therapy of patients with invasive <i>H. influenzae</i> disease.

Abbreviations: ATCC[®], American Type Culture Collection; BLNAR, β -lactamase negative, ampicillin-resistant; CFU, colony-forming unit(s); CSF, cerebrospinal fluid; HTM, *Haemophilus* test medium; I, intermediate; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; **NAD**, β -nicotinamide adenine dinucleotide; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

Reference for Table 2E

¹ CLSI. *Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently Isolated or Fastidious Bacteria.* 3rd ed. CLSI guideline M45. Wayne, PA: Clinical and Laboratory Standards Institute; 2016.

M100, 30th ed

Table 2F. Zone Diameter and MIC Breakpoints for Neisseria gonorrhoeae

Testing Conditions

Medium: Disk diffusion: GC agar base and 1% defined growth supplement. (The use of a

cysteine-free growth supplement is not required for disk diffusion testing.)

Agar dilution: GC agar base and 1% defined growth supplement. (The use of a cysteine-free growth supplement is required for agar dilution tests with carbapenems and clavulanate. Cysteine-containing defined growth supplement

does not significantly alter dilution test results with other drugs.)

Inoculum: Colony suspension, equivalent to a 0.5 McFarland standard prepared in MHB or

0.9% phosphate-buffered saline, pH 7, using colonies from an overnight (20- to

24-hour) chocolate agar plate incubated in 5% CO₂

Incubation: 36°C±1°C (do not exceed 37°C); 5% CO₂; all methods, 20–24 hours

Routine QC Recommendations (see Tables 4B and 5C for acceptable QC ranges)

N. gonorrhoeae ATCC®a 49226

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) For disk diffusion, test a maximum of 9 disks on a 150-mm plate and 4 disks on a 100-mm plate. For some agents, eg, fluoroquinolones or cephalosporins, only 2 to 3 disks may be tested per plate. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk. Hold the Petri plate a few inches above a black background illuminated with reflected light. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth.
- (2) The clinical effectiveness of cefotetan, cefoxitin, and spectinomycin for treating infections due to organisms that produce intermediate results with these agents is unknown.
- (3) For disk diffusion testing of *N. gonorrhoeae*, an intermediate result for an antimicrobial agent indicates either a technical problem that should be resolved by repeat testing or a lack of clinical experience in treating infections due to organisms with these zones. Strains with intermediate zones to agents other than cefotetan, cefoxitin, and spectinomycin have a documented lower clinical cure rate (85% to 95%) compared with > 95% for susceptible strains.
- (4) The recommended medium for testing *N. gonorrhoeae* consists of GC agar to which a 1% defined growth supplement (1.1 g L-cystine, 0.03 g guanine HCl, 0.003 g thiamine HCl, 0.013 g para-aminobenzoic acid, 0.01 g B12, 0.1 g cocarboxylase, 0.25 g NAD, 1 g adenine, 10 g L-glutamine, 100 g glucose, 0.02 g ferric nitrate, 25.9 g L-cysteine HCl [in 1 L H₂Q]) is added after autoclaving.

Table 2F. Neisseria gonorrhoeae (Continued)

Test/Report	Antimicrobial	Disk	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm Interpretive Categories and MIC Breakpoints, μg/mL						
Group	Agent	Content	s	ı	R	s	1	R	Comments
PENICILLINS			•	-					
0	Penicillin	10 units	≥47	27–46	. ≤26	≤0.06	0.12–1	≥2	See general comment (3).
				- - - - - - -					(5) A positive β-lactamase test predicts resistance to penicillin, ampicillin, and amoxicillin.
									(6) A β -lactamase test detects one form of penicillin resistance in <i>N. gonorrhoeae</i> and also may be used to provide epidemiological information. Strains with chromosomally mediated resistance can be detected only by the disk diffusion method or the agar dilution MIC method.
									(7) Gonococci that produce zones of inhibition of ≤19 mm around a 10-unit penicillin disk are likely to be β-lactamase–producing strains. However, the β-lactamase test remains preferable to other
				-					susceptibility methods for rapid, accurate recognition of this plasmid-mediated penicillin resistance.
CEPHEMS (PA	RENTERAL) (Including cepha	alosporins I, I	II, III, and I\	/. Please ref	fer to Glo	ssary I.)			
Α	Ceftriaxone	30 μg	≥35	_	-	≤0.25	-	-	
0	Cefoxitin	30 μg	≥28	24–27	≤23	≤2	4	≥8	See general comment (2).
0	Cefepime	30 μg	≥31		<u> </u>	≤0.5	<u> </u>	<u> </u>	
0	Cefotaxime	30 μg	≥31	_	<u> </u>	≤0.5	-	<u> </u>	
0	Cefotetan	30 μg	≥26	20–25	≤19	≤2	4	≥8	See general comment (2).
0	Ceftizoxime	30 μg	≥38	<u> </u>	<u> </u>	≤0.5	<u>: - </u>	<u>i – </u>	
CEPHEMS (OR		_		ı			1	1	
A	Cefixime	5 μg	≥ 31	_	 	≤ 0.25	 	-	
0	Cefpodoxime	10 μg	≥ 29		: -	≤0.5	<u> </u>	<u>i – </u>	

Test/Report	Antimicrobial	Disk	z	nterpretive Categories and Zone Diameter Interpretive Categories and Breakpoints, MIC Breakpoints, nearest whole mm µg/mL						
Group	Agent	Content	s	ı	R	S	1		R	Comments
MACROLIDES		•					•			
A	Azithromycin	_	-	-	_	≤1	-		_	(8) This breakpoint presumes that azithromycin (1 g single dose) is used in an approved regimen that includes an additional antimicrobial agent (ie, ceftriaxone 250 mg IM single dose).
TETRACYCLINE						. , .				
· / · ·	at are susceptible to tetracyclin	1	•		 		, -		. 0	(40) O and a solid solid to the control of the solid to t
A	Tetracycline	30 μg	≥38	31–37	: ≤30 : :	≤0.25	0.5–1		≥2	(10) Gonococci with 30-μg tetracycline disk zone diameters of ≤ 19 mm usually indicate a plasmid-mediated tetracycline-resistant <i>N. gonorrhoeae</i> isolate. Resistance in these strains should be confirmed by a dilution test (MIC ≥ 16 μg/mL).
FLUOROQUINC	DLONES									
See general com	nment (3).									
Α	Ciprofloxacin	5 μg	≥41	28–40	≤27	≤ 0.06	0.12-0	5	≥1	
AMINOCYCLITO	DLS									
0	Spectinomycin	100 μg	≥18	15–17	≤14	≤ 32	64		≥128	See general comment (2).

O Spectinomycin $100 \,\mu\text{g}$ ≥ 18 15-17 ≤ 14 ≤ 32 64 ≥ 128 See general comment (2). Abbreviations: ATCC®, American Type Culture Collection; I, intermediate; IM, intramuscular; MHB, Mueller-Hinton broth; MIC, minimal inhibitory concentration; NAD, β -nicotinamide adenine dinucleotide; pH, negative logarithm of hydrogen ion concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

M100, 30th ed

Table 2G. Zone Diameter and MIC Breakpoints for Streptococcus pneumoniae

Testing Conditions

Medium: Disk diffusion: MHA with 5% sheep blood or MH-F agar (MHA with 5% defibrinated

horse blood and 20 μg/mL NAD)

Broth dilution: CAMHB with LHB (2.5% to 5% v/v) (see M071 for instructions for

preparation of LHB)

Agar dilution: MHA with sheep blood (5% v/v); recent studies using the agar dilution

method have not been performed and reviewed by the subcommittee.

Inoculum: Colony suspension, equivalent to a 0.5 McFarland standard, prepared using colonies

from an overnight (18- to 20-hour) sheep blood agar plate

Incubation: 35°C±2°C

Disk diffusion: 5% CO₂; 20-24 hours

Dilution methods: ambient air; 20-24 hours (CO₂ if necessary, for growth with agar

dilution)

Routine QC Recommendations (see Tables 4B and 5B for acceptable QC ranges)

S. pneumoniae ATCC®a 49619

Disk diffusion: deterioration of oxacillin disk content is best assessed with *S. aureus* ATCC® 25923, with an acceptable range of 18–24 mm on unsupplemented MHA.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) For disk diffusion, test a maximum of 9 disks on a 150-mm plate and 4 disks on a 100-mm plate. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the *M02 Disk Diffusion Reading Guide*²). The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Do not measure the zone of inhibition of hemolysis. Measure the zones from the upper surface of the agar illuminated with reflected light, with the cover removed. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.
- (2) For pneumococci when testing chloramphenicol, clindamycin, erythromycin, linezolid, tedizolid, and tetracycline by broth microdilution MIC, trailing growth can make end-point determination difficult. In such cases, read the MIC at the lowest concentration where the trailing begins. Tiny buttons of growth should be ignored (see M07,¹ Figures 3 and 4). With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, read the end point at the concentration in which there is ≥80% reduction in growth compared with the control (see M07,¹ Figure 5).
- (3) Amoxicillin, ampicillin, cefepime, cefotaxime, ceftriaxone, cefuroxime, ertapenem, imipenem, and meropenem may be used to treat pneumococcal infections; however, reliable disk diffusion susceptibility tests with these agents do not yet exist. Their *in vitro* activity is best determined using an MIC method.
- (4) For *S. pneumoniae* isolated from CSF, penicillin and cefotaxime, ceftriaxone, or meropenem should be tested by a reliable MIC method (such as that described in M07¹) and reported routinely. Such isolates can also be tested against vancomycin using the MIC or disk diffusion method.
- (5) For disk diffusion, results using MHA with 5% sheep blood and MH-F agar were equivalent when disk contents, testing conditions, and zone diameter breakpoints in Table 2G were used. Disk diffusion QC ranges for *S. pneumoniae* ATCC® 49619 in Table 4B apply to testing using either MHA with 5% sheep blood or MH-F agar.

Test/Report Antimicrobial		Disk	Zone D	etive Catego iameter Bre arest whole	akpoints,		etive Catego IIC Breakpoi µg/mL		
Group	Agent	Content	s	1	R	s	1	R	Comments
mpicillin-sulba	ningitis isolates, a penicillin ctam, amoxicillin, amoxicill ripenem, ertapenem, imipe	in-clavulanate, cef	aclor, cefdi	nir, cefditore	0 mm) can _l n, cefepime	predict sus , cefotaxir	sceptibility to me, cefpodox	the followin	ng β-lactams: ampicillin (oral or parenteral), zil, ceftaroline, ceftizoxime, ceftriaxone,
see general co									
A	Penicillin	1 μg oxacillin	≥20		——————————————————————————————————————	_	_	_	(7) Isolates of pneumococci with oxacillin zone sizes ≥20 mm are susceptible (MIC≤0.06 μg/mL) to penicillin. Penicillin and cefotaxime, ceftriaxone, or meropenem MICs should be determined for isolates with oxacillin zone diameters ≤19 mm, because zones ≤19 mm occur with penicillin-resistant, -intermediate, o certain -susceptible strains. For isolates with oxacillin zones≤19 mm, do not report penicilli as resistant without performing a penicillin MIC test.
A	Penicillin parenteral (nonmeningitis)	-	-	-		≤2	4	≥8	(8) Rx: Doses of intravenous penicillin of at le 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 ≤ 2 μg/mL. Strains with an intermediate MIC of 4 μg/mL may necessitate penicillin doses of 18–24 million units per day (9) For all isolates other than those from CSF, report interpretations for both meningitis and nonmeningitis.
A	Penicillin parenteral (meningitis)	-	_	-	-	≤0.06	_	≥0.12	(10) Rx: Use of penicillin in meningitis require therapy with maximum doses of intravenous penicillin (eg, at least 3 million units every 4 hours in adults with normal renal function). (11) For CSF isolates, report only meningitis interpretations.
				:	1		:	:	See general comment (4).

Table 2G. Streptococcus pneumoniae (Continued)

Test/Report	J	Disk	Zo	retive Categ and one Diamete Breakpoints rest whole i	er ,	Interpre MIC			
Group		Content	S	1	R	S	ı	R	Comments
PENICILLINS				•				•	
С	Amoxicillin (nonmeningitis)	_	_	_	-	≤2	4	≥8	
С	Amoxicillin-clavulanate (nonmeningitis)					≤2/1	4/2	≥8/4	
·	ARENTERAL) (Including cepha	losporins I, II	l, III, and I	V. Please re	fer to GI	ossary I.)			
See comment				·	,				
0	Cefepime (meningitis)	_	_	-	_	≤0.5	1	≥2	(13) In the United States, for CSF isolates, report only nonmeningitis interpretations. There is not an FDA-approved indication for the use of cefepime for meningitis in the United States.
В	Cefepime (nonmeningitis)	_	-	-	-	≤1	2	≥4	(14) In the United States, report only interpretations for nonmeningitis and include the nonmeningitis notation on the report.
В	Cefotaxime (meningitis)	_	_	: -	: -	≤0.5	1	: ≥2	(15) For CSF isolates, report only meningitis
В	Ceftriaxone (meningitis)	_	-	<u>.</u> –	-	≤0.5	1	≥2	interpretations.
									(16) Rx: Use of cefotaxime or ceftriaxone in meningitis requires therapy with maximum doses.
				:	:			:	See general comment (4).
В	Cefotaxime (nonmeningitis)	_	_	-	-	≤1	2	≥4	(17) For all isolates other than those from
В	Ceftriaxone (nonmeningitis)	_	_	-	_	≤ 1	2	≥4	CSF, report interpretations for both meningitis and nonmeningitis.
С	Ceftaroline (nonmeningitis)	30 μg	≥26	-	_	≤0.5	_	_	(18) Breakpoints are based on a dosage regimen of 600 mg administered every 12 h.
С	Cefuroxime (parenteral)	_	-	-	<u> </u>	≤0.5	1	≥2	
CEPHEMS (O									
С	Cefuroxime (oral)	_	-	-	-	≤1	2	≥4	(19) Interpretations for oral cefuroxime ma be reported for isolates other than those from CSF.
0	Cefaclor	_	-	-	-	≤1	2	≥4	
0	Cefdinir	_	-		<u> </u>	≤0.5	1	≥2	
0	Cefpodoxime	_	_	-	-	≤0.5	1	≥2	
	Cefprozil	_	_	_		≤2	4	≥8	
Ο	Ceipiozii					_ ∠		0	

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Table 2G. St	reptococcus	pneumoni	ae ((Continu	ued)
					Inte

Test/Report Antimicrobial		Disk	Zo	tive Categor one Diamete Breakpoints, irest whole r	r		ive Categorie Breakpoints μg/mL		
Group	Agent	Content	S	i I	R	S	I	R	Comments
CARBAPENE	ŴS								
2	(0)								
See comment B	(b). Meropenem	_	I _	! –	! _	≤0.25	0.5	. ≥1	See general comment (4) and comment (7).
С	Ertapenem		_	: -	: -	≥0.25 ≤1	2	≥4	See general comment (4) and comment (7).
C	Imipenem	_	_	_	_	≤ 1 ≤0.12	0.25–0.5	≥4 ≥1	
0	Doripenem			! 		≥0.12 ≤1	0.25-0.5		
SLYCOPEPTI		_	_		<u> </u>	<u> </u>	_		
B	Vancomycin	30 μg	≥17			≤1	_	· _	See general comment (4).
ACROLIDES	,	30 μg	_ ∠ 17		<u>. </u>	≥1		!	Oce general comment (4).
21) Not routine A	ely reported for organisms iso Erythromycin	lated from the υ 15 μg	ırinary tract ≥21	16–20	≤15	≤0.25	0.5	≥1	
0	Azithromycin	15 μg	≥18	14–17	<u>= 13</u> ≤13	<u>≤0.25</u>	1	≥2	
0	Clarithromycin	15 μg	≥21	17–20	≤16	≤0.25	0.5	<u></u> ≥1	
0	Dirithromycin	15 μg	≥18	14–17	<u>≤13</u>	<u>=</u> 0.25 ≤0.5	1	≥2	
22) Organism esistance.	s that are susceptible to tet	racycline are a	lso conside	ered suscept	ible to d				
В	Tetracycline	30 μg	≥28	25–27	≤24	oxycycline. H ≤1	lowever, resis	tance to ≥4	doxycycline cannot be inferred from tetracyc
В	Doxycycline				T				doxycycline cannot be inferred from tetracyc
В	Doxycycline IOLONES	30 μg	≥28	25–27 25–27	≤24	≤1 ≤0.25	2 0.5	≥4	
B LUOROQUIN B	Doxycycline IOLONES Gemifloxacin	30 μg 30 μg 5 μg	≥28 ≥28 ≥23	25–27 25–27 20–22	≤24 ≤24 ≤19	≤1 ≤0.25	2 0.5	≥4 ≥1 ≥0.5	(23) S. pneumoniae isolates susceptible to
B ELUOROQUIN B B	Doxycycline OLONES Gemifloxacin Levofloxacin	30 μg 30 μg 5 μg 5 μg	≥28 ≥28 ≥23 ≥17	25–27 25–27 20–22 14–16	≤24 ≤24 ≤19 ≤13	≤1 ≤0.25 ≤0.12 ≤2	2 0.5 0.25 4	≥4 ≥1 ≥0.5 ≥8	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to
B LUOROQUIN B	Doxycycline IOLONES Gemifloxacin	30 μg 30 μg 5 μg	≥28 ≥28 ≥23	25–27 25–27 20–22	≤24 ≤24 ≤19	≤1 ≤0.25	2 0.5	≥4 ≥1 ≥0.5	(23) S. pneumoniae isolates susceptible to
B LUOROQUIN B B B	Doxycycline OLONES Gemifloxacin Levofloxacin	30 μg 30 μg 5 μg 5 μg	≥28 ≥28 ≥23 ≥17	25–27 25–27 20–22 14–16	≤24 ≤24 ≤19 ≤13	≤1 ≤0.25 ≤0.12 ≤2 ≤1	2 0.5 0.25 4	≥4 ≥1 ≥0.5 ≥8	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin of moxifloxacin cannot be assumed to be
B LUOROQUIN B B B	Doxycycline OLONES Gemifloxacin Levofloxacin Moxifloxacin	30 μg 30 μg 5 μg 5 μg 5 μg	≥28 ≥28 ≥23 ≥17 ≥18	25–27 25–27 20–22 14–16 15–17	≤24 ≤24 ≤19 ≤13 ≤14	≤1 ≤0.25 ≤0.12 ≤2 ≤1	0.5 0.25 4 2	≥4 ≥1 ≥0.5 ≥8 ≥4	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin of moxifloxacin cannot be assumed to be
B LUOROQUIN B B B B	Doxycycline OLONES Gemifloxacin Levofloxacin Moxifloxacin Gatifloxacin	30 μg 30 μg 5 μg 5 μg 5 μg	≥28 ≥28 ≥28 ≥17 ≥18	25–27 25–27 20–22 14–16 15–17	≤24 ≤24 ≤19 ≤13 ≤14	≤1 ≤0.25 ≤0.12 ≤2 ≤1	2 0.5 0.25 4 2	≥4 ≥1 ≥0.5 ≥8 ≥4	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin moxifloxacin cannot be assumed to be
B ELUOROQUIN B B B O O O	Doxycycline OLONES Gemifloxacin Levofloxacin Moxifloxacin Gatifloxacin Ofloxacin	30 µg 30 µg 5 µg 5 µg 5 µg 5 µg 5 µg	≥28 ≥28 ≥28 ≥17 ≥18 ≥21 ≥21 ≥21	25–27 25–27 20–22 14–16 15–17 18–20 13–15	≤24 ≤24 ≤19 ≤13 ≤14 ≤17 ≤12	≤1 ≤0.25 ≤0.12 ≤2 ≤1 ≤1 ≤1	2 0.5 0.25 4 2	≥4 ≥1 ≥0.5 ≥8 ≥4 ≥4	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin of moxifloxacin cannot be assumed to be
B ELUOROQUIN B B B O O O	Doxycycline OLONES Gemifloxacin Levofloxacin Moxifloxacin Gatifloxacin Ofloxacin Sparfloxacin HWAY ANTAGONISTS Trimethoprim-	30 μg 30 μg 5 μg 5 μg 5 μg 5 μg 5 μg 5 μg	≥28 ≥28 ≥28 ≥17 ≥18 ≥21 ≥21 ≥21	25–27 25–27 20–22 14–16 15–17 18–20 13–15	≤24 ≤24 ≤19 ≤13 ≤14 ≤17 ≤12	≤1 ≤0.25 ≤0.12 ≤2 ≤1 ≤1 ≤1	2 0.5 0.25 4 2	≥4 ≥1 ≥0.5 ≥8 ≥4 ≥4	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin of moxifloxacin cannot be assumed to be
B LUOROQUIN B B B O O O O O OLATE PATI	Doxycycline OLONES Gemifloxacin Levofloxacin Moxifloxacin Gatifloxacin Ofloxacin Sparfloxacin WAY ANTAGONISTS	30 µg 30 µg 5 µg 5 µg 5 µg 5 µg 5 µg	≥28 ≥28 ≥23 ≥17 ≥18 ≥21 ≥16 ≥19	25–27 25–27 20–22 14–16 15–17 18–20 13–15 16–18	≤24 ≤24 ≤19 ≤13 ≤14 ≤17 ≤12 ≤15	≤1 ≤0.25 ≤0.12 ≤2 ≤1 ≤1 ≤1 ≤2 ≤0.5	2 0.5 0.25 4 2	≥4 ≥1 ≥0.5 ≥8 ≥4 ≥4 ≥8 ≥2	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin moxifloxacin cannot be assumed to be
B ELUOROQUIN B B B O O O O COLATE PATI	Doxycycline OLONES Gemifloxacin Levofloxacin Moxifloxacin Gatifloxacin Ofloxacin Sparfloxacin HWAY ANTAGONISTS Trimethoprim-	30 μg 30 μg 5 μg 5 μg 5 μg 5 μg 5 μg 5 μg	≥28 ≥28 ≥23 ≥17 ≥18 ≥21 ≥16 ≥19	25–27 25–27 20–22 14–16 15–17 18–20 13–15 16–18	≤24 ≤24 ≤19 ≤13 ≤14 ≤17 ≤12 ≤15	≤1 ≤0.25 ≤0.12 ≤2 ≤1 ≤1 ≤1 ≤2 ≤0.5	2 0.5 0.25 4 2	≥4 ≥1 ≥0.5 ≥8 ≥4 ≥4 ≥8 ≥2	(23) S. pneumoniae isolates susceptible to levofloxacin are predictably susceptible to gemifloxacin and moxifloxacin. However, S. pneumoniae susceptible to gemifloxacin moxifloxacin cannot be assumed to be

Table 2G. Streptococcus pneumoniae (Continued)

Test/Report	Antimicrobial	Disk	Zone Dia	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm Interpretive Categories and MIC Breakpoints, µg/mL						
Group	Agent	Content			R	Comments				
ANSAMYCIN	S									
С	Rifampin	5 μg	≥19	17–18	≤16	≤1		2	≥4	(24) Rx: Rifampin should not be used alone for antimicrobial therapy.
LINCOSAMID	ES									
В	Clindamycin	2 µg	≥19	16–18	≤15	≤0.25	0).5	≥1	(25) 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 before reporting clindamycin (see Table 3H, Subchapter 3.9 in M02, ³ and Subchapter 3.12 in M07 ¹). See comment (21).
STREPTOGR	AMINS									
0	Quinupristin-dalfopristin	15 μg	≥19	16–18	≤15	≤1		2	≥4	
OXAZOLIDIN	ONES									
С	Linezolid	30 μg	≥21	<u> </u>	_	≤2		_	<u> </u>	

Abbreviations: ATCC®, American Type Culture Collection; **NAD**, **β-nicotinamide adenine dinucleotide**; CAMHB, cation-adjusted Mueller-Hinton broth; CSF, cerebrospinal fluid; **ICR**, **inducible clindamycin resistance**; FDA, US Food and Drug Administration; I, intermediate; LHB, lysed horse blood; MHA, Mueller-Hinton agar; **MH-F agar**, **Mueller-Hinton fastidious agar**; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. $ATCC^{\otimes}$ is a registered trademark of the American Type Culture Collection.

References for Table 2G

- CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ³ CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 2H-1. Zone Diameter and MIC Breakpoints for Streptococcus spp. β-Hemolytic Group

Testing Conditions

Medium: Disk diffusion: MHA with 5% sheep blood

Broth dilution: CAMHB with LHB (2.5% to 5% v/v); the CAMHB should be supplemented to 50 µg/mL calcium for daptomycin (see M07¹ for instructions for

preparation of LHB)

Agar dilution: MHA with sheep blood (5% v/v); recent studies using the agar dilution

method have not been performed and reviewed by the subcommittee.

Inoculum: Colony suspension, equivalent to a 0.5 McFarland standard, using colonies from an

overnight (18- to 20-hour) sheep blood agar plate

Incubation: $35^{\circ}C \pm 2^{\circ}C$

Disk diffusion: 5% CO₂; 20-24 hours

Dilution methods: ambient air; 20-24 hours (CO₂ if necessary, for growth with agar

dilution)

Routine QC Recommendations (see Tables 4B and 5B for acceptable QC ranges)

S. pneumoniae ATCC®a 49619

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

Refer to Table 3H for additional testing recommendations, reporting suggestions, and QC.

General Comments

- (1) For disk diffusion, test a maximum of 9 disks on a 150-mm plate and 4 disks on a 100-mm plate. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk (see the M02 Disk Diffusion Reading Guide²). The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Do not measure the zone of inhibition of hemolysis. Measure the zones from the upper surface of the agar illuminated with reflected light, with the cover removed. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth.
- (2) For β-hemolytic streptococci when testing chloramphenicol, clindamycin, erythromycin, linezolid, tedizolid, and tetracycline by broth microdilution MIC, trailing growth can make end-point determination difficult. In such cases, read the MIC at the lowest concentration where the trailing begins. Tiny buttons of growth should be ignored (see M07,¹ Figures 3 and 4).
- (3) For this table, the β-hemolytic group includes the large colony–forming pyogenic strains of streptococci with group A (*S. pyogenes*), C, or G antigens and strains with Group B (*S. agalactiae*) antigen. Small colony–forming β-hemolytic strains with group A, C, F, or G antigens (*S. anginosus* group, previously termed "*S. milleri*") are considered part of the viridans group, and breakpoints for the viridans group should be used (see Table 2H-2).
- (4) Penicillin and ampicillin are drugs of choice for treatment of β-hemolytic streptococcal infections. Susceptibility testing of penicillins and other β-lactams approved by the US Food and Drug Administration for treatment of β-hemolytic streptococcal infections does not need to be performed routinely, because nonsusceptible isolates (ie, penicillin MICs > 0.12 and ampicillin MICs > 0.25 μg/mL) are extremely rare in any β-hemolytic streptococcus and have not been reported for *S. pyogenes*. If testing is performed, any β-hemolytic streptococcal isolate found to be nonsusceptible should be re-identified, retested, and, if confirmed, submitted to a public health laboratory. See Appendix A for additional instructions.

Table 2H-1. Streptococcus spp. β-Hemolytic Group (Continued)

(5) Breakpoints for *Streptococcus* spp. β-hemolytic group are proposed based on population distributions of various species, pharmacokinetics of the antimicrobial agents, previously published literature, and the clinical experience of subcommittee members. Systematically collected clinical data were not available for review with many of the antimicrobial agents in this table.

Test/Report	Fest/Report Antimicrobial Group Agent		Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm Content S I R					ries and ts,	Comments
	Agent	Content	S	<u> </u>	R	S	<u> </u>	R	Comments
PENICILLINS									
									pproved indications and does not need to be teste
									oxicillin-clavulanate, ampicillin-sulbactam, cefazolir
						i, ertapenem	, and merop	penem. Fo	r group A β-hemolytic streptococci, penicillin is als
	cefaclor, cefdinir, cefprozil, ceftib Penicillin or	10 units		cerpodoxime •	<u>. </u>	10.40		, 	Con general comment (4)
A A		-	≥24	<u> </u>		≤0.12	_	-	See general comment (4).
	ampicillin	10 μg	≥24	<u> </u>	: -	≤0.25	: -	<u>: </u>	
CEPHEMS (PA	RENTERAL) (Including cephal	osporins I,	II, III, and I	v. Please re	ter to Glos	ssary I.)			
0 / //	2)								
See comment (- 1	0.0	. 04	:	:		:		T
B B	Cefepime or cefotaxime or	30 μg	≥24	<u> </u>	; –	≤0.5	-	-	
В	celotaxime of ceftriaxone	30 μg	≥24	<u> </u>	: ⁻	≤0.5	_	: -	
		30 μg	≥24	: -	; -	≤0.5		; -	(7) 5
С	Ceftaroline	30 μg	≥26	_	_	≤0.5	_	_	(7) Breakpoints are based on a dosage regimen of 600 mg administered every 12 h.
CARBAPENEN	IS								
See comment (6).								
0	Doripenem	_	_	<u> </u>	. –	≤0.12	-	-	
0	Ertapenem	_	_	-	-	≤1	-	-	
0	Meropenem	_	_	-	. –	≤0.5	_	-	
GLYCOPEPTIC	DES .			•		•	-	•	
В	Vancomycin	30 μg	≥17	_	_	≤1	_	-	
LIPOGLYCOPE	PTIDES			•			•		
C	Dalbavancin	_	_	: -		≤0.25	-	: -	(8) For reporting against S. pyogenes,
Ü	Balbavarioni				į	-0.20			S. agalactiae, and S. dysgalactiae.
С	Oritavancin	_	_			≤0.25	_	-	o. againetiae, and o. ajoguidondo.
C	Telavancin	_	_	: _	!	<u>≤0.20</u> ≤0.12	-	;	
LIPOPEPTIDES						_ = 0.12	'		
C	Daptomycin	_		· _	-	≤1	_	; _	(9) Daptomycin should not be reported for
J	Daptomyon		_	<u> </u>	_	_ =	_	-	isolates from the respiratory tract.
		ı	I.		:	1			

See comment (11).

С

Chloramphenicol

30 μg

≥21 18–20

≤17

≤4

8

≥16

M100, 30th ed.

Test/Report	Antimicrobial	Disk	Zone Dia	etive Categor ameter Brea arest whole	kpoints, mm	MIC	ive Categor Breakpoin μg/mL	its,	
Group	Agent	Content	S	<u> </u>	R	S	<u> </u>	R	Comments
MACROLIDES									
10) Susceptibil	ity and resistance to azithrom	ycin, clarithror	nycin, and	dirithromycin	ı can be pre	edicted by te	sting erythro	omycin.	
11) Not routine	ly reported on isolates from the	ne urinary trac	t.						
A	Erythromycin	15 μg	≥21	16–20	≤15	≤0.25	0.5	≥1	(12) Rx: Recommendations for intrapartum prophylaxis for group B streptococci are penicilli or ampicillin. Although cefazolin is recommended for penicillin-allergic women at low risk for anaphylaxis, those at high risk for anaphylaxis may receive clindamycin. Group B streptococci are susceptible to ampicillin, penicillin, and cefazolin, but may be resistant to erythromycin and clindamycin. When a group B Streptococcus is isolated from a pregnant woman with severe penicillin allergy (high risk for anaphylaxis), erythromycin and clindamycin (including ICR) should be tested, and only clindamycin should breported. Erythromycin should be tested for ICR determination only and should not be reported. See Table 3H.
0	Azithromycin	15 μg	≥18	14–17	≤13	≤0.5	1	≥2	
0	Clarithromycin	15 μg	≥ 21	17–20	≤16	≤0.25	0.5	≥1	
0	Dirithromycin	15 μg	≥ 18	14–17	≤13	≤0.5	1	≥2	
nferred from te	that are susceptible to tetracy tracycline resistance.						·		esistance to doxycycline and minocycline cannot be
O FLUOROQUIN	Tetracycline	30 μg	≥23	19–22	≤18	≤2	4	! ≥8	
C	Levofloxacin	T 5.10	>17	14–16	<12	-2	4	, , 0	
0	Gatifloxacin	5 μg	≥17 ≥21	18–20	≤13 ≤17	≤2 ≤1	2	≥8 ≥4	
0	Gatilloxacin	5 μg	≥21 ≥19	16–20	≤17 ≤15	≤ 1 ≤0.5	1	≥4 ≥2	+
0	Ofloxacin	5 μg		13–15			4	_	+
	· · · · · · · · · · · · · · · · · · ·	5 μ g	≥16		≤12	≤2		≥8	
0	Trovafloxacin	10 μg	≥19	16–18	≤15	<1	2	≥4	

Test/Report	Antimicrobial	Disk	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm		Interpretive Categories and MIC Breakpoints, µg/mL					
Group	Agent	Content	S	1	R	S	1		R	Comments
LINCOSAMIDE	S									
Α	Clindamycin	2 μg	≥19	16–18	≤15	≤0.25	0.5		≥1	See comments (11) and (12).
										(14) 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 before reporting clindamycin. See Table 3H, Subchapter 3.9 in M02,3 and Subchapter 3.12 in M07.1
STREPTOGRA	MINS									
0	Quinupristin-dalfopristin	15 μg	≥19	16–18	≤15	≤1	2	1	≥4	(15) Report against S. pyogenes.
OXAZOLIDINO	NES		•				•	, and the second	•	
С	Linezolid	30 μg	≥21	<u>-</u>	! -	≤2	-	-	_	
С	Tedizolid	_	-	-	-	≤0.5	_		-	(16) For reporting against <i>S. pyogenes</i> and <i>S. agalactiae</i> only.

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; **ICR**, **inducible clindamycin resistance**; I, intermediate; LHB, lysed horse blood; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

References for Table 2H-1

- CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. M02 Disk Diffusion Reading Guide. 1st ed. CLSI quick guide M02QG. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 3 CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

M100, 30th ed

Table 2H-2. Zone Diameter and MIC Breakpoints for Streptococcus spp. Viridans Group

Testing Conditions

Medium: Disk diffusion: MHA with 5% sheep blood

Broth dilution: CAMHB with LHB (2.5% to 5% v/v); the CAMHB should be supplemented to 50 μg/mL calcium for daptomycin (see M07¹ for instructions for

preparation of LHB)

Agar dilution: MHA with sheep blood (5% v/v); recent studies using the agar dilution

method have not been performed and reviewed by the subcommittee.

Inoculum: Colony suspension, equivalent to a 0.5 McFarland standard using colonies from an

overnight (18- to 20-hour) sheep blood agar plate

Incubation: $35^{\circ}C \pm 2^{\circ}C$

Disk diffusion: 5% CO₂; 20-24 hours

Dilution methods: ambient air: 20–24 hours (CO₂ if necessary for growth with agar

dilution)

Routine QC Recommendations (see Tables 4B and 5B for acceptable QC ranges)

S. pneumoniae ATCC®a 49619

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) For disk diffusion, measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk. The zone margin should be considered the area showing no obvious, visible growth that can be detected with the unaided eye. Do not measure the zone of inhibition of hemolysis. Measure the zones from the upper surface of the agar illuminated with reflected light, with the cover removed. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth.
- (2) For viridans streptococci when testing chloramphenicol, clindamycin, erythromycin, linezolid, tedizolid, and tetracycline by broth microdilution MIC, trailing growth can make end-point determination difficult. In such cases, read the MIC at the lowest concentration where the trailing begins. Tiny buttons of growth should be ignored (see M07, Figures 3 and 4).
- (3) The viridans group of streptococci includes the following five groups, with several species within each group: *mutans* group, *salivarius* group, *bovis* group, *anginosus* group (previously "S. *milleri*" group), and *mitis* group. The *anginosus* group includes small colony–forming β-hemolytic strains with groups A, C, F, and G antigens. For detailed information on the species within the groups, please refer to recent literature.
- (4) Breakpoints for *Streptococcus* spp. viridans group are proposed based on population distributions of various species, pharmacokinetics of the antimicrobial agents, previously published literature, and the clinical experience of subcommittee members. Systematically collected clinical data were not available for review with many of the antimicrobial agents in this table.

Table 2H-2. Streptococcus spp. Viridans Group (Continued)

Test/Report	Antimicrobial			ts,					
Group	Agent	Content	S	1	R	S	<u> </u>	R	Comments
PENICILLINS									
A A	Penicillin Ampicillin	_	_		-	≤0.12 ≤0.25	0.25–2 0.5–4	≥4 ≥8	(5) Viridans streptococci isolated from normally sterile anatomical sites (eg, CSF, blood, bone) should be tested for penicillin susceptibility using an MIC method.
									(6) A penicillin MIC of ≤0.125 μg/mL is the same as a penicillin MIC of ≤0.12 μg/mL and both should be interpreted as susceptible. Laboratories should report an MIC of ≤0.125 μg/mL as ≤0.12 μg/mL.
				:			· · ·		(7) Rx: Penicillin- or ampicillin-intermediate isolates may necessitate combined therapy with an aminoglycoside for bactericidal actio
β-LACTAM CO	MBINATION AGENTS								
С	Ceftolozane-tazobactam	_	_	-	-	≤8/4	16/4	≥32/4	(8) Breakpoints are based on a dosage regimen of 1.5 g administered every 8 h.
CEPHEMS (PA	RENTERAL) (Including cept	nalosporins I,	II, III, and I	V. Please ref	er to Gloss	ary I.)			
В	Cefepime	30 μg	≥24	22–23	≤21	≤1	2	≥4	
В	Cefotaxime	30 μg	≥28	26–27	≤25	≤1	2	≥4	
В	Ceftriaxone	30 μg	≥27	25–26	≤24	≤1	2	≥4	
CARBAPENEN	IS								
0	Doripenem	_	_	<u> </u>	-	≤1	<u> </u>	: -	
0	Ertapenem	_	_	_	_	≤1	_	-	
0	Meropenem	_	_	<u> </u>	_	≤0.5	<u> </u>	† -	
GLYCOPEPTID	DES								
В	Vancomycin	30 μg	≥17	<u> </u>	· –	≤1	. –	<u>:</u> -	
LIPOGLYCOPE	PTIDES		•						
С	Dalbavancin	_	_	-	_	≤0.25	-	-	(9) For reporting against <i>S. anginosus</i> group (includes <i>S. anginosus</i> , <i>S. intermedius</i> , and <i>S. constellatus</i>) only.
С	Oritavancin	_	_	 	_	≤0.25	<u> </u>	† -	
C	Telavancin	_	_	-		≤0.06	. –	: -	
LIPOPEPTIDES			<u> </u>			0.00			
0	Daptomycin	_	_	-	-	≤1	-	<u> </u>	(10) Daptomycin should not be reported for isolates from the respiratory tract.

See comment (12).

See comment (12).

See comment (9).

≥16

 \geq 4

Test/Report	st/Report Antimicrobial		Zone I	e Catego eter Brea st whole	Interpretive Categories and MIC Breakpoints, µg/mL							
Group	Agent	Content	S	i	ı	R	S	;	ı	•	R	Comments
MACROLIDES							•					
(11) Susceptibil	ity and resistance to azithromy	cin, clarithrom	ycin, and	dirith	romycin c	an be predi	cted by test	ing e	erythrom	ycin.		
(12) Not routine	ly reported on isolates from the	urinary tract.										
С	Erythromycin	15 μg	≥21	į	16–20	≤15	≤0.25	į	0.5	i	≥1	
0	Azithromycin	15 μg	≥18		14–17	≤13	≤0.5	i	1	- 1	≥2	
0	Clarithromycin	15 μg	≥21	-	17–20	≤16	≤0.25	-	0.5		≥1	
0	Dirithromycin	15 μg	≥18		14–17	≤13	≤0.5		1		≥2	
inferred from tet	that are susceptible to tetracy tracycline resistance.	T		ed su	•			осу	cline. Ho	weve		stance to doxycycline and minocycline cannot be
0	Tetracycline	30 μg	≥23	<u> </u>	19–22	: ≤18	≤2	<u>:</u>	4	<u>:</u>	≥8	
FLUOROQUING	OLONES											
0	Levofloxacin	5 μg	≥17	-	14–16	: ≤13	≤2	- ;	4	:	≥8	
0	Ofloxacin	5 μg	≥16		13–15	≤12	≤2		4	i_	≥8	
0	Gatifloxacin	5 μg	≥21		18–20	≤17	≤1	- !	2		≥4	
0	Grepafloxacin	5 μg	≥19		16–18	≤15	≤0.5	i	1		≥2	
0	Trovafloxacin	10 μg	≥19		16–18	≤15	≤1	i	2		≥4	
PHENICOLS			•				•	-		-		

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; CSF, cerebrospinal fluid; I, intermediate; LHB, lysed horse blood; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

≤17

≤15

≤15

≤4

≤0.25

≤1

≤2

≤0.25

0.5

Footnote

18-20

16-18

16-18

a. ATCC® is a registered trademark of the American Type Culture Collection.

30 μg

2 μα

15 μg

30 µg

≥21

≥19

≥19

≥21

Table 2H-2. Streptococcus spp. Viridans Group (Continued)

Reference for Table 2H-2

С

LINCOSAMIDES

STREPTOGRAMINS

OXAZOLIDINONES

С

Chloramphenicol

Quinupristin-dalfopristin

Clindamycin

Linezolid

Tedizolid

¹ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

M100, 30th ed

Table 2I. Zone Diameter and MIC Breakpoints for Neisseria meningitidis

Testing Conditions

Medium: Disk diffusion: MHA with 5% sheep blood

Broth microdilution: CAMHB supplemented with LHB (2.5% to

5% v/v) (see M07¹ for preparation of LHB)

Agar dilution: MHA supplemented with sheep blood (5% v/v)

Inoculum: Colony suspension from 20–24 hours growth from chocolate agar

incubated at 35°C; 5% CO₂; equivalent to a 0.5 McFarland standard. Colonies grown on sheep blood agar may be used for inoculum preparation. However, the 0.5 McFarland suspension obtained from sheep blood agar will contain approximately 50% fewer CFU/mL. This must be considered when preparing the final

dilution before panel inoculation, as guided by colony counts.

Incubation: $35^{\circ}C \pm 2^{\circ}C$; 5% CO_2 ; 20-24 hours

Routine QC Recommendations (See Tables 4A-1, 4B, 5A-1, and 5B for acceptable QC ranges.)

Streptococcus pneumoniae ATCC®a 49619:

Disk diffusion: incubate in 5% CO₂.

Broth microdilution: incubate in ambient air or CO₂ (except azithromycin QC tests that must be incubated in ambient air).

E. coli ATCC® 25922

Disk diffusion, broth microdilution or agar dilution for ciprofloxacin, nalidixic acid, minocycline, and sulfisoxazole: incubate in ambient air or CO₂.

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

Important: For complete information on safety precautions, see *Biosafety in Microbiological and Biomedical Laboratories*. 5th ed. Washington, DC: US Department of Health and Human Services; 2009. http://www.cdc.gov/biosafety/publications/bmbl5/. Accessed 10 December 2019.

- (1) Recommended precautions: Perform all AST of *N. meningitidis* in a BSC. Manipulating *N. meningitidis* outside a BSC is associated with increased risk for contracting meningococcal disease. Laboratory-acquired meningococcal disease is associated with a case fatality rate of 50%. Exposure to droplets or aerosols of *N. meningitidis* is the most likely risk for laboratory-acquired infection. Rigorous protection from droplets or aerosols is mandated when microbiological procedures (including AST) are performed on all *N. meningitidis* isolates.
- (2) If a BSC is unavailable, manipulation of these isolates should be minimized, limited to Gram staining or serogroup identification using phenolized saline solution, while wearing a laboratory coat and gloves and working behind a full face splash shield. Use BSL-3 practices, procedures, and containment equipment for activities with a high potential for droplet or aerosol production and for activities involving production quantities or high concentrations of infectious materials. If BSL-2 or BSL-3 facilities are not available, forward isolates to a referral or public health laboratory with a minimum of BSL-2 facilities.
- (3) Laboratorians who are exposed routinely to potential aerosols of *N. meningitidis* should consider vaccination according to the current recommendations of the Centers for Disease Control and Prevention Advisory Committee on Immunization Practices, available at http://www.cdc.gov/vaccines/acip/index.html.

Table 2I. Neisseria meningitidis (Continued)

- (4) For disk diffusion, test a maximum of 5 disks on a 150-mm plate and 2 disks on a 100-mm plate. Measure the diameter of the zones of complete inhibition (as judged by the unaided eye), including the diameter of the disk. Measure the zones from the upper surface of the agar illuminated with reflected light, with the cover removed. Ignore faint growth of tiny colonies that can be detected only with a magnifying lens at the edge of the zone of inhibited growth. With trimethoprim and the sulfonamides, antagonists in the medium may allow some slight growth; therefore, disregard slight growth (20% or less of the lawn of growth) and measure the more obvious margin to determine the zone diameter.
- (5) Breakpoints are based on population distributions of MICs of various agents, pharmacokinetics of the agents, previously published literature, and the clinical experience of subcommittee members. Systematically collected clinical data were not available to review with many of the antimicrobial agents in this table.
- (6) With azithromycin, breakpoints were developed initially using MICs determined by incubation in ambient air for the pharmacodynamic calculations.

Test/Report	Antimicrobial	Disk	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm			etive Categor C Breakpoin µg/mL					
Group	Agent	Content	S	I	R	S	ı	R	Comments		
PENICILLINS	PENICILLINS										
С	Penicillin		_	_	ļ –	≤0.06	0.12-0.25	≥0.5			
С	Ampicillin		-	<u> </u>	! –	≤0.12	0.25–1	≥2			
CEPHEMS											
С	Cefotaxime or	30 μg	≥34	-	-	≤0.12	-	-			
С	ceftriaxone	30 μg	≥34	<u> </u>	<u> </u>	≤0.12	_	i –			
CARBAPENE	MS										
С	Meropenem	10 μg	≥30	-	-	≤0.25	_	-			
MACROLIDES											
С	Azithromycin	15 μg	≥20	-	-	≤2	-	: -	See general comment (6).		
									(7) May be appropriate only for prophylaxis of meningococcal case contacts. These breakpoints		
					:			:	do not apply to therapy of patients with invasive meningococcal disease.		
TETRACYCLIN	NES								1 0		
С	Minocycline	30 μg	≥26	1 -	-	≤2	_	-	See comment (7).		
	FLUOROQUINOLONES (8) For surveillance purposes, a nalidixic acid MIC ≥ 8 μg/mL or a zone ≤ 25 mm may correlate with diminished fluoroquinolone susceptibility.										
C	Ciprofloxacin	5 μg	≥35	33–34	≤32	≤0.03	0.06	≥0.12	See comment (7).		
С	Levofloxacin	-	_	_	_	≤0.03	0.06	≥0.12			

Table 2I. Nei	isseria meningitidis (Ce	ontinued)							
Test/Report	Antimicrobial	Disk	Zone Di	Interpretive Categories and Zone Diameter Breakpoints, nearest whole mm		Interpret	ive Categories Breakpoints, µg/mL	and MIC	
Group	Agent	Content	S	1	R	S	; 'J	R	Comments
FOLATE PATH	HWAY ANTAGONISTS								
С	Sulfisoxazole	_	-		_	≤2	4	≥8	See comment (7).
С	Trimethoprim-	1.25/	≥30	: 26–29	: ≤25	≤0.12/	0.25/4.75	≥ 0.5/	(9) Trimethoprim-sulfamethoxazole is the
	sulfamethoxazole	23.75 μg				2.4		9.5	preferred disk for detection of sulfonamide resistance. Trimethoprim-sulfamethoxazole testing predicts susceptibility and resistance to trimethoprim-sulfamethoxazole and sulfonamides. Sulfonamides may be appropriate only for prophylaxis of meningococcal case contacts.
PHENICOLS									
С	Chloramphenicol	30 μg	≥26	20–25	≤19	≤2	4	≥8	(10) Not routinely reported on isolates from the urinary tract.
ANSAMYCINS									
С	Rifampin	5 μα	≥25	20–24	≤19	≤0.5	1	≥2	See comment (7).

Abbreviations: AST, antimicrobial susceptibility testing; ATCC®, American Type Culture Collection; BSC, biological safety cabinet; BSL-2, biosafety level 2; BSL-3, biosafety level 3; CAMHB, cation-adjusted Mueller-Hinton broth; CFU, colony-forming unit(s); I, intermediate; LHB, lysed horse blood; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection.

Reference for Table 2I

CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 2J. MIC Breakpoints for Anaerobes

Testing Conditions

Medium: Agar dilution (for all anaerobes): Brucella agar supplemented

with hemin (5 μ g/mL), vitamin K₁ (1 μ g/mL), and laked sheep

blood (5% v/v)

Broth microdilution (for *Bacteroides* spp. and *Parabacteroides* spp. only): Brucella broth supplemented with hemin (5 µg/mL),

vitamin K_1 (1 μ g/mL), and LHB (5% ν / ν)

Inoculum: Broth culture method or colony suspension, equivalent to

0.5 McFarland suspension Agar: 10⁵ CFU per spot Broth: 10⁶ CFU/mL

Incubation: 36°C±1°C, anaerobically

Broth microdilution: 46–48 hours Agar dilution: 42–48 hours **Routine QC Recommendations** (see Tables 5D and 5E for acceptable QC ranges)

Test one or more of the following organisms. The choice and number of QC strains tested should be based on obtaining on-scale end points for the antimicrobial agent tested.

B. fragilis ATCC[®] 25285 Bacteroides thetaiotaomicron ATCC[®] 29741 Clostridioides (formerly Clostridium) difficile ATCC[®] 700057 Eggerthella lenta (formerly Eubacterium lentum) ATCC[®] 43055

When a commercial test system is used for susceptibility testing, refer to the manufacturer's instructions for QC test recommendations and QC ranges.

General Comments

- (1) For isolates for which the antimicrobial agent MICs fall within the intermediate category, maximum dosages, along with proper ancillary therapy, should be used to achieve the best possible levels of drug in abscesses and/or poorly perfused tissues. If this approach is taken, organisms for which the antimicrobial agent MICs fall within the susceptible range are generally amenable to therapy. Organisms for which the antimicrobial agent MICs are in the intermediate range may respond, but in such cases, efficacy as measured by patient clinical response should be carefully monitored. Ancillary therapy, such as drainage procedures and debridement, are of great importance for proper management of anaerobic infections.
- (2) Refer to Figures 3 and 4 in CLSI document M11¹ for examples of reading end points.
- (3) MIC values using either Brucella blood agar or Wilkins Chalgren agar (former reference medium) are considered equivalent.
- (4) Broth microdilution is recommended only for testing *Bacteroides* spp. and *Parabacteroides* spp. MIC values for agar or broth microdilution are considered equivalent for those species.
- (5) Until additional studies are performed to validate broth microdilution for testing other organisms, it should be used only for testing members of *Bacteroides* spp. and *Parabacteroides* spp.

For Use With M11

Table 2J. Anaerobes (Continued)

Test/Report	Antimicrobial	Interp	retive Categorie Breakpoints µg/mL		
Group	Agent	S	ı	R	Comments
PENICILLINS					
A/C A/C	Ampicillin ^b Penicillin ^b	≤0.5 ≤0.5	1	≥2 ≥2	 (6) Ampicillin and penicillin are recommended for primary testing and reporting for grampositive organisms (group A) because most of them are β-lactamase negative, but not for gram-negative organisms (group C) because many are β-lactamase positive. (7) Bacteroides spp. are intrinsically resistant to penicillin and ampicillin. Parabacteroides spp. are presumed to be resistant to penicillin and ampicillin. Other gram-negative and gram-positive anaerobes may be screened for β-lactamase activity with a chromogenic cephalosporin; if β-lactamase positive, report as resistant to penicillin, ampicillin, and amoxicillin. Be aware that β-lactamase–negative isolates may be resistant to β-lactams by other mechanisms. Because higher blood levels are achievable with these antimicrobial agents, infection with non–β-lactamase-producing organisms with higher MICs (2–4 μg/mL) with adequate dosage regimen might be treatable. (8) Results of ampicillin testing can be used to predict results for amoxicillin.
0	Piperacillin	≤32	64	≥128	(6) Results of amplifiant testing can be used to predict results for amoxicians.
<u> </u>	OMBINATION AGENTS		,	120	
A	Amoxicillin-clavulanate	≤4/2	8/4	≥16/8	
A	Ampicillin-sulbactam	≤8/4	16/8	≥32/16	
A	Piperacillin-tazobactam	≤16/4	32/4–64/4	≥128/4	
0	Ticarcillin-clavulanate	≤32/2	64/2	≥128/2	
CEPHEMS (PA	ARENTERAL) (Including cep		•		er to Glossary I.)
C	Cefotetan	<u>≤</u> 16	32	≥64	
С	Cefoxitin	≤16	32	≥64	
С	Ceftizoxime	≤32	64	≥128	
C	Ceftriaxone	≤16	32	≥64	
0	Cefmetazole	≤16	32	≥64	
0	Cefoperazone	≤16	32	≥64	
0	Cefotaxime	≤16	32	≥64	
CARBAPENE	MS				
Α	Doripenem	≤2	4	. ≥8	
Α	Ertapenem	≤4	8	≥16	
Α	Imipenem	≤4	8	≥16	
Α	Meropenem	≤4	8	≥16	
TETRACYCLI	INES				
С	Tetracycline	≤4	8	≥16	
FLUOROQUIN					
С	Moxifloxacin	≤2	4	≥8	

Table 2J. Anaerobes (Continued)

14510 2017111	able 20. Anacrobes (Continued)												
Test/Report	Fest/Report Antimicrobial			ive Cate Breakp μg/mL	oin	ries and ts,							
Group	Agent	S	į	ı	i	R	Comments						
LINCOSAMIDE	S												
Α	Clindamycin	≤2	;	4	-	≥8							
PHENICOLS													
С	Chloramphenicol	≤8	i	16	÷	≥32							
NITROIMIDAZO	OLES												
Α	Metronidazole	≤8	į	16		≥32	(9) Many non–spore-forming, gram-positive anaerobic rods are resistant to metronidazole.						

Abbreviations: ATCC®, American Type Culture Collection; CFU, colony-forming unit(s); I, intermediate; LHB, lysed horse blood; MIC, minimal inhibitory concentration; QC, quality control; R, resistant; S, susceptible.

Footnotes

- a. ATCC® is a registered trademark of the American Type Culture Collection.
- b. A/C: Group A for gram-positive anaerobes and group C for gram-negative organisms. Refer to Table 1C.

Reference for Table 2J

¹ CLSI. *Methods for Antimicrobial Susceptibility Testing of Anaerobic Bacteria.* 9th ed. CLSI standard M11. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

For Use With M11

Table 3A. Tests for Extended-Spectrum β-Lactamases in Klebsiella pneumoniae, Klebsiella oxytoca, Escherichia coli, and Proteus mirabilis

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 no longer necessary before reporting results (ie, it is no longer necessary to edit results for cephalosporins, aztreonam, or penicillins to resistant). However, ESBL testing may still be useful for epidemiological or infection **prevention** purposes. For laboratories that have not implemented the current breakpoints, ESBL testing should be performed as described in this table.

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.

Test	Criteria for Performa	nce of ESBL Test	ESBL 1	Test
Test method	Disk diffusion	Broth microdilution	Disk diffusion	Broth microdilution
Medium	MHA	CAMHB	MHA	САМНВ
Antimicrobial concentration	For K. pneumoniae, K. oxytoca, and E. coli: Cefpodoxime 10 μg or Ceftazidime 30 μg or Aztreonam 30 μg or Cefotaxime 30 μg or Ceftriaxone 30 μg For P. mirabilis: Cefpodoxime 10 μg or Ceftazidime 30 μg or Ceftazidime 30 μg (Testing more than one antimicrobial agent improves the sensitivity of ESBL detection.)	For K. pneumoniae, K. oxytoca, and E. coli: Cefpodoxime 4 μg/mL or Ceftazidime 1 μg/mL or Aztreonam 1 μg/mL or Cefotaxime 1 μg/mL or Ceftriaxone 1 μg/mL For P. mirabilis: Cefpodoxime 1 μg/mL or Ceftazidime 1 μg/mL or Ceftazidime 1 μg/mL (Testing more than one antimicrobial agent improves the sensitivity of ESBL detection.)	Ceftazidime 30 μg Ceftazidime-clavulanate ^a 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.)	Ceftazidime 0.25–128 μg/mL Ceftazidime-clavulanate 0.25/4–128/4 μg/mL and Cefotaxime 0.25–64 μg/mL Cefotaxime-clavulanate 0.25/4–64/4 μg/mL (Testing necessitates using both cefotaxime and ceftazidime, alone and in combination with clavulanate.)
Inoculum	Standard disk diffusion procedure	Standard broth dilution procedure	Standard disk diffusion procedure	Standard broth dilution procedure
Incubation conditions	35°C±2°C; ambient air	35°C±2°C; ambient air	35°C±2°C; ambient air	35°C±2°C; ambient air
Incubation length	16–18 hours	16–20 hours	16–18 hours	16–20 hours

Table 3A. (Continued)

Test	Criteria for Performa	ance of ESBL Test	ESBL 1	Test Test
Test method	Disk diffusion	Broth microdilution	Disk diffusion	Broth microdilution
Results	For K. pneumoniae, K. oxytoca, and E. coli: Cefpodoxime zone ≤17 mm Ceftazidime zone ≤22 mm Aztreonam zone ≤27 mm Cefotaxime zone ≤27 mm Ceftriaxone zone ≤25 mm For P. mirabilis: Cefpodoxime zone ≤22 mm Ceftazidime zone ≤22 mm Cefotaxime zone ≤27 mm Zones above may indicate ESBL production.	Growth at or above the concentrations listed may indicate ESBL production (ie, for <i>E. coli, K. pneumoniae</i> , and <i>K. oxytoca</i> , MIC ≥8 µg/mL for cefpodoxime or MIC ≥2 µg/mL for ceftazidime, aztreonam, cefotaxime, or ceftriaxone; and for <i>P. mirabilis</i> , MIC ≥2 µg/mL for cefpodoxime, ceftazidime, or cefotaxime).	A ≥5-mm 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).	A \geq 3 twofold concentration decrease in an MIC for either antimicrobial agent tested in combination with clavulanate vs the MIC of the agent when tested alone = ESBL (eg, ceftazidime MIC = 8 μ g/mL; ceftazidime-clavulanate MIC = 1 μ g/mL).
Reporting			For all confirmed ESBL-producing strategy of the strategy of t	nalosporin and aztreonam ould be reported as resistant for reonam. rin and aztreonam breakpoints,

Table 3A. (Continued)

Test	Criteria for Perform	ance of ESBL Test	ESBL	Test
Test method	Disk diffusion	Broth microdilution	Disk diffusion	Broth microdilution
QC recommendations	When testing antimicrobial agents used for ESBL detection, <i>K. pneumoniae</i> ATCC®b 700603 is provided as a supplemental QC strain (eg, for training, competence assessment, or test evaluation). Either strain, <i>K. pneumoniae</i> ATCC® 700603 or <i>E. coli</i> ATCC® 25922, may then be used for routine QC (eg, weekly or daily).	When testing antimicrobial agents used for ESBL detection, <i>K. pneumoniae</i> ATCC® 700603 is provided as a supplemental QC strain (eg, for training, competence assessment, or test evaluation). Either strain, <i>K. pneumoniae</i> ATCC® 700603 or <i>E. coli</i> ATCC® 25922, may then be used for routine QC (eg, weekly or daily).	When performing the ESBL test, K. pneumoniae ATCC® 700603 and E. coli ATCC® 25922 should be used for routine QC (eg, weekly or daily).	When performing the ESBL test, <i>K. pneumoniae</i> ATCC® 700603 and <i>E. coli</i> ATCC® 25922 should be tested routinely (eg, weekly or daily).
	E. coli ATCC® 25922 (see acceptable QC ranges in Table 4A-1)	E. coli ATCC® 25922 = no growth (see acceptable QC ranges listed in Table 5A-1)	Acceptable QC: E. coli ATCC® 25922: ≤2-mm increase in zone diameter for antimicrobial agent tested in combination with clavulanate vs the zone diameter when tested alone.	Acceptable QC: E. coli ATCC® 25922: <3 twofold concentration decrease in MIC for antimicrobial agent tested in combination with clavulanate vs the MIC of the agent when tested alone.
Abbreviskings ATOOR	K. pneumoniae ATCC® 700603: Cefpodoxime zone 9–16 mm Ceftazidime zone 10–18 mm Aztreonam zone 10–16 mm Cefotaxime zone 17–25 mm Ceftriaxone zone 16–24 mm	$\it K. pneumoniae ATCC^{\$}$ 700603 = Growth: Cefpodoxime MIC ≥8 μg/mL Ceftazidime MIC ≥2 μg/mL Aztreonam MIC ≥2 μg/mL Cefotaxime MIC ≥2 μg/mL Ceftriaxone MIC ≥2 μg/mL	K. pneumoniae ATCC® 700603: ≥5-mm increase in zone diameter of ceftazidime- clavulanate vs ceftazidime alone; ≥3-mm increase in zone diameter of cefotaxime- clavulanate vs cefotaxime alone.	K. pneumoniae ATCC® 700603: ≥3 twofold concentration decrease in MIC for an antimicrobial agent tested in combination with clavulanate vs the MIC of the agent when tested alone.

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; ESBL, extended-spectrum β-lactamase; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; PK-PD, pharmacokinetic-pharmacodynamic; QC, quality control.

Footnotes

- a. Preparation of ceftazidime-clavulanate (30 μg/10 μg) and cefotaxime-clavulanate (30 μg/10 μg) disks: Using a stock solution of clavulanate at 1000 μg/mL (either freshly prepared or taken from small aliquots that have been frozen at -70°C), add 10 μL of clavulanate to ceftazidime (30 μg) and cefotaxime (30 μg) disks. Use a micropipette to apply the 10 μL of stock solution to the ceftazidime and cefotaxime disks within one hour before they are applied to the plates, allowing about 30 minutes for the clavulanate to absorb and the disks to be dry enough for application. Use disks immediately after preparation or discard; do not store.
- b. ATCC® is a registered trademark of the American Type Culture Collection.

Introduction to Tables 3B and 3C. Tests for Carbapenemases in Enterobacterales and Pseudomonas aeruginosa

Institutional infection **prevention** procedures or epidemiological investigations may necessitate identification of carbapenemase-producing **Enterobacterales** and *P. aeruginosa*. Such testing is not currently recommended for routine use.

Carbapenemase-producing isolates of **Enterobacterales** usually test intermediate or resistant to one or more carbapenems using the current breakpoints as listed in Table 2A (**NOTE: Testing not susceptible to ertapenem is often the most sensitive indicator of carbapenemase production**) and usually test resistant to one or more agents in cephalosporin subclass III (eg, cefoperazone, cefotaxime, ceftazidime, ceftizoxime, and ceftriaxone). However, some isolates that produce carbapenemases such as SME or IMI often test susceptible to these cephalosporins.

Laboratories using **Enterobacterales** MIC breakpoints for carbapenems described in M100-S20 (January 2010) should perform the CarbaNP test, mCIM, eCIM, and/or a molecular assay (refer to Tables 3B and 3C for methods) when isolates of **Enterobacterales** are suspicious for carbapenemase production based on imipenem or meropenem MICs 2–4 µg/mL or ertapenem MIC 2 µg/mL (refer to Tables 3B-1 and 3C-1 for guidance on reporting). After implementing the current breakpoints, these additional tests may not need to be performed other than for epidemiological or infection **prevention** purposes (ie, it is no longer necessary to edit results for the carbapenems to resistant if a carbapenemase producer is detected).

Introduction to Tables 3B and 3C. (Continued)

	Tests	Used for Epidemiological or Infe	ection Prevention-Related	Testing
	CarbaNP	mCIM	mCIM With eCIM	
	(Table 3B)	(Table 3C)	(Table 3C)	Other (eg, molecular assays)
Organisms	Enterobacterales and P. aeruginosa that are not susceptible to one or more carbapenems	Enterobacterales and P. aeruginosa that are not susceptible to one or more carbapenems	Enterobacterales that are positive by mCIM	Enterobacterales and P. aeruginosa that are not susceptible to one or more carbapenems to determine the presence of a carbapenemase, or to determine carbapenemase type in isolates positive by CarbaNP or mCIM.
Strengths	Rapid	No special reagents or media necessary	No special reagents or media necessary	Determines type of carbapenemase in addition to absence or presence of the enzyme
Limitations	Special reagents are needed, some of which necessitate inhouse preparation (and have a short shelf life). Invalid results occur with some isolates. Certain carbapenemase types (eg, OXA-type, chromosomally encoded) are not consistently detected.	Requires overnight incubation	Requires overnight incubation	Special reagents and equipment are needed. Specific to targeted genes; false-negative result if specific carbapenemase gene present is not targeted.

Abbreviations: eCIM, EDTA-modified carbapenem inactivation method; mCIM, modified carbapenem inactivation method, MIC, minimal inhibitory concentration.

NOTE: Information in boldface type is new or modified since the previous edition.

Tables 3B and 3B-1 CarbaNP Test for Suspected Carbapenemase Production and Modifications When Using MIC Breakpoints Described in M100-S20 (January 2010)

Table 3B. CarbaNP Test for Suspected Carbapenemase Production in Enterobacterales and Pseudomonas aeruginosa¹⁻⁷

NOTE: If using FORMER MIC breakpoints for carbapenems described in M100-S20 (January 2010), please refer to modifications in Table 3B-1 below.

Test	CarbaNP Test	
When to perform this test	For epidemiological or infection prevention purposes. NOTE: No change in the interpretation of carbapenem susceptibility test results is necessary for CarbaNP–positive isolates. Such testing is not currently recommended for routine use.	
Test method	Colorimetric microtube assay	
Test reagents and	Clinical laboratory reagent water	
materials	Imipenem reference standard powder	
	Commercially available bacterial protein extraction reagent in Tris HCl buffer, pH 7.4	
	Zinc sulfate heptahydrate	
	Phenol red powder	
	1 N NaOH solution	
	10% HCl solution	
	Microcentrifuge tubes 1.5 mL, clear	
	• 1-µL inoculation loops	
	Containers to store prepared solutions	
	Use reagents above to prepare the following solutions (instructions for preparation are provided below this table):	
	10 mM zinc sulfate heptahydrate solution	
	0.5% phenol red solution	
	0.1 N sodium hydroxide solution	
	CarbaNP Solution A	
	CarbaNP Solution B (solution A + imipenem)	
Test procedure	1. Label two microcentrifuge tubes (one "a" and one "b") for each patient isolate, QC organism, and uninoculated reagent control.	
	2. Add 100 μL of bacterial protein extraction reagent to each tube.	
	3. For each isolate to be tested, emulsify a 1-µL loopful of bacteria from an overnight blood agar plate in both tubes "a" and "b."	
	Vortex each tube for 5 seconds. (Uninoculated reagent control tubes should contain only bacterial protein extraction	
	reagent, no organism.) NOTE : Do not use growth from selective media or plates containing antibiotics or other agents that	
	select for certain bacteria.	
	4. Add 100 μL of solution A to tube "a." 5. Add 100 μL of solution B to tube "b."	
	6. Vortex tubes well.	
	7. Incubate at 35°C ± 2°C for up to 2 hours. Isolates that demonstrate positive results before 2 hours can be reported as	
	carbapenemase producers.	

Table 3B. (Continued)

Test			CarbaNP Test	
Test interpretation	1. Read uning Both to If either 2. Read inocu Tube "	eading (see Figure 1, below): oculated reagent control tubes "a" a ubes must be red or red-orange. er tube is any other color, the test is ulated tube "a." a" must be red or red-orange. "a" is any other color, the test is inv	nd "b" (ie, "blanks"). invalid.	
	Red or Light or Orang	3. Read inoculated tube "b." Red or red-orange = negative		
		Re	esults for Patient and QC Tubes	
		Tube "a": Solution A	results for Patient and QC Tubes Tube "b": Solution B	Interpretation
		Tube "a":	Tube "b":	Interpretation Negative, no carbapenemase detected
		Tube "a": Solution A (serves as internal control)	Tube "b": Solution B	
		Tube "a": Solution A (serves as internal control) Red or red-orange	Tube "b": Solution B Red or red-orange Light orange, dark yellow, or	Negative, no carbapenemase detected Positive, carbapenemase

CarbaNP Test for Suspected Carbapenemase Production and Modifications When Using MIC
Breakpoints Described in M100-S20 (January 2010)

Table 3B. (Continued)

Test	CarbaNP Test	
Test interpretation (Continued)	NOTES:	
(common)	A slight color change may be observed with the addition of imipenem to solution A. Compare patient tubes to the uninoculated reagent control tubes when interpreting questionable results.	
	For invalid results:	
	Check reagents for QC strains and uninoculated reagent controls.	
	Reagent deterioration can cause invalid results. An invalid result for an uninoculated reagent control test indicates a problem with solution A and/or solution B. Check the pH of solution A. If pH is < 7.8, prepare fresh solution A and solution B.	
	Repeat the test, including the uninoculated reagent controls.	
	If the repeat test is invalid, perform molecular assay.	
Reporting	Report positive as "Carbapenemase producer."	
	Report negative as "No carbapenemase detected."	
QC recommendations	Test positive and negative QC strains and uninoculated reagent control tubes each day of testing.	
	K. pneumoniae ATCC®a BAA-1705™—Carbapenemase positive	
	K. pneumoniae ATCC [®] BAA-1706™—Carbapenemase negative	
	Results for uninoculated reagent control tubes "a" and "b" must be negative (ie, red or red-orange). Any other result invalidates all tests performed on that day with the same lot of reagents.	
	The addition of imipenem to tube "b" might cause tube "b" to appear red-orange when tube "a" is red.	

Abbreviations: ATCC®, American Type Culture Collection; MIC, minimal inhibitory concentration; pH, negative logarithm of hydrogen ion concentration; QC, quality control.

Footnote

a. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.

Table 3B. (Continued)

NOTE 1: Test recommendations were largely derived following testing of US isolates of **Enterobacterales** and *P. aeruginosa* and provide for a high level of sensitivity (>90%) and specificity (>90%) in detecting KPC, **NDM**, VIM, IMP, SPM, and SME-type carbapenemases in these isolates. The sensitivity and specificity of the test for detecting other carbapenemase production can vary. **The ability of this test, as listed in the above procedure, to detect OXA-48-like producers is poor.^{6,7}**

NOTE 2: In CLSI studies, two KPC-positive strains with low carbapenem MICs (one *E. cloacae* susceptible by MIC to all three carbapenems and one *E. coli* that was susceptible to meropenem and intermediate to imipenem and ertapenem) were not detected by this test.

NOTE 3: Additional investigations of CarbaNP with *Acinetobacter* spp. showed poor sensitivity (ie, 21.3% for *A. baumannii*); therefore, the previous recommendation for use of CarbaNP with *Acinetobacter* spp. was removed.

NOTE 4: Information in boldface type is new or modified since the previous edition.

CarbaNP Test for Suspected Carbapenemase Production and Modifications When Using MIC Breakpoints Described in M100-S20 (January 2010)

Table 3B-1. Modifications of Table 3B When Using MIC Breakpoints for Carbapenems Described in M100-S20 (January 2010)¹⁻⁵

Table ob 1. Modifications of Table ob When Coming who breakpoints for Garbapenettic Described in Milos CEC (Garbary 2010)		
Test	CarbaNP Test	
When to perform this test:	Until laboratories can implement the revised carbapenem MIC breakpoints, this test (or an alternative confirmatory test for carbapenemases) should be performed when isolates of Enterobacterales are suspicious for carbapenemase production based on imipenem or meropenem MICs of 2–4 µg/mL or ertapenem MIC of 2 µg/mL.	
Reporting	For isolates that are CarbaNP positive, report all carbapenems as resistant, regardless of MIC. If the CarbaNP test is negative, interpret the carbapenem MICs using CLSI breakpoints as listed in Table 2A in M100-S20 (January 2010). If the CarbaNP test is negative, interpret the carbapenem MICs using CLSI breakpoints as listed in Table 2A in M100-S20 (January 2010). NOTE: Not all carbapenemase-producing isolates of Enterobacterales are CarbaNP positive.	

Abbreviation: MIC, minimal inhibitory concentration.

Tables 3B and 3B-1 – Instructions for Preparing Test Components

The steps for preparing 10 mM zinc sulfate heptahydrate solution are listed below.

Step	Action	Comment
1	Weigh out 1.4 g of ZnSO ₄ • 7H2O.	
2	Add the powder to 500 mL clinical laboratory reagent water.	
3	Mix the solution.	
4	Store the solution at room temperature.	Expiration is 1 year or not to exceed expiration of individual
		components

The steps for preparing 0.5% phenol red solution are listed below.

Step	Action	Comment
1	Weigh out 1.25 g of phenol red powder.	
2	Add the powder to 250 mL clinical laboratory reagent water.	
3	Mix the solution.	
4	Store the solution at room temperature.	Expiration is 1 year or not to exceed expiration of individual components.
		NOTE: This solution does not remain in solution. Mix well before use.

The steps for preparing 0.1 N sodium hydroxide solution are listed below.

Step	Action	Comment
1	Add 20 mL of 1 N NaOH to 180 mL clinical laboratory reagent water.	
2	Store the solution at room temperature.	Expiration is 1 year or not to exceed expiration of individual
		components

Tables 3B and 3B-1. (Continued)

The steps for preparing CarbaNP solution A are listed below.

Step	Action	Comment
1	To a 25- to 50-mL beaker, add 2 mL of 0.5% phenol red solution to 16.6	
	mL clinical laboratory reagent water.	
2	Add 180 µL of 10 mM zinc sulfate solution.	
3	Adjust the pH to 7.8 ± 0.1 with 0.1 N NaOH solution (or 10% HCl	10% HCl solution can be used if the pH is too high.
	solution if pH is too high).	
4	Store the solution at 4 to 8°C in a small vial or bottle.	Protect the solution from prolonged light exposure.
		Expiration is 2 weeks or not to exceed expiration of individual components
		(solution should remain red or red-orange; do not use if solution turns any
		other color).

The steps for preparing CarbaNP solution B (solution A+6 mg/mL imipenem) are listed below.

Step	Action	Comment
1	Determine the amount of solution B needed, allowing 100 µL per tube for each patient, QC strain, and uninoculated reagent control.	Example: To test 2 patient isolates, positive and negative controls and an uninoculated reagent control, 500 µL of solution B is needed.
2	Weigh out approximately 10–20 mg of imipenem powder.	It is advisable to weigh out at least 10 mg of powder. Divide the actual weight by 6 to determine the amount (in mL) of solution A to add to the powder. Example: 18 mg of imipenem / 6 = 3 mL of solution A, which is sufficient
		for 30 tubes.
3	Store the solution at 4 to 8°C for up to 3 days.	

NOTE: Information in boldface type is new or modified since the previous edition.

Tables 3B and 3B-1 CarbaNP Test for Suspected Carbapenemase Production and Modifications When Using MIC Breakpoints Described in M100-S20 (January 2010)

Tables 3B and 3B-1. (Continued)

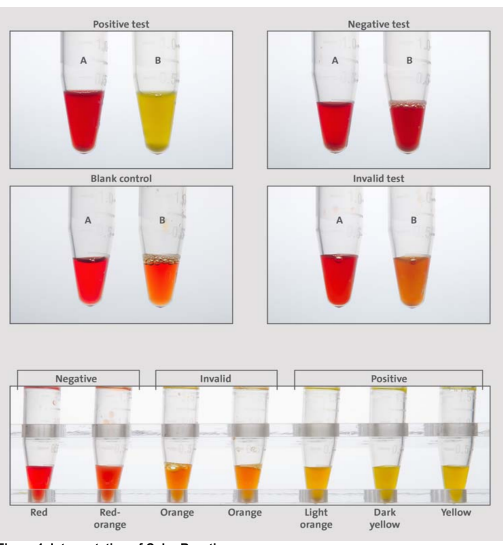


Figure 1. Interpretation of Color Reactions

Tables 3B and 3B-1. (Continued)

References for Tables 3B and 3B-1

- Nordmann P, Poirel L, Dortet L. Rapid detection of carbapenemase-producing *Enterobacteriaceae. Emerg Infect Dis.* 2012;18(9):1503-1507.
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- ⁷ Cunningham SA, Limbago B, Traczewski M, et al. Multicenter performance assessment of Carba NP test. J Clin Microbiol. 2017;55(6):1954-1960.

M100, 30th ed.

Table 3C. Modified Carbapenem Inactivation Methods for Suspected Carbapenemase Production in Enterobacterales and *Pseudomonas aeruginosa*¹⁻⁶

NOTE: If using FORMER MIC breakpoints for carbapenems described in M100-S20 (January 2010), please refer to modifications in Table 3C-1 below.

Test	mCIM Only or in Conjunction With eCIM
When to perform this test:	For epidemiological or infection prevention purposes.
	NOTE: No change in the interpretation of carbapenem susceptibility test results is necessary for mCIM positive and/or eCIM results. mCIM with or without eCIM testing is not currently recommended for routine use.
	mCIM is used for detecting carbapenemases in Enterobacterales and <i>P. aeruginosa</i> whereas eCIM is used together with mCIM to differentiate metallo-β-lactamases from serine carbapenemases in Enterobacterales .
	mCIM can be performed alone; however, eCIM must be performed together with mCIM.
	eCIM is only valid if mCIM is positive.
Test method	Meropenem disk inactivation
Test reagents and	TSB (2 mL aliquots)
materials	Meropenem disks (10 μg)
	• 1-μL and 10-μL inoculation loops
	Nutrient broth (eg, Mueller-Hinton, TSB) or normal saline (3.0–5.0 mL aliquots)
	MHA plates (100 mm or 150 mm)
	Meropenem-susceptible indicator strain – E. coli (ATCC®a 25922)
	0.5 M EDTA (only for eCIM)

Table 3C. (Continued)

Table 3C. (Continued) Test	mCIM Only or in Conjunction With aCIM
Test procedure: mCIM	mCIM Only or in Conjunction With eCIM 1. For each isolate to be tested, emulsify a 1-μL loopful of bacteria for Enterobacterales or 10-μL loopful of bacteria for <i>P. aeruginosa</i> from an overnight blood agar plate in 2 mL TSB.
	2. Vortex for 10–15 seconds.
	3. Add a 10-µg meropenem disk to each tube using sterile forceps or a single disk dispenser. Ensure the entire disk is immersed in the suspension.
	4. Incubate at 35°C±2°C in ambient air for 4 hours±15 minutes.
	5. Just before or immediately following completion of the TSB-meropenem disk suspension incubation, prepare a 0.5 McFarland suspension (using the colony suspension method) of <i>E. coli</i> ATCC® 25922 in nutrient broth or saline.
	6. Inoculate an MHA plate with <i>E. coli</i> ATCC [®] 25922 as for the routine disk diffusion procedure (see M02 ⁴) making sure the inoculum suspension preparation and MHA plate inoculation steps are each completed within 15 minutes. Allow the plates to dry for 3–10 minutes before adding the meropenem disks.
	7. Remove the meropenem disk from each TSB-meropenem disk suspension using a 10-µL loop by placing the flat side of the loop against the flat edge of the disk and using surface tension to pull the disk out of the liquid. Carefully drag and press the loop along the inside edge of the tube to expel excess liquid from the disk. Continue using the loop to remove the disk from the tube and then place it on the MHA plate previously inoculated with the meropenem-susceptible <i>E. coli</i> ATCC® 25922 indicator strain. Disk capacity: 4 disks on a 100 mm MHA plate; 8 disks on a 150 mm MHA plate (see Figure 1).
	8. Invert and incubate the MHA plates at 35°C±2°C in ambient air for 18–24 hours.
	9. Following incubation, measure the zones of inhibition as for the routine disk diffusion method (see M02 ⁴).
Test procedure: eCIM for Enterobacterales only;	For each isolate, label a second 2-mL TSB tube for the eCIM test.
optional	2. Add 20 μL of the 0.5 M EDTA to the 2-mL TSB tube to obtain a final concentration of 5 mM EDTA.
	3. Follow steps 1 through 9 above as for mCIM procedure. Process the mCIM and eCIM tubes in parallel.
	4. Place the meropenem disks from the mCIM and eCIM tubes on the same MHA plate inoculated with the meropenem-susceptible <i>E. coli</i> ATCC [®] 25922 indicator strain.
	NOTE: Additional QC is needed for the eCIM test (see QC recommendations).

M100, 30th ed.

Table 3C. (Continued)

Test	Test mCIM Only or in Conjunction With eCIM				
Test interpretation	For additional explanations, refer to Figures 2A, 2B, and 3A through 3D, as well as the notes section below.				
	mCIM Carbapenemase positive (see Figures 2A and 2B): Zone diameter of 6–15 mm or presence of pinpoint colonies within a 16–18 mm zone				
	 If the test isolate produces a carbapenemase, the meropenem in the disk will be hydrolyzed and there will be no inhibition or limited growth inhibition of the meropenem-susceptible <i>E. coli</i> ATCC[®] 25922. 				
	 Carbapenemase negative (see Figure 2A): Zone diameter of ≥ 19 mm (clear zone) 				
	 If the test isolate does not produce carbapenemase, the meropenem in the disk will not be hydrolyzed and will inhibit growth of the meropenem-susceptible E. coli ATCC® 25922. 				
	 Carbapenemase indeterminate: Zone diameter of 16–18 mm Zone diameter of ≥ 19 mm and the presence of pinpoint colonies within the zone The presence or absence of a carbapenemase cannot be confirmed. 				
	eCIM – Interpret only when mCIM test is positive • Metallo-β-lactamase positive: - A ≥ 5-mm increase in zone diameter for eCIM vs zone diameter for mCIM (eg, mCIM = 6 mm; eCIM = 15 mm; zone diameter difference = 9 mm). For only the eCIM test, ignore pinpoint colonies within any zone of inhibition (see Figures 3B and 3C).				
	 If the test isolate produces a metallo-β-lactamase, the activity of the carbapenemase will be inhibited in the presence of EDTA such that the meropenem in the disk will not be hydrolyzed as efficiently as in the tube without EDTA. The result is inhibition of the meropenem-susceptible <i>E. coli</i> and an increase in the zone diameter for the eCIM zone diameter compared with the mCIM zone diameter. 				
	 Metallo-β-lactamase negative: A ≤4-mm increase in zone diameter for the eCIM vs zone diameter of mCIM (eg, mCIM = 6 mm; eCIM = 8 mm; zone diameter difference = 2 mm). For only the eCIM test, ignore pinpoint colonies within any zone of inhibition (see Figure 3D). 				
	 If the test isolate produces a serine carbapenemase, the activity of the carbapenemase will not be affected by the presence of EDTA and there will be no or marginal (≤4 mm) increase in zone diameter in the presence of EDTA compared with the mCIM zone diameter. 				

Table 3C. (Continued)

Test	mCIM Only or in Conjunction With eCIM				
Reporting	mCIM Only				
	mCIM Result	eCIM Result	Report		
	Negative	Not set up	Carbapenemase not detected		
	Positive	Not set up	Carbapenemase detected		
	Indeterminate	Not set up	Testing inconclusive for the presence of carbapenemase.		
			Call laboratory to discuss.*		
		mCIM ar	nd eCIM Combination Test		
	mCIM Result	eCIM Result	Report		
	Negative	Do not interpret	Carbapenemase not detected		
	Positive	Negative	Serine carbapenemase detected		
	Positive	Positive	Metallo-β-lactamase detected		
	Indeterminate	Do not interpret	Testing inconclusive for the presence of carbapenemase. Call laboratory to discuss.*		
	* If indeterminate results are obtained on repeat testing, consider performing a different phenotypic test for carbapenemase detection (ie, CarbaNP), a test for carbapenemase genes or send isolate to a referral laboratory for further testing.				
	If both a serine carbapenemase and a metallo-β-lactamase are co-produced by one organism, differentiation between enzymes will not be possible and false-negative eCIM results may occur.				

Table 3C. (Continued)

Test					
NOTES					
QC recommendations		Organism Characteristic	and 2B for examples of positive and negative Expected Result		
	K. pneumoniae	KPC positive	mCIM positive		
	ATCC® BAA-1705™	Serine carbapenemase producer	eCIM negative		
	K. pneumoniae ATCC® BAA-1706™	Carbapenemase negative	mCIM negative		
	 K. pneumoniae ATCC® BAA-2146™* NDM positive Metallo-β-lactamase producer mCIM positive eCIM positive 				
Abbreviations: ATCC® Amer	In addition, perform QC of meroper and handle disks as described in M0 cartridge of disks used for the run an	2.4 Alternatively, perform QC of meropenem of placing it on the MHA plate inoculated with a	owing the routine disk diffusion QC procedure, disks with each run by removing a disk from the E. coli ATCC [®] 25922; incubate as above. thod; mCIM, modified carbapenem inactivatio		

Abbreviations: ATCC®, American Type Culture Collection; eCIM, EDTA-modified carbapenem inactivation method; mCIM, modified carbapenem inactivation method; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; ; QC, quality control; TSB, trypticase soy broth.

Table 3C. (Continued)

Footnotes

- a. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- b. The AR Isolate Bank (http://www.cdc.gov/drugresistance/resistance-bank/overview.html) is a centralized repository of microbial pathogens with well-characterized resistance profiles that are assembled by the Centers for Disease Control and Prevention in collaboration with the US Food and Drug Administration.

NOTE 1: mCIM: This method demonstrated a sensitivity > 99% and specificity > 99% for detection of KPC, NDM, VIM, IMP, IMI, SPM, SME and OXA-type carbapenemases among **Enterobacterales** isolates investigated by CLSI.^b This method demonstrated a sensitivity > 97% and specificity 100% for detection of KPC, NDM, VIM, IMP, IMI, SPM and OXA-type carbapenemases among *P. aeruginosa* isolates investigated by CLSI.^b Performance for other carbapenemases or for testing isolates of non-**Enterobacterales** other than *P. aeruginosa* has not been established. Investigations of mCIM with *Acinetobacter* spp. showed poor specificity and poor reproducibility between laboratories, and performing mCIM with *Acinetobacter* spp. is not endorsed by CLSI. In CLSI studies, one OXA-232–producing *K. pneumoniae* isolate was negative by this assay at 4 of 9 validation sites.

NOTE 2: eCIM: This method demonstrated a sensitivity > 95% and specificity > 92% for differentiation of metallo-β-lactamases (NDM, VIM, and IMP) from serine carbapenemases (KPC, OXA, and SME) among **Enterobacterales** isolates investigated by CLSI.^b In CLSI studies, one *K. pneumoniae* co-producing NDM and OXA-181 yielded a false-negative result at 3 of 4 validation sites.

NOTE 3: Information in boldface type is new or modified since the previous edition.

Table 3C. (Continued)







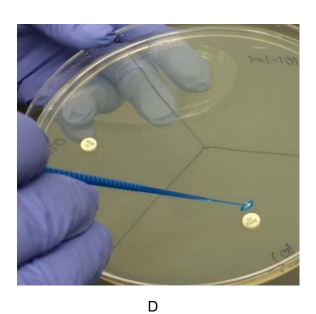


Figure 1. Procedure for Placing Meropenem Disks for the mCIM. Remove the meropenem disk with a 10-μL loop (A) and drag the loop against the inside edge of the tube to expel any excess liquid (B). Use the same loop to remove the disk from the tube (C) and place it on the MHA plate (D) previously inoculated with the meropenem-susceptible *E. coli* (ATCC® 25922) indicator strain.

Table 3C. (Continued)



Figure 2A. mCIM Results for QC Strains: Negative Control K. pneumoniae ATCC® BAA-1706™ (A) and Positive Control K. pneumoniae ATCC® BAA-1705™ (B). NOTE: A narrow ring of growth around the meropenem disk as seen with the negative control (A) results from carryover of the test organism in the TSB and should be ignored.

Table 3C. (Continued)



Figure 2B. mCIM Test Interpretation

- Result: positive mCIM
- Report: carbapenemase detected

NOTE: A narrow ring of growth around the meropenem disk results from carryover of the test organism in the TSB and should be ignored.

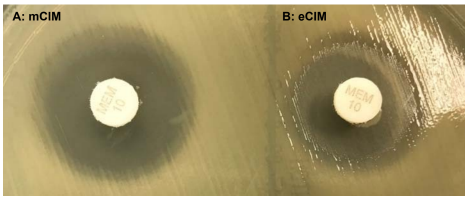


Figure 3A. mCIM and eCIM Test Interpretation: Negative mCIM. "A" shows an mCIM negative result (zone diameter = 20 mm) and "B" shows an eCIM invalid result. Do not interpret the eCIM result when the mCIM is negative as the isolate is negative for carbapenemase production.

- Result: negative for carbapenemase production
- Report: carbapenemase not detected

Table 3C. (Continued)

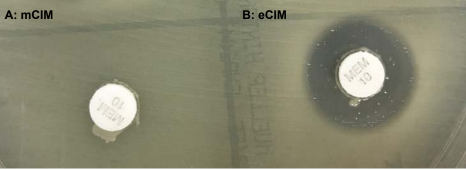


Figure 3B. mCIM and eCIM Test Interpretation: Positive mCIM and eCIM. "A" shows an mCIM positive result (zone diameter of 6 mm) and "B" shows an eCIM positive result (zone diameter = 15 mm with pinpoint colonies throughout the zone of inhibition). NOTE: The pinpoint colonies throughout the zone of inhibition are ignored when measuring the zone for the eCIM test. $A \ge 5$ -mm increase in zone diameter for eCIM vs zone diameter for mCIM (15 mm - 6 mm = 9 mm) demonstrates the inhibition of the metallo-β-lactamase in the presence of EDTA.

- · Result: positive mCIM and eCIM
- Report: metallo-β-lactamase detected

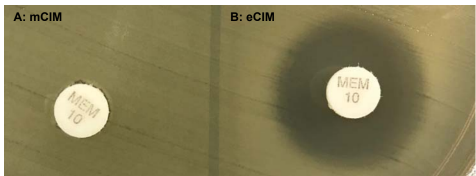


Figure 3C. mCIM and eCIM Test Interpretation: Positive mCIM and eCIM. "A" shows an mCIM positive result (zone diameter = 6 mm) and "B" shows an eCIM positive result (zone diameter = 19 mm). A \geq 5-mm increase in zone diameter for eCIM vs diameter for mCIM zone (19 mm – 6 mm = 13 mm) demonstrates the inhibition of the metallo-β-lactamase in the presence of EDTA.

- Result: positive mCIM and eCIM
- Report: metallo-β-lactamase detected

Modified Carbapenem Inactivation Methods and Modifications When Using MIC Breakpoints
Described in M100-S20 (January 2010)

Table 3C. (Continued)

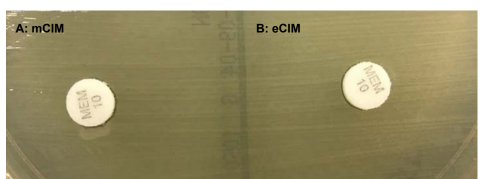


Figure 3D. mCIM and eCIM Test Interpretation: Positive mCIM and Negative eCIM. "A" shows an mCIM positive result (zone diameter = 6 mm) and "B" shows an eCIM negative result (zone diameter = 6 mm). Serine carbapenemases are not inhibited by EDTA and demonstrate a ≤ 4-mm increase in zone diameter for eCIM vs zone diameter for mCIM.

- Result: positive mCIM and negative eCIM
- Report: serine carbapenemase detected

References for Table 3C

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- Sfeir MM, Hayden JA, Fauntleroy KA, et al. EDTA-modified carbapenem inactivation method: a phenotypic method for detecting metallo-β-lactamase-producing Enterobacteriaceae. J Clin Microbiol. 2019;57(5):pii: e01757-18.

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Tables 3C and 3C-1
Modified Carbapenem Inactivation Methods and Modifications When Using MIC Breakpoints
Described in M100-S20 (January 2010)

Table 3C-1 Modifications of Table 3C When Using MIC Breakpoints for Carbanenems Described in M100-S20 (January 2010)

Test	mCIM				
When to perform this test:	Until laboratories can implement the revised carbapenem MIC breakpoints, this test (or an alternative confirmatory test for carbapenemases) should be performed when isolates of Enterobacterales are suspicious for carbapenemase production based on imipenem or meropenem MICs of $2-4 \mu g/mL$ or ertapenem MIC of $2 \mu g/mL$.				
Reporting	For isolates that are mCIM positive, report all carbapenems as resistant, regardless of MIC. If the mCIM test is negative, interpret the carbapenem MICs using CLSI breakpoints as listed in Table 2A in M100-S20 (January 2010).				
	If the mCIM test is negative, interpret the carbapenem MICs using CLSI breakpoints as listed in Table 2A in M100-S20 (January 2010).				
	NOTE: Not all carbapenemase-producing isolates of Enterobacterales are mCIM positive.				

Abbreviations: mCIM, modified carbapenem inactivation method; MIC, minimal inhibitory concentration.

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Table 3D. Tests for Colistin Resistance for Enterobacterales and Pseudomonas aeruginosa

The polymyxins (colistin and polymyxin B) are antimicrobial agents of last resort for treating multidrug-resistant infections. Clinical and PK-PD data suggest that these agents have limited clinical efficacy. Alternative agents are strongly preferred. If these agents are not available, knowledge of the colistin MIC may be helpful to inform treatment decisions.

For colistin, broth microdilution, broth disk elution and agar dilution MIC methods are acceptable. Broth microdilution is the only approved method for polymyxin B. Disk diffusion and gradient diffusion methods should not be performed.

Colistin and polymyxin B are considered equivalent agents, so MICs obtained from testing colistin predict MICs to polymyxin B and vice versa. At this time, CLSI has not evaluated polymyxin B testing methods, and the procedures below should not be adapted to polymyxin B. The methods below were evaluated for *Acinetobacter* spp. by CLSI and found to yield inaccurate results.

These methods were established with limited disk and/or media manufacturers and are considered provisional until additional data are evaluated by CLSI and shown to meet CLSI document M23¹ guidelines.

Test	Colistin Broth Disk Elution	Colistin Agar Test
Approved organisms	Enterobacterales and Pseudomonas aeruginosa	Enterobacterales and P. aeruginosa
Strengths	No special reagents or media necessary	Ability to test up to 10 isolates at one time
Limitations	Hands-on time and cost	Requires special media (colistin agar plate)
When to perform this test	Testing multidrug-resistant isolates for clinical or infection prevention purposes	Testing multidrug-resistant isolates for clinical or infection prevention purposes
Test method	Tube dilution using colistin disk as the colistin source	Agar dilution: slight variation of method described in M07 ² (ie, different inoculum and different approach to interpreting results)
Organism group	Enterobacterales and P. aeruginosa	Enterobacterales and P. aeruginosa
Medium	CAMHB (10-mL tubes)	MHA (20 mL in 100-mm Petri plate) ^a
Antimicrobial concentration	10-μg colistin disks Final concentration: 0 μg/mL (growth control), 1 μg/mL, 2 μg/mL, and 4 μg/mL colistin	Final concentration: 0 μg/mL (growth control), 1 μg/mL, 2 μg/mL, and 4 μg/mL colistin ^a
Inoculum	 Using a loop or swab, pick 3–5 colonies from a fresh (18–24 hours) nonselective agar plate and transfer to sterile saline (4–5 mL). Adjust turbidity to equivalent of a 0.5 McFarland turbidity 	 Using a loop or swab, pick 3–5 colonies from a fresh (18–24 hours) nonselective agar plate and transfer to sterile saline (4–5 mL). Adjust turbidity to equivalent of a 0.5 McFarland turbidity standard.
	standard.	3. Dilute the standardized inoculum 1:10 in saline.

Table 3D. (Continued)

Test	Colistin Broth Disk Elution	Colistin Agar Test
Test Test procedure	 Let the CAMHB tubes (10 mL) and colistin disks warm to room temperature. Label 4 tubes of CAMHB for each isolate to be tested with 1, 2, and 4 μg/mL and control (see Figure 1). Using aseptic technique, carefully add: 1 colistin disk to the tube labeled "1 μg/mL" 2 colistin disks to tube labeled "2 μg/mL" 4 colistin disks to the tube labeled "4 μg/mL" Gently vortex the tubes with the added disk and let the colistin elute from the disks for at least 30 minutes but no longer than 60 minutes at room temperature. Prepare the standardized inoculum. Add 50 μL standardized inoculum to the control 	 Colistin Agar Test Divide each colistin agar plate with increasingly doubled dilutions of colistin in up to 10 parts, with a marker to test up to 10 isolates per plate. Label each part with the appropriate isolate number (see Figure 2). Using a pipette or a 10-μL loop, streak 10 μL of the 1:10 dilution onto the appropriate part of each colistin agar plate. Using a 10-μL loop, subculture from the original inoculum tube to a blood agar plate as a purity check. Incubate the colistin agar plates and purity plate
	 6. Add 50 μL standardized inoculum to the control and 1-, 2-, and 4-μg/mL tubes to attain a final inoculum concentration of approximately 7.5 × 10⁵ CFU/mL. 7. Using a 10-μL loop, subculture from the original inoculum tube to a blood agar plate as a purity 	
	check. 8. Cap the tubes tightly and vortex each inoculated tube on slow speed to mix. Slow speed is suggested to prevent colistin from sticking to the cap and glass surface above the meniscus of liquid.	
	9. Loosen the caps slightly before incubation.10. Incubate the tubes and purity plate.	
Incubation conditions	33 to 35°C; ambient air	33 to 35°C; ambient air
Incubation length	16–20 hours	16–20 hours

Table 3D. (Continued)

Test	Colistin Broth Disk Elution	Colistin Agar Test
Results	Examine the purity plate to ensure inoculum was pure.	Examine the purity plate to ensure inoculum was pure.
	Examine the growth control tube, which must demonstrate obvious turbidity for the test to be valid. NOTE: Some <i>P. aeruginosa</i> isolates may grow only near	Examine the growth control plate, which must demonstrate confluent growth for the test to be valid.
	the meniscus.	3. Examine the colistin plates carefully with transmitted light for colony or light film of growth.
	Read the MIC as the lowest concentration that completely inhibits growth of the test isolate. (See Figure 1 for examples.) For Enterobacterales and <i>P. aeruginosa:</i>	4. Read the MIC as the lowest colistin agar plate concentration that completely inhibits growth of the test isolate (eg, even 1 colony would be considered growth). See Figure 2 for examples.
	≤ 2 µg/mL = intermediate ≥ 4 µg/mL = resistant	For Enterobacterales and <i>P. aeruginosa:</i>
	2 4 µg/mz = 103/3tum	≤2 μg/mL = intermediate ≥4 μg/mL = resistant
Additional testing and reporting	If there is an inconsistent growth pattern (eg, no growth in 2 μg/mL but growth at 1 μg/mL and 4 μg/mL), repeat the test. An inconsistent growth pattern may occur as a result of: Contamination at higher dilutions Heteroresistance Improper concentrations of antimicrobial agent in the	If there is an inconsistent growth pattern (eg, no growth in 2 μg/mL but growth at 1 μg/mL and 4 μg/mL), repeat the test. An inconsistent growth pattern may occur as a result of: Contamination at higher dilutions Heteroresistance Improper concentrations of antimicrobial agent in the
	tubesError inoculating the tubes	colistin agar plates • Error inoculating the plates
QC recommendations – routine ^b	Escherichia coli AR Bank #0349 mcr-1 (≤1–4 μg/mL, with a target of 2 μg/mL) ^c and <i>P. aeruginosa</i> ATCC ^{®d} 27853 (≤1–4 μg/mL)	E. coli AR Bank #0349 mcr-1 (≤1–4 μg/mL, with a target of 2 μg/mL) ^c and P. aeruginosa ATCC [®] 27853 (≤ 1–4 μg/mL)

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; CFU, colony-forming unit(s); MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; PK-PD, pharmacokinetic-pharmacodynamic; QC, quality control.

Table 3D. (Continued)

Footnotes

- a. Refer to M07² for preparation of media and antimicrobial agents.
- b. QC recommendations routine

Test recommended routine QC strains:

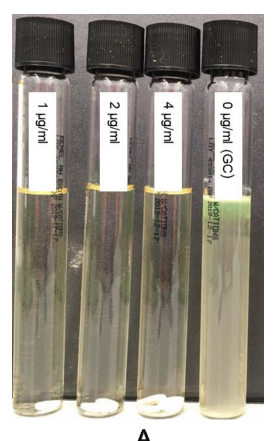
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02³ and M07²) and the individualized QC plan is complete
- . Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met

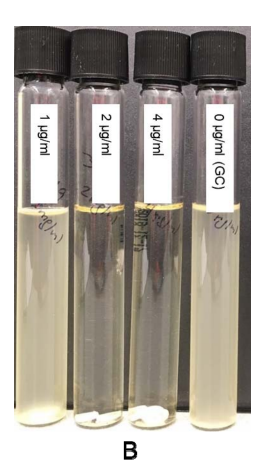
Perform QC of colistin disks and test media daily or weekly following the routine disk diffusion QC procedure and handle disks as described in M02.3

- c. The QC ranges were established with disks (colistin broth disk elution) and media from a limited number of manufacturers and are considered provisional until additional data are evaluated by CLSI and shown to meet CLSI document M23¹ guidelines.
- d. ATCC® is a registered trademark of the American Type Culture Collection.

NOTE: Information in boldface type is new or modified since the previous edition.

Table 3D. (Continued)

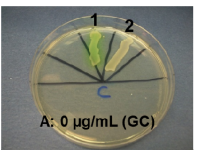


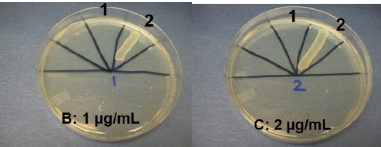


Abbreviation: GC, growth control.

Figure 1. Colistin Broth Disk Elution. Results for routine QC strain *P. aeruginosa* ATCC[®] 27853 with an MIC \leq 1 µg/mL (A) and supplemental QC strain *E. coli* AR Bank #0349 *mcr-1* with an MIC 2 µg/mL (B). (Courtesy of Patricia J. Simner, Johns Hopkins University School of Medicine. Used with permission.)

Table 3D. (Continued)





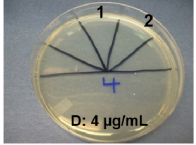


Figure 2. Colistin Agar Test. The plates need to be examined carefully with transmitted light for confluent growth, individual colonies, or light film of growth to determine the MIC. Colistin agar test results for routine QC strain *P. aeruginosa* ATCC[®] 27853 (position 1) with an MIC ≤ 1 μg/mL and for supplemental QC strain *E. coli* AR Bank #0349 *mcr-1* (position 2) with an MIC 4 μg/mL. The plates shown contain 0 μg/mL (control) (A), 1 μg/mL (B), 2 μg/mL (C), and 4 μg/mL (D) colistin. (Courtesy of Patricia J. Simner, Johns Hopkins University School of Medicine. Used with permission.)

References for Table 3D

- ¹ CLSI. Development of In Vitro Susceptibility Testing Criteria and Quality Control Parameters. 5th ed. CLSI guideline M23. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ³ CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 3E. Test for Detection of β-Lactamase Production in *Staphylococcus* spp.

Test	Test β-Lactamase Production				
Test method	Disk Diffusion (penicillin zone-edge test)	Nitrocefin-based Test			
Organism group	S. aureus with penicillin MICs ≤ 0.12 μg/mL or zones ≥ 29 mm ^a	Staphylococcus spp.a,b with penicillin MICs ≤ 0.12 μg/mL or zones ≥ 29 mm			
Medium	MHA	N/A			
Antimicrobial concentration	10 units penicillin disk	N/A			
Inoculum	Standard disk diffusion procedure	Induced growth (ie, growth taken from the zone margin surrounding a penicillin or cefoxitin disk test on either MHA or a blood agar plate after 16–18 hours of incubation)			
Incubation conditions	35°C±2°C; ambient air	Room temperature			
Incubation length	16–18 hours	Up to 1 hour for nitrocefin-based test or follow manufacturer's directions			
Results	Sharp zone edge ("cliff") = β-lactamase positive (see Figure 1 below this table) Fuzzy zone edge ("beach") = β-lactamase negative (see Figure 2 below this table)	Nitrocefin-based test: conversion from yellow to red/pink = β-lactamase positive.			
Additional testing and reporting	β-lactamase-positive staphylococci are resistant to penicillin, amino-, carboxy-, and ureidopenicillins.	Nitrocefin-based tests can be used for <i>S. aureus</i> , but negative results should be confirmed with the penicillin zone-edge test before reporting penicillin as susceptible. β-lactamase-positive staphylococci are resistant to penicillin, amino-, carboxy-, and ureidopenicillins.			
QC recommendations – routine ^c	S. aureus ATCC®d 25923 for routine QC of penicillin disk to include examination of zone-edge test (fuzzy edge = "beach")	and diedopeniciims.			
QC recommendations – lot/shipment ^e		S. aureus ATCC® 29213 – positive S. aureus ATCC® 25923 – negative			
QC recommendations – supplemental ^f	S. aureus ATCC® 29213 – positive penicillin zone- edge test (sharp edge = "cliff")	(or see local regulations and manufacturers' recommendations)			

Abbreviations: ATCC®, American Type Culture Collection; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; N/A, not applicable; QC, quality control.

Table 3E. (Continued)

Footnotes

- a. The penicillin disk diffusion zone-edge test was shown to be more sensitive than nitrocefin-based tests for detection of β-lactamase production in *S. aureus*. The penicillin zone-edge test is recommended if only one test is used for β-lactamase detection. However, some laboratories may choose to perform a nitrocefin-based test first and, if this test is positive, report the results as positive for β-lactamase (or penicillin resistant). If the nitrocefin test is negative, the penicillin zone-edge test should be performed before reporting the isolate as penicillin susceptible in cases in which penicillin may be used for therapy (eg, endocarditis).^{1,2}
- b. For *S. lugdunensis*, tests for β-lactamase detection are not necessary because isolates producing a β-lactamase will test penicillin resistant (MIC > 0.12 µg/mL and zone diameters < 29 mm). If a laboratory is using a method other than the CLSI disk diffusion or MIC reference methods and is unsure if the method can reliably detect penicillin resistance with contemporary isolates of *S. lugdunensis*, the laboratory should perform an induced nitrocefin assay or other CLSI reference method on isolates that test penicillin susceptible before reporting the isolate as penicillin susceptible.
- c. QC recommendations routine

Test negative (susceptible) QC strain:

- With each new lot/shipment of testing materials
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02³ and M07⁴)
- Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met
- d. ATCC® is a registered trademark of the American Type Culture Collection.
- e. QC recommendations lot/shipment

Test positive (resistant) QC strain at minimum with each new lot/shipment of testing materials.

- f. QC recommendations supplemental
 - Supplemental QC strains can be used to assess a new test, for training personnel, and for competence assessment. It is not necessary to include supplemental QC strains in routine daily or weekly antimicrobial susceptibility testing QC programs. See Appendix C, which describes use of QC strains.

Table 3E. (Continued)



Figure 1. Positive Penicillin Disk Zone-Edge Test for β-Lactamase Detection. The zone edge is sharp or like a "cliff" indicating β-lactamase production.



Figure 2. Negative Penicillin Disk Zone-Edge Test for β-Lactamase Detection. The zone edge is fuzzy or like a "beach," indicating no β-lactamase production.

Table 3E. (Continued)

References for Table 3E

- Kaase M, Lenga S, Friedrich S, et al. Comparison of phenotypic methods for penicillinase detection in *Staphylococcus aureus*. *Clin Microbiol Infect*. 2008;14(6):614-616.
- Gill VJ, Manning CB, Ingalls CM. Correlation of penicillin minimum inhibitory concentrations and penicillin zone edge appearance with staphylococcal beta-lactamase production. *J Clin Microbiol.* 1981;14(4):437-440.
- 3 CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ⁴ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

mecA negative M100, 30th ed

Table 3F. Test for Detecting Methicillin (Oxacillin) Resistance in Staphylococcus spp. Detecting mecA-mediated Detecting mecA-Mediated Resistance Detecting mecA-Mediated Resistance Resistance Using Oxacillin Salt Agar Test **Using Cefoxitin Using Oxacillin** Test method Disk Diffusion **Disk Diffusion Broth Microdilution Broth Microdilution and Agar Dilution Agar Dilution** Other Staphylococcus Staphylococcus Organism S. aureus and S. aureus and S. epidermidis, S. S. aureus and S. aureus pseudintermedius. group spp. (excluding S. lugdunensis S. lugdunensis spp. (excluding luadunensis S. pseudintermedius and S. schleiferi S. aureus and and S. schleiferi) S. lugdunensis) MHA CAMHB MHA CAMHB with 2% NaCl (broth MHA with 4% NaCl Medium microdilution) MHA with 2% NaCl (agar dilution) Antimicrobial 0.25 µg/mL oxacillin 30 µg cefoxitin disk 4 µg/mL cefoxitin 1-µg oxacillin disk 2 µg/mL 6 μg/mL oxacillin oxacillin concentration Inoculum Standard disk diffusion Standard broth Standard disk Standard broth microdilution procedure Colony suspension to obtain diffusion 0.5 McFarland turbidity procedure microdilution or standard agar dilution procedure Procedure procedure Using a 1-µL loop that was dipped in the suspension, spot an area 10-15 mm in diameter. Alternatively, using a swab dipped in the suspension and the excess liquid expressed, spot a similar area or streak an entire quadrant. Incubation 33 to 35°C; ambient 33 to 35°C; ambient 33 to 35°C: ambient aira 33 to 35°C: ambient aira 33 to 35°C: ambient aira conditions air^a aira Incubation 16-18 hours 24 hours (may be 16-20 hours 16-18 hours 24 hours (may be reported after 24 hours; length reported after 18 18 hours, if resistant) read with transmitted light hours, if resistant) ≥ 0.5 µg/mL = *mecA* Results ≤ 21 mm = ≤ 24 mm = *mecA* ≥ **8** µg/mL =*mecA* ≤ 17 mm = mecA ≥4 µg/mL = Examine carefully with mecA positive positive positive positive mecA positive positive transmitted light for > 1 colony or light film of growth. ≥ 22 mm = ≥ 25 mm = mecA \leq 4 µg/mL = mecA≥ 18 mm = mecA ≤ 2 µg/mL = \leq 0.25 µg/mL = mecAnegative negative negative mecA negative negative > 1 colony = oxacillin resistant

Table 3F. (Continued)

Test	Detecting <i>mecA-</i> Mediated Resistance Using Cefoxitin		Detecting <i>mecA</i> -Mediated Resistance Using Oxacillin		Detecting <i>mecA</i> -mediated Resistance Using Oxacillin Salt Agar
Additional testing and reporting	Cefoxitin is used as a surrogate for mecA-mediated methicillin (oxacillin) resistance. Isolates that test as mecA positive should be reported as methicillin (oxacillin) (not cefoxitin) resistant; other β-lactam agents, except ceftaroline, should be reported as resistant or should not be reported.	Ed as a surrogate for ad methicillin (oxacillin) Cefoxitin is used as a surrogate for mecA-mediated methicillin (oxacillin) resistance. Est as mecA positive stream methicillin (oxacillin) resistant; other β-lactam agents, except ceftaroline, should be reported as resistant or should not be reported. Because of the rare occurrence of methicillin (oxacillin) resistance mechanisms other than mecA, isolates that test as mecA negative but for which the oxacillin MICs are resistant (MIC ≥ 4 μg/mL) should be reported as methicillin (oxacillin) resistant; routine Isolates that test as mecA positive should be reported as resistant or should not be reported. Because of the rare occurrence of methicillin (oxacillin) resistance mechanisms other than mecA, isolates that test as mecA negative but for which the oxacillin MICs are resistant (MIC ≥ 4 μg/mL) should be reported as methicillin (oxacillin) resistant.		MRS are resistant to all β-lactam agents with the exception of ceftaroline; other β-lactam agents should be reported as resistant or should not be reported	
		B-lactam agents, except ceftaroline , is not advised. Because of the rare occurrence of methicillin (oxacillin) resistance mechanisms other than <i>mecA</i> , isolates that test as <i>mecA</i> negative, but for which the oxacillin MICs are resistant (MIC ≥ 4 μg/mL), should be reported as methicillin (oxacillin) resistant.		spp., excluding S. aureus, S. lugdunensis, S. pseudintermedius, and S. schleiferi, oxacillin MIC breakpoints may overcall resistance, and some isolates for which the oxacillin MICs are 0.5–2 µg/mL may be mecA negative. Isolates from serious infections for which oxacillin MICs are 0.5–2 µg/mL may be tested for mecA or for PBP2a. Isolates that test mecA or PBP2a negative should be reported as methicillin (oxacillin) susceptible.	

Table 3F
Test for Methicillin (Oxacillin) Resistance
in Staphylococcus spp.

Table 3F. (Continued)

Test	Detecting <i>mecA</i> -Mediated Resistance Using Cefoxitin			Detecting <i>mecA</i> -Mediated Resistance Using Oxacillin			Detecting Oxacillin Resistance Using Oxacillin Salt Agar
Test method	Disk Diffusion Broth Microdi		Broth Microdilution	Disk Diffusion	Broth Microdilution and Agar Dilution		Agar Dilution
Organism group	S. aureus and S. lugdunensis	Other Staphylococcus spp., excluding S. pseudintermedius S. schleiferi	S. aureus and S. lugdunensis	S. epidermidis, S. pseudintermedius, and S. schleiferi	S. aureus and S. lugdunensis	Staphylococcus spp., excluding S. aureus and S. lugdunensis)	S. aureus
QC recommendations – routine ^b	S. aureus ATCC® 25923 – mecA negative (cefoxitin zone 23–29 mm)		S. aureus ATCC® 29213 – mecA negative (cefoxitin MIC 1–4 µg/mL)	S. aureus ATCC® 25923 - mecA negative (oxacillin zone 18-24 mm)	S. aureus ATCC® 2 negative (oxacillin µg/mL)		S. aureus ATCC®c 29213 – susceptible (with each test day)
QC recommendations – lot/shipment ^d			S. aureus ATCC® 43300 – mecA positive (MIC > 4 μg/mL)	S. aureus ATCC® 43300 - mecA positive (zone ≤ 24 mm)	S. aureus ATCC® 4 positive (MIC > 4 p	ıg/mL)	S. aureus ATCC® 43300 – resistant

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; MRS, methicillin (oxacillin)-resistant staphylococci; PBP2a, penicillin-binding protein 2a; QC, quality control.

Footnotes

- a. Testing at temperatures above 35°C may not detect MRS.
- b. QC recommendations routine

Test negative (susceptible) QC strain:

- With each new lot/shipment of testing materials
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02¹ and M07²)
- Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met
- c. ATCC® is a registered trademark of the American Type Culture Collection.
- d. QC recommendations lot/shipment

Test positive (resistant) QC strain at minimum with each new lot/shipment of testing materials.

NOTE: Information in boldface type is new or modified since the previous edition.

For Use With M02 and M07

Table 3F. (Continued)

References for Table 3F

- ¹ CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically.* 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 3G
Vancomycin Agar Screen for Staphylococcus aureus and Enterococcus spp.

Table 3G. Vancomycin Agar Screen for Staphylococcus aureus and Enterococcus spp.

Screen Test	Vancomycin MIC ≥8 μg/mL					
Test method	Agar Dilution	Agar Dilution				
Organism group	S. aureus	Enterococcus spp.				
Medium	BHI agar	BHI ^a agar				
Antimicrobial concentration	6 μg/mL vancomycin	6 μg/mL vancomycin				
Inoculum	Colony suspension to obtain 0.5 McFarland turbidity Preferably, using a micropipette, spot a 10-μL drop onto agar surface. Alternatively, using a swab dipped in the suspension and the excess liquid expressed, spot an area 10–15 mm in diameter or streak a portion of the plate.	1–10 µL of a 0.5 McFarland suspension spotted onto agar surface. Alternatively, using a swab dipped in the suspension and the excess liquid expressed, spot an area 10–15 mm in diameter or streak a portion of the plate.				
Incubation conditions	35°C±2°C; ambient air	35°C±2°C; ambient air				
Incubation length	24 hours	24 hours				
Results	Examine carefully with transmitted light for > 1 colony or light film of growth. > 1 colony = Presumptive reduced susceptibility to vancomycin	>1 colony = Presumptive vancomycin resistance				
Additional testing and reporting	Perform a vancomycin MIC using a validated MIC method to determine vancomycin MICs on <i>S. aureus</i> that grow on BHI–vancomycin screening agar. Testing on BHI–vancomycin screening agar does not reliably detect all vancomycin-intermediate <i>S. aureus</i> strains. Some strains for which the vancomycin MICs are 4 μg/mL will fail to grow.	Perform vancomycin MIC on <i>Enterococcus</i> spp. that grow on BHI–vancomycin screening agar and test for motility and pigment production to distinguish species with acquired resistance (eg, <i>vanA</i> and <i>vanB</i>) from those with intrinsic, intermediate-level resistance to vancomycin (eg, <i>vanC</i>), such as <i>Enterococcus gallinarum</i> and <i>Enterococcus casseliflavus</i> , which often grow on the vancomycin screen plate. In contrast to other enterococci, <i>E. casseliflavus</i> and <i>E. gallinarum</i> with vancomycin MICs of 8–16 µg/mL (intermediate) differ from vancomycin-resistant enterococci for infection prevention purposes.				
QC recommendations – routine ^b	E. faecalis ATCC®c 29212 – susceptible	E. faecalis ATCC® 29212 – susceptible				
QC recommendations – lot/shipment ^d	E. faecalis ATCC® 51299 – resistant	E. faecalis ATCC® 51299 – resistant				

Abbreviations: ATCC®, American Type Culture Collection; BHI, brain heart infusion; MIC, minimal inhibitory concentration; QC, quality control.

For Use With M02 and M07

Table 3G. (Continued)

Footnotes

- a. BHI: Even though not as widely available, dextrose phosphate agar and broth have been shown in limited testing to perform comparably.
- b. QC recommendations routine

Test negative (susceptible) QC strain:

- With each new lot/shipment of testing materials
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02¹ and M07²)
- Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met
- c. ATCC® is a registered trademark of the American Type Culture Collection.
- d. QC recommendations lot/shipment

Test positive (resistant) QC strain at minimum with each new lot/shipment of testing materials.

NOTE: Information in boldface type is new or modified since the previous edition.

References for Table 3G

- 1 CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 3H Test for Inducible Clindamycin Resistance in *Staphylococcus* spp., *Streptococcus pneumoniae*, and *Streptococcus* spp. β-Hemolytic Group

Table 3H. Test for Detecting Inducible Clindamycin Resistance in *Staphylococcus* spp., *Streptococcus pneumoniae*, and *Streptococcus* spp. 8-Hemolytic Group^{a,b}

Test	ICR							
Test method	Disk Diffusior	n (D-zone test)	Broth Microdilution					
Organism group (applies only to organisms resistant to erythromycin and susceptible or intermediate to clindamycin)	All Staphylococcus spp.	S. pneumoniae and β-hemolytic Streptococcus spp.	All Staphylococcus spp.c	S. pneumoniae and β-hemolytic Streptococcus spp.				
Medium	MHA or blood agar purity plate used with MIC tests	MHA supplemented with sheep blood (5% v/v) or TSA supplemented with sheep blood (5% v/v)	САМНВ	CAMHB with LHB (2.5% to 5% v/v)				
Antimicrobial concentration	15-µg erythromycin and 2-µg clindamycin disks spaced 15–26 mm apart	15-µg erythromycin and 2-µg clindamycin disks spaced 12 mm apart	4 μg/mL erythromycin and 0.5 μg/mL clindamycin in same well	1 μg/mL erythromycin and 0.5 μg/mL clindamycin in same well				
Inoculum	Standard disk diffusion procedure or	Standard disk diffusion procedure	Standard broth microdilu	ition procedure				
	heavily inoculated area of purity plate							
Incubation conditions	35°C±2°C; ambient air	35°C±2°C; 5% CO ₂	35°C±2°C; ambient air					
Incubation length	16–18 hours	20–24 hours	18–24 hours	20-24 hours				
Results	Flattening of the zone of inhibition disk (referred to as a D-zone) = I Hazy growth within the zone of it clindamycin resistance, even if r	CR. nhibition around clindamycin=	Any growth=ICR. No growth=no ICR.	,				

For Use With M02 and M07

Table 3H. (Continued)

Test		ICR			
Test method	Disk Diffusio	n (D-zone test)	Broth Microdilution		
Organism group (applies only	All Staphylococcus spp.	S. pneumoniae and	All Staphylococcus	S. pneumoniae and β-hemolytic	
to organisms resistant to		β-hemolytic Streptococcus	spp.c	Streptococcus spp.	
erythromycin and susceptible		spp.			
or intermediate to clindamycin)					
Additional testing and	Report isolates with ICR as "cli	ndamycin resistant."			
reporting					
				esistant based on detection of ICR,	
	as determined by testing clin	damycin in combination with e	rythromycin."		
QC recommendations – routine ^c	S. aureus ATCC®d 25923 for	S. pneumoniae ATCC® 49619	S. aureus ATCC® BAA-976™ or	S. pneumoniae ATCC® 49619 or	
Toutine	routine QC of erythromycin and clindamycin disks	for routine QC of erythromycin	S. aureus ATCC®	S. aureus ATCC® BAA-976™ –	
	and childaniyon disks	and clindamycin disks	29213 – no growth	no growth	
QC recommendations – lot/shipment ^e			S. aureus ATCC® BAA-	977™ – growth	
QC recommendations – supplemental ^f	S. aureus ATCC® BAA-976™ (D-zone test negative)	S. aureus ATCC® BAA-	976™ (no growth)	
	S. aureus ATCC® BAA-977™ (D-zone test positive)	S. aureus ATCC® BAA-	977™ (growth)	
	Use of unsupplemented MHA is	s acceptable for these strains.			
Abbroviations: ATCC® Amorican To	! !	•	eroth: ICD inducible eli	ndomicain registeres. LUD brood	

Abbreviations: ATCC[®], American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; **ICR**, **inducible clindamycin resistance**; LHB, lysed horse blood; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control; TSA, tryptic soy agar.

Footnotes

- a. Antimicrobial susceptibility testing of β-hemolytic streptococci does not need to be performed routinely (see general comment [4] in Table 2H-1). When susceptibility testing is clinically indicated, **test** for **ICR** in strains that are erythromycin resistant and clindamycin susceptible or intermediate.
- b. In accordance with 2010 guidance from the Centers for Disease Control and Prevention, colonizing isolates of group B streptococci from penicillin-allergic pregnant women should be tested for clindamycin (including ICR) (see comment [12] in Table 2H-1). For isolates that test susceptible to clindamycin (with erythromycin induction), consider adding the following comment to the patient's report: "This group B Streptococcus does not demonstrate inducible clindamycin resistance as determined by testing clindamycin in combination with erythromycin."

Test for Inducible Clindamycin Resistance in Staphylococcus spp., Streptococcus pneumoniae, and Streptococcus spp. β-Hemolytic Group

Table 3H. (Continued)

c. QC recommendations - routine

Test negative (susceptible) QC strain:

- With each new lot/shipment of testing materials
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02² and M07³)
- Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met
- d. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- e. QC recommendations lot/shipment

Test positive (resistant) QC strain at minimum with each new lot/shipment of testing materials.

- f. QC recommendations supplemental
 - Supplemental QC strains can be used to assess a new test, for training personnel, and for competence assessment. It is not necessary to include supplemental QC strains in routine daily or weekly AST QC programs. See Appendix C, which describes use of QC strains.

NOTE: Information in boldface type is new or modified since the previous edition.

References for Table 3H

- Verani JR, McGee L, Schrag SJ; Division of Bacterial Diseases, National Center for Immunization and Respiratory Diseases, Centers for Disease Control and Prevention (CDC). Prevention of perinatal group B streptococcal disease revised guidelines from CDC, 2010. MMWR Recomm Rep. 2010;59(RR-10):1-36.
- ² CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 3 CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

For Use With M02 and M07

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Table 31 Test for High-Level Mupirocin Resistance in *Staphylococcus aureus*

Table 3I. Test for Detecting High-Level Mupirocin Resistance in Staphylococcus aureus

Test		High-Level Mupirocin Resistance ^{a,1-3}
Test method	Disk Diffusion	Broth Microdilution
Organism group	S. aureus	
Medium	MHA	CAMHB
Antimicrobial concentration	200-μg mupirocin disk	Single mupirocin 256-μg/mL well
Inoculum	Standard disk diffusion procedure	Standard broth microdilution procedure
Incubation conditions	35°C±2°C; ambient air	35°C±2°C; ambient air
Incubation length	24 hours; read with transmitted light	24 hours
Additional testing and reporting	Examine carefully with transmitted light for light growth within the zone of inhibition. No zone = high-level mupirocin resistance. Any zone = the absence of high-level mupirocin resistance. Report isolates with no zone as high-level mupirocin resistant. Report any zone of inhibition as the absence	For single 256-µg/mL well: Growth = high-level mupirocin resistance. No growth = the absence of high-level mupirocin resistance. Report growth in the 256-µg/mL well as high-level mupirocin resistant. Report no growth in the 256-µg/mL well as the absence of high-level resistance.
QC recommendations – routine ^b QC recommendations – lot/shipment ^d	of high-level resistance. S. aureus ATCC®c 25923 (200-µg disk) – mupA negative (zone 29–38 mm) S. aureus ATCC® BAA-1708™ – mupA positive (no zone)	S. aureus ATCC® 29213 – mupA negative (MIC 0.06–0.5 μg/mL) or E. faecalis ATCC® 29212 – mupA negative (MIC 16–128 μg/mL) S. aureus ATCC® BAA-1708™ – mupA positive (growth in 256-μg/mL well)

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control.

Footnotes

a. Although not formally validated by CLSI document M23¹-based analyses, some studies have linked a lack of response to mupirocin-based decolonization regimens with isolates for which the mupirocin MICs are \geq 512 µg/mL.²⁻⁴ Although this document does not provide guidance on breakpoints for mupirocin, disk-based testing and the MIC test described here identify isolates for which the mupirocin MICs are \geq 512 µg/mL.

For Use With M02 and M07

Table 3I. (Continued)

b. QC recommendations – routine

Test negative (susceptible) QC strain:

- With each new lot/shipment of testing materials
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02⁵ and M07⁶)
- Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met
- c. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- d. QC recommendations lot/shipment

Test positive (resistant) QC strain at minimum with each new lot/shipment of testing materials.

References for Table 3I

- ¹ CLSI. Development of In Vitro Susceptibility Testing Criteria and Quality Control Parameters. 5th ed. CLSI guideline M23. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² Simor AE, Phillips E, McGeer A, et al. Randomized controlled trial of chlorhexidine gluconate for washing, intranasal mupirocin, and rifampin and doxycycline versus no treatment for the eradication of methicillin-resistant *Staphylococcus aureus* colonization. *Clin Infect Dis.* 2007;44(2):178-185.
- Harbarth S, Dharan S, Liassine N, Herrault P, Auckenthaler R, Pittet D. Randomized, placebo-controlled, double-blind trial to evaluate the efficacy of mupirocin for eradicating carriage of methicillin-resistant Staphylococcus aureus. Antimicrob Agents Chemother. 1999;43(6):1412-1416.
- Walker ES, Vasquez JE, Dula R, Bullock H, Sarubbi FA. Mupirocin-resistant, methicillin-resistant Staphylococcus aureus; does mupirocin remain effective? Infect Control Hosp Epidemiol. 2003;24(5):342-346.
- ⁵ CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 6 CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 3J Test for High-Level Aminoglycoside Resistance in *Enterococcus* spp.

Table 3J. Test for Detecting High-Level Aminoglycoside Resistance in *Enterococcus* spp.^a (Includes Disk Diffusion)

Test		Gentamicin HLAR			Streptomycin HLAR			
Test method	Disk diffusion	Broth microdilution	Agar dilution	Disk diffusion	Broth microdilution	Agar dilution		
Medium	MHA	BHI ^b broth	BHI ^b agar	MHA	BHI ^b broth	BHI ^b agar		
Antimicrobial concentration	120-µg gentamicin disk	Gentamicin, 500 μg/mL	Gentamicin, 500 μg/mL	300-µg streptomycin disk	Streptomycin, 1000 µg/mL	Streptomycin, 2000 µg/mL		
Inoculum	Standard disk diffusion procedure	Standard broth dilution procedure	10 μL of a 0.5 McFarland suspension spotted onto agar surface	Standard disk diffusion procedure	Standard broth dilution procedure	10 μL of a 0.5 McFarland suspension spotted onto agar surface		
Incubation conditions	35°C±2°C; ambient air	35°C±2°C; ambient air	35°C±2°C; ambient air	35°C±2°C; ambient air	35°C±2°C; ambient air	35°C±2°C; ambient air		
Incubation length	16–18 hours	24 hours	24 hours	16–18 hours	24–48 hours (if susceptible at 24 hours, reincubate)	24–48 hours (if susceptible at 24 hours, reincubate)		
Results	6 mm=resistant	Any growth = resistant	>1 colony = resistant	6 mm=resistant	Any growth = resistant	>1 colony=resistant		
	7–9 mm=inconclusive			7–9 mm=inconclusive				
	≥10 mm=susceptible			≥10 mm=susceptible				
	MIC correlates: R=>500 μg/mL S=≤500 μg/mL			MIC correlates: R=>1000 µg/mL (broth) and>2000 µg/mL (agar) S=≤1000 µg/mL (broth) and≤2000 µg/mL (agar)				
Additional testing and reporting	Susceptible: is synergis: If disk diffusion result is Strains of enterococci w	tic with cell wall–active age inconclusive: perform an agith ampicillin and penicillin Cs 16–32 µg/mL) resistancel resistance to gentamicins of penicillin (MICs ≥ 128 µ	gar dilution or broth dilution MICs ≥ 16 µg/mL are categ the may be susceptible to syrth or streptomycin, see Subchag/mL) or ampicillin (MICs ≥	n, and vancomycin). and vancomycin) that is also	enterococci with low levels cillins in combination with g h doses of penicillin or amp of be susceptible to the syn	entamicin or streptomycin (ir picillin are used. Enterococci		
QC recommendations – routine ^c	E. faecalis ATCC®d 29212: 16–23 mm	E. faecalis ATCC® 29212 – Susceptible	E. faecalis ATCC® 29212 – Susceptible	E. faecalis ATCC® 29212: 14–20 mm	E. faecalis ATCC® 29212 – Susceptible	E. faecalis ATCC® 29212 Susceptible		
QC recommendations – lot/shipment ^e		E. faecalis ATCC® 51299 – Resistant	E. faecalis ATCC® 51299 – Resistant		E. faecalis ATCC® 51299 – Resistant	E. faecalis ATCC® 51299 Resistant		

Abbreviations: ATCC®, American Type Culture Collection; BHI, brain heart infusion; CSF, cerebrospinal fluid; HLAR, high-level aminoglycoside resistance; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; QC, quality control.

For Use With M02 and M07

Table 3J. (Continued)

Footnotes

- a. Other aminoglycosides do not need to be tested, because their activities against enterococci are not superior to gentamicin and streptomycin.
- b. BHI: Even though not as widely available, dextrose phosphate agar and broth have been shown in limited testing to perform comparably.
- c. QC recommendations routine

Test negative (susceptible) QC strain:

- · With each new lot/shipment of testing materials
- Weekly if the test is performed at least once a week and criteria for converting from daily to weekly QC testing have been met (see Subchapter 4.7.2.3 in M02⁴ and M07¹)
- Daily if the test is performed less than once per week and/or if criteria for converting from daily to weekly QC testing have not been met
- d. ATCC® is a registered trademark of the American Type Culture Collection.
- e. QC recommendations lot/shipment

Test positive (resistant) QC strain at minimum with each new lot/shipment of testing materials.

References for Table 3J

- ¹ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically.* 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² Torres C, Tenorio C, Lantero M, Gastañares MJ, Baquero F. High-level penicillin resistance and penicillin-gentamicin synergy in *Enterococcus faecium*. *Antimicrob Agents Chemother*. 1993;37(11):2427-2431.
- Murray BE. Vancomycin-resistant enterococci. *Am J Med.* 1997;102(3):284-293.
- 4 CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute: 2018.

Table 4A-1. Disk <u>Diffusion QC Ranges for Nonfastidious Organisms and Antimicrobial Agents Excluding β-Lactam Combination Agents</u>

		Disk Diffusion QC Ranges, mm					
Antimicrobial Agent	Disk Content	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 25923			
Amikacin	30 μg	19–26	18–26	20–26			
Ampicillin	10 μg	15–22	-	27–35			
Azithromycin	15 μg	_	_	21–26			
Azlocillin	75 μg	-	24–30	-			
Aztreonam	30 μg	28–36	23–29	-			
Carbenicillin	100 μg	23–29	18–24	-			
Cefaclor	30 μg	23–27	_	27–31			
Cefamandole	30 μg	26–32	-	26–34			
Cefazolin	30 μg	21–27	_	29–35			
Cefdinir	5 μg	24–28	-	25–32			
Cefditoren	5 μg	22–28	_	20–28			
Cefepime	30 μg	31–37	25–31	23–29			
Cefetamet	10 μg	24–29	_	-			
Cefiderocol	30 μg	25–31	22–31	-			
Cefixime	5 μg	20–26	-	-			
Cefmetazole	30 μg	26–32	-	25–34			
Cefonicid	30 μg	25–29	_	22–28			
Cefoperazone	75 μg	28–34	23–29	24–33			
Cefotaxime	30 μg	29–35	18–22	25–31			
Cefotetan	30 μg	28–34	-	17–23			
Cefoxitin	30 μg	23–29	-	23–29			
Cefpodoxime	10 μg	23–28	-	19–25			
Cefprozil	30 μg	21–27	_	27–33			
Ceftaroline	30 μg	26–34	-	26–35			
Ceftazidime	30 μg	25–32	22–29	16–20			
Ceftibuten	30 μg	27–35	-	-			
Ceftizoxime	30 μg	30–36	12–17	27–35			
Ceftobiprole	30 μg	30–36	24–30	26–34			
Ceftriaxone	30 μg	29–35	17–23	22–28			
Cefuroxime	30 µg	20–26	-	27–35			
Cephalothin	30 μg	15–21	-	29–37			
Chloramphenicol	30 μg	21–27	-	19–26			
Cinoxacin	100 μg	26–32	-	-			

Table 4A-1. (Continued)

		Disk Diffusion QC Ranges, mm					
Antimicrobial Agent	Disk Content	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 25923			
Ciprofloxacin	5 μg	29 –38	25–33	22–30			
Clarithromycin	15 μg	_	-	26–32			
Clinafloxacin	5 μg	31–40	27–35	28–37			
Clindamycin ^c	2 μg	-	-	24–30			
Colistin	10 μg	11–17	11–17	-			
Delafloxacin ^d	5 μg	28–35	23–29	32–40			
Dirithromycin	15 μg	-	-	18–26			
Doripenem	10 μg	27–35	28–35	33–42			
Doxycycline	30 μg	18–24	-	23–29			
Enoxacin	10 μg	28–36	22–28	22–28			
Eravacycline	20 μg	16–23	-	19–26			
Ertapenem	10 μg	29–36	13–21	24–31			
Erythromycin ^c	15 μg	-	-	22–30			
Faropenem	5 μg	20–26	-	27–34			
Fleroxacin	5 μg	28–34	12–20	21–27			
Fosfomycin ^e	200 μg	22–30	-	25–33			
Fusidic acid	10 μg	-	-	24–32			
Garenoxacin	5 μg	28–35	19–25	30–36			
Gatifloxacin	5 μg	30–37	20–28	27–33			
Gemifloxacin	5 μg	29–36	19–25	27–33			
Gentamicin ^f	10 μg	19–26	17–23	19–27			
Gepotidacin	10 μg	18–26	-	23–29			
Grepafloxacin	5 μg	28–36	20–27	26–31			
Iclaprim	5 μg	14–22	-	25–33			
Imipenem ^g	10 μg	26–32	20–28	-			
Kanamycin	30 μg	17–25	-	19–26			
Lefamulin	20 μg	-	-	26–32			
Levofloxacin	5 μg	29–37	19–26	25–30			
Levonadifloxacin	10 μg	27–33 ^d	17–23 ^d	32–39 ^d			
Linezolid	30 μg	-	-	25–32 ^h			
Lomefloxacin	10 μg	27–33	22–28	23–29			
Loracarbef	30 μg	23–29	-	23–31			
Mecillinam	10 μg	24–30	-	-			

Table 4A-1. (Continued)

		Disk Diffusion QC Ranges, mm				
Antimicrobial Agent	Disk Content	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC [®] 27853	Staphylococcus aureus ATCC [®] 25923		
Meropenem	10 μg	28–35	27–33	29–37		
Minocycline	30 μg	19–25	-	25–30		
Moxalactam	30 μg	28–35	17–25	18–24		
Moxifloxacin	5 μg	28–35	17–25	28–35		
Nafcillin	1 μg	<u> </u>	_	16–22		
Nafithromycin	15 μg	_	_	25–31 ^d		
Validixic acid	30 μg	22–28	_			
Netilmicin	30 μg	22–30	17–23	22–31		
Nitrofurantoin	300 μg	20–25	_	18–22		
Norfloxacin	10 µg	28–35	22–29	17–28		
Ofloxacin	5 μg	29–33	17–21	24–28		
Omadacycline	30 μg	22–28	-	22–30		
Oxacillin	1 μg		-	18–24		
Pefloxacin	5 μg	25–33	-			
Penicillin	10 units	_	_	26–37		
Piperacillin	100 μg	24–30	25–33	-		
Plazomicin	30 μg	21–27	15–21	19–25		
Polymyxin B	300 units	13–19	14–18			
Quinupristin-dalfopristin	15 μg		-	21–28		
Razupenem	10 μg	21–26	-	_i		
Rifampin	5 μg	8–10	-	26–34		
Solithromycin	15 μg	-	-	22–30		
Sparfloxacin	5 μg	30–38	21–29	27–33		
Streptomycin ^f	10 μg	12–20	-	14–22		
Sulfisoxazole ^j	250 μg or 300 μg	15–23	_	24–34		
Sulopenem	2 µg	24-30 ^d	-	_		
Tebipenem ^g	10 µg	30–37	20–26	-		
Tedizolid ^k	2 μg	_	-	18–24 ^h		
Teicoplanin	30 μg	_	_	15–21		
Telithromycin	15 μg	_	_	24–30		
Tetracycline	30 μg	18–25	_	24–30		
Ticarcillin	75 μg	24–30	21–27			
Гigecycline	15 μg	20–27	9–13	20–25		
Tobramycin	10 μg	18–26	20–26	19–29		
Frimethoprim ^j	5 μg	21–28	_	19–26		
Frimethoprim- sulfamethoxazole ^j	1.25/23.75 μg	23–29	-	24–32		
Trospectomycin	30 μg	10–16	_	15–20		
Trovafloxacin	10 μg	29–36	21–27	29–35		
Ulifloxacin (prulifloxacin)	5 μg	32–38	27–33	20–26		
Vancomycin	30 μg	_	_	17–21		

Abbreviations: ATCC®, American Type Culture Collection, QC, quality control.

Table 4A-1. (Continued)

Footnotes

- a. Refer to Table 4A-2 for QC of β -lactam combination agents.
- b. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- c. When disk approximation tests are performed with erythromycin and clindamycin, *S. aureus* ATCC® BAA-977™ (containing inducible *erm*A-mediated resistance) and *S. aureus* ATCC® BAA-976™ (containing *msr*A-mediated macrolide-only efflux) are recommended as supplemental QC strains (eg, for training, competence assessment, or test evaluation). *S. aureus* ATCC® BAA-977™ should demonstrate inducible clindamycin resistance (ICR) (ie, a positive D-zone test), whereas *S. aureus* ATCC® BAA-976™ should not demonstrate ICR. *S. aureus* ATCC® 25923 should be used for routine QC (eg, weekly or daily) of erythromycin and clindamycin disks using standard Mueller-Hinton agar.
- d. QC ranges were established using data from only one disk manufacturer. Disks from other manufacturers were not available at the time of testing.
- e. The 200- μ g fosfomycin disk contains 50 μ g of glucose-6-phosphate.
- f. For control ranges of gentamicin 120-μg and streptomycin 300-μg disks, use *E. faecalis* ATCC® 29212 (gentamicin: 16–23 mm; streptomycin: 14–20 mm).
- g. Klebsiella pneumoniae ATCC® 700603 is a supplemental QC strain for testing QC of imipenem (25–33 mm) and tebipenem (26–32 mm).
- h. Zones of inhibition for linezolid and tedizolid with S. aureus ATCC® 25923 should be read using transmitted light.
- Razupenem tested with S. aureus ATCC[®] 25923 can often produce the double or target zone phenomenon. For accurate QC results, use S. aureus ATCC[®] 29213 (no double zones) with acceptable range 33–39 mm.
- j. These agents can be affected by excess levels of thymidine and thymine. See M02,1 Subchapter 3.1.1.2 for guidance, should a problem with QC occur.
- k. E. faecalis ATCC® 29212 is a supplemental QC strain for testing QC of tedizolid (14–21 mm) to assist with reading.
- I. Ulifloxacin is the active metabolite of the prodrug prulifloxacin. Only ulifloxacin should be used for antimicrobial susceptibility testing.

NOTE: Information in boldface type is new or modified since the previous edition.

Reference for Table 4A-1

CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

M100, 30th ed.

Table 4A-2. Disk Diffusion QC Ranges for Nonfastidious Organisms and β-Lactam Combination Agents^a

		QC Organisms and Characteristics									
		Escherichia coli ATCC®b 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 25923	Escherichia coli ATCC® 35218 ^{c,d}	Klebsiella pneumoniae ATCC® 700603 ^{c,d} SHV-18 OXA-2 Mutations in	Escherichia coli NCTC 13353 ^{c,d}	Klebsiella pneumoniae ATCC® BAA- 1705™c,d	Klebsiella pneumoniae ATCC® BAA-2814™	Acinetobacter baumannii NCTC 13304 ^{c,d}	
Antimicrobial Agent	Disk Content	β- lactamase negative	Inducible AmpC	β-lactamase negative, <i>mec</i> A negative	TEM-1	OmpK35 and OmpK37 TEM-1 meter QC Range	CTX-M-15	KPC-2 SHV	KPC-3 SHV-11 TEM-1	OXA-27	
Amoxicillin-clavulanate	20/10 μg	18–24	_	28–36	17–22	lineter QO Kange	- -	_	_	_	
(2:1)		_				_					
Ampicillin	10 μg	15–22	_	27–35	6	-	_	_	_	_	
Ampicillin-sulbactam (2:1)	10/10 μg	19–24	_	29–37	13–19	-	_	-	-	-	
Aztreonam	30 μg	28–36	23–29	_	31–38	10–16	_	-	_	_	
Aztreonam-avibactam	30/20 μg	32–38	24–30	-	31–38	26–32 ^e	_	_	-	-	
Cefepime	30 μg	31–37	25–31	23–29	31–37	23–29	6–15 ^f	_	-	6–16 ^f	
Cefepime- enmetazobactam ^e	30/20 μg	32–38	26–32	-	32–38	26–32	27–33	-	-	-	
Cefepime- taniborbactam	30/20 μg	31–37	25–31	-	31–37	24–31	24–30	22–27	-	-	
Cefepime-tazobactam	30/20 μg	32–37	27–31	24–30	-	25–30 ^e	27–31	-	-	_	
Cefepime-zidebactam	30/30 µg	33–40	29–35	_	_	28–34	29–35	-	_	19–25	
Cefotaxime	30 μg	29–35	18–22	25–31	-	17–25	ı	_	-	_	
Cefpodoxime	10 μg	23–28	ı	19–25	-	9–16	ı	_		-	
Ceftaroline	30 μg	26–34	-	26–35	-	_	_	_	-	-	
Ceftaroline-avibactam	30/15 μg	27–34	17–26	25–34	27–35	21–27 ^e	_	-	-	-	
Ceftazidime	30 μg	25–32	22–29	16–20	-	10–18	1	-	-	_	
Ceftazidime-avibactam	30/20 μg	27–35	25–31	16–22	28–35	21–27 ^e	-	-	-	-	
Ceftolozane-tazobactam	30/10 μg	24–32	25–31	10–18	25–31	17–25	1	_	-	_	
Ceftriaxone	30 μg	29–35	17–23	22–28	_	16–24		_	_	_	
Imipenem	10 μg	26-32	20-28	_		25–33	ı	11–22	6–14	_	
Imipenem-relebactam ^{e,g}	10/25 μg	27–33	26–31	-		26–32	1	23–29	22–28	_	
Meropenem ^f	10 μg	28–35	27–33	29–37	_	-	_	11–18 ^e	6 ^e	-	

Table 4A-2. (Continued)

					QC Organis	ms and Chara	cteristics			
		Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 25923	Escherichia coli ATCC [®] 35218 ^{c,d}	Klebsiella pneumoniae ATCC® 700603 ^{c,d}	Escherichia coli NCTC 13353 ^{c,d}	Klebsiella pneumoniae ATCC [®] BAA- 1705™ ^{c,d}	Klebsiella pneumoniae ATCC® BAA-2814™	Acinetobacter baumannii NCTC 13304 ^{c,d}
Antimicrobial	Disk	β-lactamase negative	Inducible AmpC	β-lactamase negative, <i>mec</i> A negative	TEM-1	SHV-18 OXA-2 Mutations in OmpK35 and OmpK37 TEM-1	CTX-M-15	KPC-2 SHV	KPC-3 SHV-11 TEM-1	OXA-27
Agent	Content				Zone Dia	meter QC Range	es, mm			
Meropenem- vaborbactam ^g	20/10 μg	31–37	29–35	32–38	-	29–35	-	21–27	16–20	-
Piperacillin	100 μg	24-30	25-33	-	12–18	-	_	_	-	_
Piperacillin- tazobactam	100/10 μg	24–30	25–33	27–36	24–30	_	ı	ı	-	-
Sulbactam- durlobactam	10/10 μg	26–32	1	_	ı	_	-	-	-	24–30
Ticarcillin	75 μg	24–30	21–27	-	6	_	-	-	-	_
Ticarcillin- clavulanate	75/10 μg	24–30	20–28	29–37	21–25	-	-	-	-	-

Abbreviations: ATCC®, American Type Culture Collection; MIC, minimal inhibitory concentration; N/A, not applicable; NCTC, National Collection of Type Cultures; QC, quality control.

QC strain selection codes:

QC strain is recommended for routine QC.

Test one of these agents by a disk diffusion or MIC method to confirm the integrity of the respective QC strain.c,d

Footnotes

- a. Unsupplemented Mueller-Hinton medium. See Table 4A-1 for QC ranges for combination agents from other drug classes.
- b. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- c. Careful attention to organism maintenance (eg, minimal subcultures) and storage (eg, -60°C or below) is especially important for these QC strains because spontaneous loss of the plasmid encoding the β-lactamase has been documented. If stored at temperatures above -60°C or if repeatedly subcultured, these strains may lose their resistance characteristics and QC results may be outside the acceptable ranges.
- d. To confirm the integrity of the QC strain, test one of the single β-lactam agents highlighted in orange by either a disk diffusion or MIC test method when the strain is first subcultured from a frozen or lyophilized stock culture. In some cases, only MIC ranges are available to accomplish this confirmation (see Table 5A-2). In-range results for the single agent indicate the QC strain is reliable for QC of β-lactam combination agents. It is not necessary to check the QC strain again with a single agent until a new frozen or lyophilized stock culture is put into use, providing recommendations for handling QC strains as described in M02¹ and M07² are followed.

Table 4A-2. (Continued)

- e. QC ranges were established using data from only one disk manufacturer. Disks from other manufacturers were not available at the time of testing.
- If discrete colonies or a haze of growth are present inside the zone of inhibition, measure the colony-free inner zone.
- Either strain highlighted in green may be used for routine QC of this antimicrobial agent.

NOTE: Information in boldface type is new or modified since the previous edition.

References for Table 4A-2

- CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

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Table 4B. Disk Diffusion QC Ranges for Fastidious Organisms

		Disk Diffusion QC Ranges, mm						
Antimicrobial Agent	Disk Content	Haemophilus influenzae ATCC® ^a 49247	Haemophilus influenzae ATCC® 49766	Neisseria gonorrhoeae ATCC® 49226	Streptococcus pneumoniae ATCC® 49619 ^b			
Amoxicillin-clavulanate ^c	20/10 μg	15–23	_	_	_			
Ampicillin	10 μg	13–21	_	-	30–36			
Ampicillin-sulbactam	10/10 μg	14–22	_	-	-			
Azithromycin	15 μg	13–21	_	30–38	19–25			
Aztreonam	30 μg	30–38	_	-	-			
Cefaclor	30 μg	-	25–31	-	24–32			
Cefdinir	5 μg	_	24–31	40–49	26–31			
Cefditoren	5 μg	25–34	_	-	27–35			
Cefepime	30 μg	25–31	_	37–46	28-35			
Cefetamet	10 μg	23–28	_	35–43	_			
Cefixime	5 μg	25–33	_	37–45	16–23			
Cefmetazole	30 µg	16–21	_	31–36	_			
Cefonicid	30 μg	-	30–38	-	_			
Cefotaxime	30 µg	31–39	_	38–48	31–39			
Cefotetan	30 µg	_	_	30–36	_			
Cefoxitin	30 µg	_	_	33–41	_			
Cefpodoxime	10 μg	25–31	-	35–43	28–34			
Cefprozil	30 μg	_	20–27	_	25–32			
Ceftaroline	30 μg	29–39	_	_	31–41			
Ceftaroline-avibactam ^d	30/15 μg	30–38	_	_	_			
Ceftazidime	30 μg	27–35	_	35–43	_			
Ceftazidime-avibactam ^d	30/20 μg	28–34	_	-	23–31			
Ceftibuten	30 μg	29–36	-	_				
Ceftizoxime	30 μg	29–39	-	42–51	28–34			
Ceftobiprole ^e	30 μg	28–36	30–38	-	33–39			
	30/10 μg	23–29	_	_	21–29			
Ceftolozane-tazobactam ^d								
Ceftriaxone	30 μg	31–39	-	39–51	30–35			
Cefuroxime	30 μg	_	28–36	33–41				
Cephalothin	30 μg	-	_	-	26–32			
Chloramphenicol	30 μg	31–40		-	23–27			
Ciprofloxacin	5 μg	34–42	_	48–58				
Clarithromycin	15 μg	11–17		-	25–31			
Clinafloxacin	5 μg	34–43	<u> </u>	-	27–34			
Clindamycin	2 μg	-		-	19–25			
Delafloxacin	5 μg	40–51		-	28-36 ^f			
Dirithromycin	15 μg	_	_	-	18–25			
Doripenem	10 μg	21–31	<u> </u>	_	30–38			
Doxycycline	30 μg	-	-	_	25–34			
Enoxacin	10 μg	_	<u> </u>	43–51	_			
Eravacycline	20 μg	_		_	23–30			
Ertapenem ^e	10 μg	20–28	27–33	-	28–35			
Erythromycin	15 μg	_	_	_	25–30			

Table 4B. (Continued)

		Disk Diffusion QC Ranges, mm					
Antimicrobial Agent	Disk Content	Haemophilus influenzae ATCC®a 49247	Haemophilus influenzae ATCC® 49766	Neisseria gonorrhoeae ATCC® 49226	Streptococcus pneumoniae ATCC® 49619 ^b		
Faropenem	5 μg	15–22	_	_	27–35		
Fleroxacin	5 μg	30–38	_	43–51	_		
Fusidic acid	10 μg	_	_	_	9–16		
Garenoxacin	5 μg	33–41	_	_	26–33		
Gatifloxacin	5 μg	33–41	_	45–56	24–31		
Gemifloxacin	5 μg	30–37	_	_	28–34		
Gepotidacin	10 μg	_	_	32–40	22–28		
Grepafloxacin	5 μg	32–39	_	44–52	21–28		
Iclaprim	5 μg	24–33	_	_	21–29		
Imipenem	10 μg	21–29	_	_	-		
Lefamulin	20 μg	22–28	_	_	19–27		
Levofloxacin	5 μg	32–40	_	_	20–25		
Levonadifloxacin	10 μg	33–41 ^f	-	-	24–31 ^f		
Linezolid	30 μα	_	-	_	25–34		
Lomefloxacin	10 μg	33–41	-	45–54	_		
Loracarbef	30 μg	_	26–32	_	22–28		
Meropenem	10 μg	20–28	_	_	28–35		
Moxifloxacin	5 μg	31–39	_	_	25–31		
Nafithromycin	15 μg	16–20 ^f	_	_	25–31 ^f		
Nitrofurantoin	300 μg	-	_	_	23–29		
Norfloxacin	10 μg	_	_	_	15–21		
Ofloxacin	5 μg	31–40	=	43–51	16–21		
Omadacycline	30 µg	21–29	_	_	24–32		
Oxacillin	1 μg		_	_	≤12 ^g		
Penicillin	10 units	_		26–34	24–30		
Piperacillin-tazobactam	100/10 μg	33–38		-			
Quinupristin-dalfopristin	15 μg	15–21	_	_	19–24		
Razupenem	10 μg	24–30	_	_	29–36		
Rifampin	5 μg	22–30	_	_	25–30		
Solithromycin	15 μg	16–23	_	33–43	25–33		
Sparfloxacin	5 μg	32–40	_	43–51	21–27		
Spectinomycin	100 μg	-	_	23–29			
Tedizolid	2 μg	_	_	_	18–25		
Telithromycin	15 μg	17–23	_	_	27–33		
Tetracycline	30 μg	14–22	_	30–42	27–31		
Tigecycline	15 μg	23–31		30–40	23–29		
Trimethoprim-sulfamethoxazole	1.25/23.75 µg	24–32		-	20–28		
Trospectomycin	30 μg	22–29		28–35			
Trovafloxacin	30 μg 10 μg	32–39		42–55	25–32		
Vancomycin	30 μg	-		-	20–27		

Table 4B. (Continued)

Disk Diffusion Testing Conditions for Clinical Isolates and Performance of QC

bisk billusion resting conditions for clinical isolates and reflormance of wo								
Organism	H. influenzae	N. gonorrhoeae	Streptococci and <i>N. meningitidis</i>					
Medium	НТМ	GC agar base and 1% defined growth supplement. The use of a cysteine-free growth supplement is not required for disk diffusion testing.	MHA supplemented with 5% defibrinated sheep blood MH-F agar for <i>S. pneumoniae</i> only					
Inoculum	Colony suspension	Colony suspension	Colony suspension					
Incubation characteristics	5% CO ₂ ; 16–18 hours; 35°C	5% CO ₂ ; 20–24 hours; 35°C	5% CO ₂ ; 20–24 hours; 35°C					

Abbreviations: ATCC®, American Type Culture Collection; HTM, *Haemophilus* test medium; MHA, Mueller-Hinton agar; **MH-F agar, Mueller-Hinton fastidious** agar; QC, quality control.

Footnotes

- a. ATCC® is a registered trademark of the American Type Culture Collection.
- b. Despite the lack of reliable disk diffusion breakpoints for *S. pneumoniae* with certain β-lactams, *S. pneumoniae* ATCC® 49619 is the strain designated for QC of all disk diffusion tests with all *Streptococcus* spp.
- c. When testing on HTM incubated in ambient air, the acceptable QC limits for E. coli ATCC® 35218 are 17–22 mm for amoxicillin-clavulanate.
- d. QC limits for *E. coli* ATCC® 35218 in HTM: ceftaroline-avibactam 26–34 mm; ceftazidime-avibactam 27–34 mm; ceftolozane-tazobactam 25–31 mm.
- e. Either H. influenzae ATCC® 49247 or 49766 may be used for routine QC testing.
- f. QC ranges for delafloxacin, levonadifloxacin, and nafithromycin were established using data from only one disk manufacturer. Disks from other manufacturers were not available at the time of testing.
- g. Deterioration in oxacillin disk content is best assessed with QC organism S. aureus ATCC® 25923, with an acceptable zone diameter of 18–24 mm.

NOTE: Information in boldface type is new or modified since the previous edition.

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Table 4C. Disk Diffusion Reference Guide to QC Frequency

This table summarizes the suggested QC frequency when modifications are made to antimicrobial susceptibility test systems (refer to CLSI document EP23 $^{\text{TM}1}$). It applies only to antimicrobial agents for which satisfactory results have been obtained with either the 15-replicate (3- × 5-day) plan or 20 or 30 consecutive test day plan. Otherwise QC is required each test day.

		Recommended (QC Frequency	
T of Market and a			15-Replicate Plan or	0
Test Modification	1 Day	5 Days	20- or 30-Day Plan	Comments
Disks				
Use new shipment or lot number.	X			
Use new manufacturer.	Х			
Addition of new antimicrobial agent to			×	
existing system.				In addition, perform in-house verification studies.
Media (prepared agar plates)				
Use new shipment or lot number.	X			
Use new manufacturer.		Х		
Inoculum preparation				
Convert inoculum preparation/				Example:
standardization to use of a device that has				Convert from visual adjustment of turbidity to use of a
its own QC protocol.		X		photometric device for which a QC procedure is
40 process				provided.
Convert inoculum preparation/				Example:
standardization to a method that depends			X	Convert from visual adjustment of turbidity to another
on user technique.				method that is not based on a photometric device.
Measuring zones				
Change method of measuring zones.				Example:
Change meaned of meadaring zeroe.				Convert from manual zone measurements to automated
			×	zone reader.
			,	Zono roddor.
				In addition, perform in-house verification studies.
Instrument/software (eg, automated zone	reader)			in addition, portorn in nodoc vornication stadios.
Software update that affects AST results				Monitor all drugs, not just those implicated in software
Contware update that affects ACT results		Х		modification.
Repair of instrument that affects AST				Depending on extent of repair (eg, critical component
results	X			such as the photographic device), additional testing may
				be appropriate (eg, 5 days).
Abbreviations: AST, antimicrobial susceptibility	testing: OC	quality control		· - · ·

Abbreviations: AST, antimicrobial susceptibility testing; QC, quality control.

Table 4C. (Continued)

NOTE 1: QC can be performed before or concurrent with testing patient isolates. Patient results can be reported for that day if QC results are within the acceptable limits.

NOTE 2: Manufacturers of commercial or in-house-prepared tests should follow their own internal procedures and applicable regulations.

NOTE 3: For troubleshooting out-of-range results, refer to M02,² Subchapter 4.8 and M100 Table 4D. Additional information is available in Appendix C (eg, QC organism characteristics, QC testing recommendations).

NOTE 4: Broth, saline, and/or water used to prepare an inoculum does not need routine QC.

References for Table 4C

- CLSI. Laboratory Quality Control Based on Risk Management; Approved Guideline. CLSI document EP23-A™. Wayne, PA: Clinical and Laboratory Standards Institute; 2011.
- ² CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 4D. Disk Diffusion Troubleshooting Guide

This table provides guidance for troubleshooting and corrective action for out-of-range QC, primarily using antimicrobial susceptibility tests with MHA. Refer to M02,¹ Chapter 4, for additional information. Out-of-range QC tests are often the result of contamination or the use of an incorrect QC strain; corrective action should first include repeating the test with a pure culture of a freshly subcultured QC strain. If the issue is unresolved, this troubleshooting guide should be consulted regarding additional suggestions for troubleshooting out-of-range QC results and unusual clinical isolate results. In addition, see general corrective action outlined in M02¹ and notify manufacturers of potential product problems.

General Comment

(1) QC organism maintenance: Avoid repeated subcultures. Retrieve new QC strain from stock (refer to M02, Subchapter 4.4). If using lyophilized strains, follow the maintenance recommendations of the manufacturer.

Antimicrobial Agent	QC Strain	Observation	Probable Cause	Comments/Suggested Actions
β-LACTAMS				
β-lactam combination agents	A. baumannii ATCC®a 13304 E. coli ATCC® 35218 E. coli ATCC® 13353 K. pneumoniae ATCC® 700603 K. pneumoniae ATCC® BAA-1705™	Zone too large or susceptible for single β -lactam agent; in range for combination β -lactam agent	Spontaneous loss of the plasmid encoding the β -lactamase	Obtain new frozen or lyophilized stock culture. Use other routine QC strains (if available). These strains should be stored at −60°C or below, and frequent subcultures should be avoided. NOTE: <i>K. pneumoniae</i> BAA-2814™ is stable and does not require QC integrity check.
β-lactam combination agents	A. baumannii ATCC® 13304 E. coli ATCC® 35218 E. coli ATCC® 13353 K. pneumoniae ATCC® 700603 K. pneumoniae ATCC® BAA-1705™ K. pneumoniae ATCC® BAA-2814™	Zone too small or resistant for both the single β-lactam agent and the combination β-lactam agent	Antimicrobial agent is degrading.	Use alternative lot of test materials. Check storage and package integrity. Imipenem and clavulanate are especially labile.
Carbenicillin	P. aeruginosa ATCC® 27853	Zone too small	QC strain develops resistance after repeated subculture.	See general comment (1) on QC strain maintenance.
Cefepime	A. baumannii NCTC 13304 E. coli NCTC 13353	QC strain integrity test	Discrete colonies may grow within the zone of inhibition when this organism is tested with cefepime 30-µg disk.	If this occurs, measure the colony-free inner zone.
Imipenem	K. pneumoniae ATCC® BAA-1705™ K. pneumoniae ATCC® BAA-2814™	QC strain integrity test	Discrete colonies may grow within the zone of inhibition when this organism is tested with cefepime. 30-µg disk.	If this occurs, measure the colony-free inner zone.
Penicillins	Any	Zone too large	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO ₂ incubation, which lowers pH.
Penicillins	Any	Zone too small	pH of media too high	Acceptable pH range = 7.2–7.4

Table 4D. (Continued)

Antimicrobial Agent	QC Strain	Observation	Probable Cause	Comments/Suggested Actions
NON-β-LACTAMS		·		
β-lactam group	Any	Zone initially acceptable, but decreases to possibly be out of range over time	Imipenem, clavulanate, and cefaclor are especially labile. Disks have lost potency.	Use alternative lot of disks. Check storage conditions and package integrity.
Aminoglycosides Quinolones	Any	Zone too small	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO ₂ incubation, which lowers pH.
Andrew where a distant	Any	Zone too large	pH of media too high	Acceptable pH range = 7.2–7.4
Aminoglycosides	P. aeruginosa ATCC® 27853	Zone too small	Ca++ and/or Mg++ content too high	Use alternative lot of media.
Aminoglycosides	P. aeruginosa ATCC® 27853	Zone too large	Ca++ and/or Mg++ content too low	Use alternative lot of media.
Clindamycin Macrolides	S. aureus ATCC® 25923	Zone too small	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO ₂ incubation, which lowers pH.
	S. aureus ATCC® 25923	Zone too large	pH of media too high	Acceptable pH range = 7.2–7.4
Quinolones	Any	Zone too small	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO ₂ incubation, which lowers pH.
Quinolones	Any	Zone too large	pH of media too high	Acceptable pH range = 7.2–7.4
Tedizolid	E. faecalis ATCC® 29212	Zone with Enterococcus spp. is difficult to read	Light growth on MHA	E. faecalis ATCC® 29212 is provided as supplemental QC to assist in personnel training and assessment of proper reading. Measure zone edge where there is a significant decrease in density of growth when using transmitted light as illustrated in the photographs.b
Tetracyclines	Any	Zone too large	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO ₂ incubation, which lowers pH.
Tetracyclines	Any	Zone too small	pH of media too high	Acceptable pH range = 7.2–7.4
Tetracyclines	Any	Zone too small	Ca++ and/or Mg++ content too high	Use alternative lot of media.
Tetracyclines	Any	Zone too large	Ca++ and/or Mg++ content too low	Use alternative lot of media.
Sulfonamides Trimethoprim Trimethoprim- sulfamethoxazole	E. faecalis ATCC® 29212	Zone ≤ 20 mm	Media too high in thymidine content	Use alternative lot of media.

Table 4D. (Continued)

Antimicrobial Agent	QC Strain	Observation	Probable Cause	Comments/Suggested Actions
ALL AGENTS				
Various	Various	Zone too small	Contamination	Measure zone edge with visible growth detected with unaided eye. Subculture to
			Use of magnification to read zones	determine purity and repeat if necessary.
Various	Any	Many zones too small	Inoculum too heavy Error in inoculum preparation	Repeat using McFarland 0.5 turbidity standard or standardizing device. Check expiration date and proper storage if using
			Media depth too thick	barium sulfate or latex standards. Use agar with depth approximately 4 mm. Recheck alternate lots of MHA.
Various	Any	One or more zones too small or too large	Measurement error	Recheck readings for measurement or transcription errors.
			Transcription error	
			Random defective disk	Retest. If retest results are out of range and no errors are detected, initiate corrective action.
			Disk not pressed firmly against agar	action.
Various	Various	Zone too large	Did not include lighter growth in zone measurement (eg, double zone, fuzzy	Measure zone edge with visible growth detected with unaided eye.
Various	S. pneumoniae ATCC®	Zones too large	zone edge) Inoculum source plate too old and	Subculture QC strain and repeat QC test or
various	49619	Zones too large	contains too many nonviable cells.	retrieve new QC strain from stock.
	49019	Lawn of growth scanty	Plate used to prepare inoculum	Tetrieve new QC strain nom stock.
		Lawii oi giowai ocanty	should be 18–20 hours.	
Various	Any	QC results from one	One QC strain may be a better	Retest this strain to confirm reproducibility of
		strain are out of range,	indicator of a QC problem.	acceptable results.
		but results from other QC		
		strain(s) is in range with the same antimicrobial		Evaluate with alternative strains with known MICs.
		agent.		Initiate corrective action with problem QC strain/antimicrobial agent(s).
Various	Any	QC results from two strains are out of range	A problem with the disk	Use alternative lot of disks.
		with the same antimicrobial agent.		Check storage conditions and package integrity.
Various	Any	Zones overlap.	Too many disks per plate	Place no more than 12 disks on a 150-mm plate and 5 disks on a 100-mm plate; for some fastidious bacteria that produce large zones, use fewer.

Abbreviations: ATCC®, American Type Culture Collection; MHA, Mueller-Hinton agar; MIC, minimal inhibitory concentration; NCTC, National Collection of Type Cultures; pH, negative logarithm of hydrogen ion concentration; QC, quality control.

Footnotes

- a. ATCC® is a trademark of the American Type Culture Collection.
- b. Figure 1 shows examples of tedizolid disk diffusion results for *E. faecalis*.

Table 4D. (Continued)

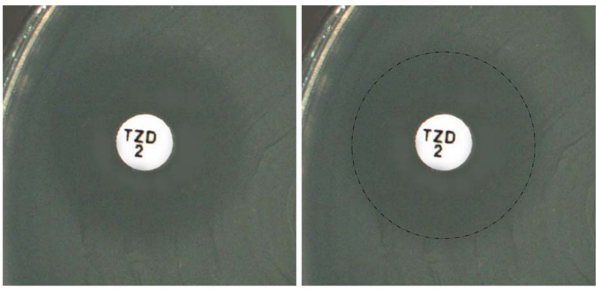


Figure 1. Measuring the Tedizolid Zone for *E. faecalis* ATCC[®] 29212 When Light Growth Is Observed. (Courtesy of Laura M. Koeth, Laboratory Specialists, Inc. Used with permission.)

NOTE: Information in boldface type is new or modified since the previous edition.

Reference for Table 4D

¹ CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 5A-1. MIC QC Ranges for Nonfastidious Organisms and Antimicrobial Agents Excluding β-Lactam Combination Agents^a

	MIC QC Ranges, μg/mL							
Antimicrobial Agent	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 29213	Enterococcus faecalis ATCC® 29212				
Amikacin	0.5–4	1–4	1–4	64–256				
Amikacin-fosfomycin (5:2) ^c	0.25/0.1–2/0.8	1/0.4-8/3.2	0.5/0.2-4/1.6	32/12.8-128/51.2				
Amoxicillin	_	_	_	_				
Ampicillin	2–8	_	0.5–2	0.5–2				
Azithromycin	_	_	0.5–2	_				
Azlocillin	8–32	2–8	2–8	1–4				
Aztreonam	0.06-0.25	2–8	_	_				
Besifloxacin	0.06-0.25	1–4	0.016-0.06	0.06-0.25				
Biapenem	0.03-0.12	0.5–2	0.03-0.12	_				
Cadazolid	_	=	0.06-0.5	0.06-0.25				
Carbenicillin	4–16	16–64	2–8	16–64				
Cefaclor	1–4	=	1–4	_				
Cefamandole	0.25–1	ı	0.25–1	_				
Cefazolin	1–4	=	0.25–1	_				
Cefdinir	0.12-0.5	ı	0.12-0.5	_				
Cefditoren	0.12–1	-	0.25–2	_				
Cefepime	0.016-0.12	0.5–4	1–4	_				
Cefetamet	0.25–1	ı	_	_				
Cefiderocol ^d	0.06-0.5	0.06–0.5	_	_				
Cefixime	0.25–1	_	8–32	_				
Cefmetazole	0.25–1	>32	0.5–2	_				
Cefonicid	0.25–1	_	1–4	_				
Cefoperazone	0.12-0.5	2–8	1–4	_				
Cefotaxime	0.03-0.12	8–32	1–4	_				
Cefotetan	0.06-0.25	_	4–16	_				
Cefoxitin	2–8	=	1–4	_				
Cefpodoxime	0.25–1	ı	1–8	_				
Cefprozil	1–4	-	0.25–1	-				
Ceftaroline	0.03-0.12	_	0.12–0.5	0.25–2 ^e				
Ceftazidime	0.06-0.5	1–4	4–16	_				
Ceftibuten	0.12-0.5	_	_	_				
Ceftizoxime	0.03-0.12	16–64	2–8	_				
Ceftobiprole	0.03-0.12	1–4	0.12–1	0.06-0.5				
Ceftriaxone	0.03-0.12	8–64	1–8	_				
Cefuroxime	2–8		0.5–2	_				
Cephalothin	4–16	-	0.12-0.5	_				

For Use With M07—MIC Testing

Table 5A-1. (Continued)

		MIC QC	Ranges, µg/mL	
Antimicrobial Agent	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 29213	Enterococcus faecalis ATCC® 29212
Chloramphenicol	2–8	_	2–16	4–16
Cinoxacin	2–8	_	-	_
Ciprofloxacin ^f	0.004–0.016	0.12–1	0.12-0.5	0.25–2
Clarithromycin	_	_	0.12-0.5	_
Clinafloxacin	0.002–0.016	0.06–0.5	0.008-0.06	0.03-0.25
Clindamycin ^g	-	_	0.06-0.25	4–16
Colistin	0.25–2	0.5–4	=	_
Dalbavancin ^h	-	_	0.03-0.12	0.03-0.12
Daptomycin ⁱ	_	_	0.12–1	1–4
Delafloxacin	0.008-0.03	0.12-0.5	0.001-0.008	0.016-0.12
Dirithromycin	-	_	1–4	_
Doripenem	0.016–0.06	0.12–0.5	0.016–0.06	1–4
Doxycycline	0.5–2	-	0.12-0.5	2–8
Enoxacin	0.06-0.25	2–8	0.5–2	2–16
Eravacycline	0.016 –0.12	2–16	0.016-0.12	0.016-0.06
Ertapenem	0.004-0.016	2–8	0.06-0.25	4–16
Erythromycin ^g	-	_	0.25–1	1–4
Exebacase ^j	-	_	0.25–2	8–64
Faropenem	0.25–1	_	0.03-0.12	_
Fidaxomicin	_	_	2–16	1–4
Finafloxacin	0.004-0.03	1–8	0.03-0.25	0.25–1
Fleroxacin	0.03-0.12	1–4	0.25–1	2–8
osfomycin ^k	0.5–2	2–8	0.5–4	32–128
Fusidic acid	-	_	0.06-0.25	_
Garenoxacin	0.004-0.03	0.5–2	0.004-0.03	0.03-0.25
Gatifloxacin	0.008-0.03	0.5–2	0.03-0.12	0.12-1.0
Gemifloxacin	0.004-0.016	0.25–1	0.008-0.03	0.016-0.12
Gentamicin ^I	0.25–1	0.5–2	0.12–1	4–16
Gepotidacin	1–4	_	0.12–1	_
Grepafloxacin	0.004-0.03	0.25–2.0	0.03–0.12	0.12–0.5
claprim	1–4	-	0.06-0.25	0.004-0.03
mipenem	0.06-0.25	1–4	0.016–0.06	0.5–2
Kanamycin	1–4		1–4	16–64
_efamulin		_	0.06–0.25	-
evofloxacin	0.008-0.06	0.5–4	0.06-0.5	0.25–2
Levonadifloxacin	0.03-0.25	0.5–4	0.008-0.03	-
_inezolid ^m	_	_	1–4	1–4
_omefloxacin	0.03-0.12	1–4	0.25–2	2–8
_oracarbef	0.5–2	> 8	0.5–2	_

	MIC QC Ranges, μg/mL							
Antimicrobial Agent	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 29213	Enterococcus faecalis ATCC® 29212				
Mecillinam	0.03–0.25 ⁿ	-	_	-				
Meropenem	0.008-0.06	0.12–1	0.03-0.12	2–8				
Minocycline ^f	0.25–1	-	0.06-0.5	1–4				
Moxalactam	0.12–0.5	8–32	4–16	_				
Moxifloxacin	0.008-0.06	1–8	0.016-0.12	0.06-0.5				
Nafcillin	_	_	0.12-0.5	2–8				
Nafithromycin	_	_	0.06-0.25	0.016-0.12				
Nalidixic acid ^f	1–4	_	_	-				
Netilmicin	≤ 0.5–1	0.5–8	≤ 0.25	4–16				
Nitrofurantoin	4–16	=	8–32	4–16				
Norfloxacin	0.03-0.12	1–4	0.5–2	2–8				
Ofloxacin	0.016-0.12	1–8	0.12–1	1–4				
Omadacycline ^o	0.25–2	_	0.12–1	0.06-0.5				
Oritavancin ^h	_	-	0.016-0.12	0.008-0.03				
Oxacillin	_	_	0.12-0.5	8–32				
Ozenoxacin	_	1	0.001-0.004	0.015-0.06				
Penicillin	_		0.25–2	1–4				
Pexiganan	2–8	2–16	8–32	16–64				
Piperacillin	1–4	1–8	1–4	1–4				
Plazomicin	0.25–2	1–4	0.25–2	-				
Polymyxin B	0.25–2	0.5–2	_	_				
Quinupristin-dalfopristin	_	ı	0.25–1	2–8				
Razupenem	0.06-0.5	ı	0.008-0.03	0.25–1				
Rifampin	4–16	16–64	0.004-0.016	0.5–4				
Solithromycin	_	_	0.03–0.12	0.016-0.06				
Sparfloxacin	0.004–0.016	0.5–2	0.03-0.12	0.12-0.5				
Sulfisoxazole ^{f,p}	8–32	ı	32–128	32–128				
Sulopenem	0.016–0.06		0.016–0.12	2–8				
Tebipenem	0.008-0.03	1–8	0.015–0.06	0.25–1				
Tedizolid ^q	_	1	0.12–1	0.25–1				
Teicoplanin	-	1	0.25–1	0.25–1				
Telavancin ^h	-	=	0.03-0.12	0.03-0.12				
Telithromycin	_	_	0.06-0.25	0.016-0.12				
Tetracycline	0.5–2	8–32	0.12–1	8–32				
Ticarcillin	4–16	8–32	2–8	16–64				
Tigecycline ^o	0.03-0.25	_	0.03-0.25	0.03-0.12				
Tobramycin	0.25–1	0.25–1	0.12–1	8–32				

For Use With M07—MIC Testing

Table 5A-1. (Continued)

		MIC QC Ranges, μg/mL							
Antimicrobial Agent	Escherichia coli ATCC ^{®b} 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 29213	Enterococcus faecalis ATCC® 29212					
Trimethoprim ^p	0.5–2	> 64	1–4	0.12-0.5					
Trimethoprim- sulfamethoxazole ^p (1:19)	≤ 0.5/9.5	8/152–32/608	≤ 0.5/9.5	≤ 0.5/9.5					
Trospectomycin	8–32	_	2–16	2–8					
Trovafloxacin	0.004-0.016	0.25–2	0.008-0.03	0.06-0.25					
Ulifloxacin (prulifloxacin) ^r	0.004–0.016	0.12-0.5	-	-					
Vancomycin ^s	-	-	0.5–2	1–4					
Zidebactam	0.06-0.25	1–8	_	_					
Zoliflodacin	1–4	-	0.12-0.5	0.25-2					

Abbreviations: ATCC[®], American Type Culture Collection; **CAMHB, cation-adjusted Mueller-Hinton broth;** MHB, Mueller-Hinton broth; MIC, minimal inhibitory concentration; QC, quality control.

Footnotes

- a. Refer to Table 5A-2 for QC of β-lactam combination agents.
- b. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- c. QC ranges reflect MICs obtained when medium is supplemented with 25 μg/mL of glucose-6-phosphate.
- d. QC ranges reflect MICs obtained when CAMHB is iron depleted. Chelation is used for iron depletion, which also removes other cations (ie, calcium, magnesium, and zinc). Following this process, cations are added back to concentrations of calcium 20–25 mg/L, magnesium 10–12.5 mg/L, and zinc 0.5–1.0 mg/L.
- e. Testing this strain with this antimicrobial agent is considered supplemental QC only and is not required as routine user QC testing.
- f. QC limits for *E. coli* ATCC[®] 25922 with ciprofloxacin, nalidixic acid, minocycline, and sulfisoxazole when tested in CAMHB with 2.5% to 5% lysed horse blood incubated either in ambient air or 5% CO₂ (when testing *N. meningitidis*) are the same as those listed in Table 5A-1.
- g. When the erythromycin/clindamycin combination well for detecting inducible clindamycin resistance (ICR) is used, *S. aureus* ATCC® BAA-977™ (containing inducible *erm*A-mediated resistance) and *S. aureus* ATCC® 29213 or *S. aureus* ATCC® BAA-976™ (containing *msr*A-mediated macrolide-only efflux) are recommended for QC purposes. *S. aureus* ATCC® BAA-977™ should demonstrate ICR (ie, growth in the well), whereas *S. aureus* ATCC® 29213 and *S. aureus* ATCC® BAA-976™ should not demonstrate ICR (ie, no growth in the well).

Table 5A-1. (Continued)

- h. QC ranges reflect MICs obtained when CAMHB is supplemented with 0.002% polysorbate-80.
- QC ranges reflect MICs obtained when MHB is supplemented with calcium to a final concentration of 50 μg/mL. Agar dilution has not been validated for daptomycin.
- j. Exebacase QC ranges reflect MICs obtained when CAMHB is supplemented with 25% horse serum plus 0.5 mM DL-dithiothreitol (pH 7.2-7.4).
- k. The approved MIC susceptibility testing method is agar dilution. Agar media should be supplemented with 25 μg/mL of glucose-6-phosphate. Broth dilution should not be performed.
- I. For control organisms for gentamicin and streptomycin high-level aminoglycoside tests for enterococci, see Table 3J.
- m. QC range for S. aureus ATCC® 25923 with linezolid is 1–4 μg/mL; this strain exhibits less trailing, and MIC end points are easier to interpret. S. aureus ATCC® 25923 is considered a supplemental QC strain and is not required for routine QC of linezolid MIC tests.
- n. This test should be performed by agar dilution only.
- o. For broth microdilution testing of omadacycline and tigecycline, when MIC panels are prepared, the medium must be prepared fresh on the day of use. The medium must be no more than 12 hours old at the time the panels are made; however, the panels may then be frozen for later use.
- p. Very medium-dependent, especially with enterococci.
- q. QC range for *S. aureus* ATCC[®] 25923 with tedizolid is 0.12–0.5 μg/mL; this strain exhibits less trailing, and MIC end points are easier to interpret. *S. aureus* ATCC[®] 25923 is considered a supplemental QC strain and is not required for routine QC of tedizolid MIC tests.
- r. Ulifloxacin is the active metabolite of the prodrug prulifloxacin. Only ulifloxacin should be used for antimicrobial susceptibility testing.
- s. For QC organisms for vancomycin screen test for enterococci, see Table 3G.
- NOTE 1: These MICs were obtained in several referral laboratories by dilution methods. If four or fewer concentrations are tested, QC may be more difficult.
- **NOTE 2:** Information in boldface type is new or modified since the previous edition.

For Use With M07—MIC Testing

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Table 5A-2 Nonfastidious MIC QC for β-Lactam Combination Agents M07

Table 5A-2. MIC QC Ranges for Nonfastidious Organisms and β-Lactam Combination Agents^a

				QC Orq	anisms and	Characteristi	cs			
	Escherichia coli ATCC®b 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 29213	Enterococcus faecalis ATCC® 29212	Escherichia coli ATCC® 35218 ^{c,d}	Klebsiella pneumoniae ATCC 700603 ^{c,d}	Escherichia coli NCTC 13353 ^{c,d}	Klebsiella pneumoniae ATCC [®] BAA- 1705™ ^{c,d}	Klebsiella pneumoniae ATCC® BAA-2814™	Acinetobacter baumannii NCTC 13304 ^{c,d}
Antimicrobial	β-lactamase negative	Inducible Amp C	Weak β-lactamase <i>mecA</i> negative		TEM-1	SHV-18 OXA-2 Mutations in OmpK35 and OmpK37	CTX-M-15	KPC-2 TEM SHV	KPC-3 SHV-11 TEM-1	OXA-27
Agent					MIC QC Range					
Amoxicillin	_	_	-	-	-	> 128	-	_	_	_
Amoxicillin-clavulanate (2:1) ^e	2/1–8/4	-	0.12/0.06– 0.5/0.25	0.25/0.12– 1.0/0.5	4/2–16/8	4/2–16/8	_	_	_	_
Ampicillin	2–8	_	0.5–2	0.5–2	> 32	> 128	_	_	-	_
Ampicillin-sulbactam (2:1) ^e	2/1-8/4	ı	-	ı	8/4–32/16	8/4–32/16	-	-	_	ı
Aztreonam	0.06-0.25	2–8	_	-	0.03-0.12	8–64	-	_	-	_
Aztreonam-avibactam	0.03/4- 0.12/4	2/4–8/4	_	_	0.016/4- 0.06/4	0.06/4-0.5/4	_	_	_	_
Cefepime	0.016-0.12	0.5–4	1–4	_	0.008-0.06	0.5–2	≥64	_	_	16–128
Cefepime- enmetazobactam	0.03/8- 0.12/8	0.5/8–2/8	-	-	0.008/8- 0.06/8	0.12/8-0.5/8	0.03/8- 0.12/8	-	-	-
Cefepime- taniborbactam	0.03/4- 0.12/4	0.5/4-4/4	-	-	0.016/4- 0.06/4	0.12/4-0.5/4	0.12/4-1/4	0.12/4-0.5/4	-	-
Cefepime-tazobactam	0.03/8-0.12/8	0.5/8-4/8	1/8–4/8	-	-	0.12/8-0.5/8	0.06/8– 0.25/8	-	_	-
Cefepime-zidebactam (1:1)	0.016–0.06	0.5–2	_	-	_	0.06–0.25	0.06-0.5	_	_	4–16
Zidebactam ^f	0.06-0.25	1–8	-	-	-	-	0.06-0.5	-	-	≥128
Cefotaxime	0.03-0.12	8–32	1–4	_	_	-	-	_	-	-
Cefpodoxime	0.25–1	_	1–8	_	0.12-0.5	4–32	32–128	-	-	-
Ceftaroline	0.03-0.12	-	0.12-0.5	0.25–2	_	2–8	_	-	-	-
Ceftaroline-avibactam	0.03/4- 0.12/4	-	0.12/4-0.5/4	-	0.016/4- 0.06/4	0.25/4–1/4	-	-	_	-
Ceftazidime	0.06-0.5	1–4	4–16	_	_	16–64	-	_	-	_
Ceftazidime-avibactam	0.06/4-0.5/4	0.5/4-4/4	4/4–16/4	-	0.03/4- 0.12/4	0.25/4–2/4	=	-	_	_
Ceftolozane-tazobactam	0.12/4-0.5/4	0.25/4–1/4	16/4–64/4	-	0.06/4- 0.25/4	0.5/4–2/4	_	-	_	_
Ceftriaxone	0.03-0.12	8–64	1–8	-	-	_	-	_	-	_
Durlobactam	0.12-0.5	_	_	-	_	_	_	_	_	32–128

Table 5A-2. (Continued)

,	,			QC Org	anisms and	Characteristi	cs			
	Escherichia coli ATCC®b 25922	Pseudomonas aeruginosa ATCC® 27853	Staphylococcus aureus ATCC® 29213	Enterococcus faecalis ATCC® 29212	Escherichia coli ATCC [®] 35218 ^{c,d}	Klebsiella pneumoniae ATCC 700603 ^{c,d}	Escherichia coli NCTC 13353 ^{c,d}	Klebsiella pneumoniae ATCC [®] BAA- 1705™ ^{c,d}	Klebsiella pneumoniae ATCC® BAA-2814™	A. baumannii NCTC 13304 ^{c,d}
Antimicrobial	β-lactamase negative	Inducible Amp C	Weak β-lactamase <i>m</i> ecA negative		TEM-1	SHV-18 OXA-2 Mutations in OmpK35 and OmpK37	CTX-M-15	KPC-2 TEM SHV	KPC-3 SHV-11 TEM-1	OXA-27
Agent		r		1	MIC QC Range		T			
Imipenem	0.06-0.25	1–4	0.016–0.06	0.5–2	-	0.03-0.25	_	4–16	16–64	_
Imipenem- relebactam ^e	0.06/4– 0.25/4	0.25/4–1/4	0.008/4-0.03/4	0.5/4–2/4	0.06/4– 0.25/4	0.03/4- 0.25/4	_	0.03/4– 0.25/4	0.06/4 -0.5 /4	_
Meropenem	0.008-0.06	0.12-1	0.03-0.12	2–8	0.008-0.06	_	-	8–64	32-256	_
Meropenem- nacubactam (1:1)	0.015/0.015- 0.06/0.06	0.12/0.12–1/1	ı	-	ı	-	-	ı	0.5/0.5–2/2	ı
Meropenem- vaborbactam ^e	0.008/8- 0.06/8	0.12/8–1/8	0.03/8-0.12/8	-	0.008/8– 0.06/8	0.016/8– 0.06/8	_	0.008/8– 0.06/8	0.12/8-0.5/8	-
Nacubactam ^f	0.5–4	64–256	_	-	-	-	_	_	0.5–4	_
Piperacillin	1–4	1–8	1–4	1–4	>64	_	_	-	_	_
Piperacillin- tazobactam ^e	1/4-4/4	1/4-8/4	0.25/4–2/4	1/4-4/4	0.5/4–2/4	8/4-32/4	-	-	-	-
Sulbactam	16-64					32-128				16-64
Sulbactam- durlobactam	-	-	-	-		-	-	ı	-	0.5–2
Ticarcillin	4–16	8–32	2–8	16–64	> 128	> 256	_	_	_	_
Ticarcillin- clavulanate ^e	4/2–16/2	8/2–32/2	0.5/2–2/2	16/2–64/2	8/2–32/2	32/2-128/2	-	-	-	-

Abbreviations: ATCC®, American Type Culture Collection; MIC, minimal inhibitory concentration; NCTC, National Collection of Type Cultures; QC, quality control; R, resistant; S, susceptible.

QC strain selection codes:

QC strain is recommended for routine QC.

Test one of these agents by a disk diffusion or MIC method to confirm the integrity of the respective QC strain.c,d

Table 5A-2. (Continued)

Footnotes

- a. Unsupplemented Mueller-Hinton medium (cation-adjusted if broth). See Table 5A-1 for QC ranges for combination agents from other drug classes.
- b. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- c. Careful attention to organism maintenance (eg, minimal subcultures) and storage (eg, -60°C or below) is especially important for these QC strains because spontaneous loss of the plasmid encoding the β-lactamase has been documented. If stored at temperatures above -60°C or if repeatedly subcultured, these strains may lose their resistance characteristics and QC results may be outside the acceptable ranges.
- d. To confirm the integrity of the QC strain, test one of the single β-lactam agents highlighted in orange by either a disk diffusion or MIC test method when the strain is first subcultured from a frozen or lyophilized stock culture. In-range results for the single agent indicate the QC strain is reliable for QC of β-lactam combination agents. It is not necessary to check the QC strain again with a single agent until a new frozen or lyophilized stock culture is put into use, providing recommendations for handling QC strains as described in M02¹ and M07² are followed. If the highest concentration tested on a panel is lower than the QC range listed for the particular antimicrobial agent and the MIC result obtained for the QC strain is interpreted as resistant, the QC strain can be considered reliable for QC of β-lactam combination agents (eg, ampicillin panel concentrations 1–16 μg/mL; ampicillin **Enterobacterales** breakpoints [μg/mL]: ≤8 [S], 16 [I], ≥32 [R]; MIC of >16 μg/mI [R] would be acceptable for *K. pneumoniae* ATCC® 700603).
- e. Either strain highlighted in green may be used for routine QC of this antimicrobial agent.
- f. Not tested as a single agent routinely.

NOTE: Information in boldface type is new or modified since the previous edition.

References for Table 5A-2

- 1 CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

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Table 5B. MIC QC Ranges for Fastidious Organisms (Broth Dilution Methods)

	MIC QC Ranges, µg/mL						
Antimicrobial Agent	Haemophilus influenzae ATCC ^{®a} 49247	Haemophilus influenzae ATCC® 49766	Streptococcus pneumoniae ATCC® 49619				
Amikacin-fosfomycin (5:2) ^b	0.5/0.2-4/1.6	_	8/3.2–64/25.6				
Amoxicillin ^b	-	_	0.03-0.12				
Amoxicillin-clavulanate (2:1) ^c	2/1–16/8	_	0.03/0.016-0.12/0.06				
Ampicillin	2–8	_	0.06-0.25				
Ampicillin-sulbactam (2:1)	2/1–8/4	_	_				
Azithromycin	1–4	_	0.06-0.25				
Aztreonam	0.12-0.5	_	-				
Besifloxacin	0.016-0.06	_	0.03-0.12				
Cefaclor	-	1–4	1–4				
Cefamandole	_	0.25–1	_				
Cefdinir	-	0.12-0.5	0.03-0.25				
Cefditoren	0.06-0.25	_	0.016–0.12				
Cefepime	0.5–2	_	0.03-0.25				
Cefepime-tazobactam	0.5/8–2/8	_	0.03/8-0.12/8				
Cefetamet	0.5–2	_	0.5–2				
Cefixime	0.12–1	_	<u> </u>				
Cefmetazole	2–16	_					
Cefonicid	-	0.06-0.25	-				
Cefotaxime	0.12–0.5	_	0.03-0.12				
Cefotetan	-	_	-				
Cefoxitin	_	_	-				
Cefpirome	0.25–1	_	-				
Cefpodoxime	0.25–1	_	0.03-0.12				
Cefprozil	-	1–4	0.25–1				
Ceftaroline	0.03-0.12	_	0.008-0.03				
Ceftaroline-avibactam	0.016/4-0.12/4	_					
Ceftazidime	0.12–1	-	-				
Ceftazidime-avibactam ^d	0.06/4-0.5/4	0.016/4-0.06/4	0.25/4–2/4				
Ceftibuten	0.25–1	_	-				
Ceftizoxime	0.06-0.5	_	0.12–0.5				
Ceftobiprole ^e	0.12–1	0.016–0.06	0.004–0.03				
Ceftolozane-tazobactam	0.5/4–2/4	_	0.25/4–1/4				
Ceftriaxone	0.06-0.25	_	0.03-0.12				
Cefuroxime	-	0.25–1	0.25–1				
Cephalothin	-	_	0.5–2				
Chloramphenicol	0.25–1	_	2–8				
Ciprofloxacin ^f	0.004–0.03	_	<u> </u>				
Clarithromycin	4–16	_	0.03-0.12				
Clinafloxacin	0.001-0.008	_	0.03-0.12				
Clindamycin	_	_	0.03-0.12				
Dalbavancin ^g	_	_	0.008-0.03				

Table 5B. (Continued)

Table 3B. (Continued)	MIC QC Ranges, μg/mL							
Antimicrobial Agent	Haemophilus influenzae ATCC ^{®a} 49247	Haemophilus influenzae ATCC® 49766	Streptococcus pneumoniae ATCC® 49619					
Daptomycin ^h	-	-	0.06-0.5					
Delafloxacin	0.00025-0.001	_	0.004-0.016					
Dirithromycin	8–32	_	0.06-0.25					
Doripenem	_	0.06-0.25	0.03-0.12					
Doxycycline	_	_	0.016-0.12					
Enoxacin	_	-	-					
Eravacycline	0.06–0.5	-	0.004-0.03					
Ertapenem	_	0.016-0.06	0.03-0.25					
Erythromycin	_	_	0.03-0.12					
Faropenem	_	0.12–0.5	0.03-0.25					
Finafloxacin	_	0.002-0.008	0.25–1					
Fleroxacin	0.03-0.12	-	-					
Fusidic acid	_	-	4–32					
Garenoxacin	0.002-0.008	ı	0.016-0.06					
Gatifloxacin	0.004-0.03	ı	0.12-0.5					
Gemifloxacin	0.002-0.008	ı	0.008-0.03					
Gentamicin	_	ı	-					
Gepotidacin	0.25–1	1	0.06-0.25					
Grepafloxacin	0.002-0.015	1	0.06-0.5					
claprim	0.12–1	1	0.03-0.12					
mipenem	_	0.25–1	0.03-0.12					
mipenem-relebactam	_	0.25/4-1/4	0.016/4-0.12/4					
Lefamulin	0.5–2	-	0.06-0.5					
Levofloxacin	0.008-0.03	-	0.5–2					
_evonadifloxacin	0.008-0.06	-	0.12–0.5					
Linezolid	_	-	0.25–2					
omefloxacin	0.03-0.12	_	_					
_oracarbef	_	0.5–2	2–8					
Meropenem	_	0.03-0.12	0.03-0.25					
Metronidazole	-	-	_					
Minocycline ^f	_	-	-					
Moxifloxacin	0.008-0.03	_	0.06-0.25					
Nafithromycin	2–8	_	0.008-0.03					
Nalidixic acid ^f	_	_	-					
Nitrofurantoin	_	_	4–16					
Norfloxacin	_	_	2–8					
Ofloxacin	0.016-0.06	_	1–4					
Omadacycline ⁱ	0.5–2	_	0.016–0.12					

Table 5B. (Continued)

	MIC QC Ranges, µg/mL						
Antimicrobial Agent	Haemophilus influenzae ATCC ^{®a} 49247	Haemophilus influenzae ATCC® 49766	Streptococcus pneumoniae ATCC® 49619				
Oritavancin ^g	_	-	0.001-0.004				
Ozenoxacin	-	-	0.008-0.06				
Penicillin	_	-	0.25–1				
Pexiganan	8–32	-	16–64				
Piperacillin-tazobactam	0.06/4-0.5/4	-	-				
Quinupristin-dalfopristin	2–8	-	0.25–1				
Razupenem	_	0.008-0.03	0.008-0.06				
Rifampin	0.25–1	-	0.016-0.06				
Solithromycin	1–4	-	0.004-0.016				
Sparfloxacin	0.004-0.016	-	0.12-0.5				
Spectinomycin	_	-	-				
Sulfisoxazole ^f	_	_	-				
Sulopenem	_	0.06-0.25	0.03-0.12				
Tedizolid	_	-	0.12–0.5				
Telavancin ^g	_	-	0.004–0.016				
Telithromycin	1–4	_	0.004-0.03				
Tetracycline	4–32	_	0.06-0.5				
Tigecycline ⁱ	0.06-0.5	-	0.016-0.12				
Trimethoprim-sulfamethoxazole (1:19)	0.03/0.59-0.25/4.75	_	0.12/2.4-1/19				
Trospectomycin	0.5–2	_	1–4				
Trovafloxacin	0.004-0.016	_	0.06-0.25				
Vancomycin	_	-	0.12-0.5				
Zoliflodacin	0.12–1	-	0.12-0.5				

MIC Testing Conditions for Clinical Isolates and Performance of QC

		Streptococcus	
Organism	Haemophilus influenzae	pneumoniae and streptococci	Neisseria meningitidis
Medium	Broth dilution: HTM broth	Broth dilution: CAMHB with LHB	Broth dilution: CAMHB with LHB
		(2.5% to 5% v/v)	(2.5% to 5% v/v)
Inoculum	Colony suspension	Colony suspension	Colony suspension
Incubation	Ambient air; 20–24 hours; 35°C	Ambient air; 20–24 hours; 35°C	5% CO ₂ ; 20–24 hours; 35°C
characteristics			
			(for QC with S. pneumoniae ATCC® 49619,
			5% CO ₂ or ambient air, except for
			azithromycin, ambient air only)

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; HTM, *Haemophilus* test medium; LHB, lysed horse blood; MIC, minimal inhibitory concentration; QC, quality control.

Table 5B. (Continued)

Footnotes

- a. ATCC® is a registered trademark of the American Type Culture Collection.
- b. QC ranges reflect MICs obtained when medium is supplemented with 25 μg/mL of glucose-6-phosphate.
- c. QC limits for *E. coli* ATCC[®] 35218 when tested on HTM are 4/2–16/8 μg/mL for amoxicillin-clavulanate and ≥ 256 μg/mL for amoxicillin; testing amoxicillin may help to determine if the isolate has maintained its ability to produce β-lactamase.
- d. QC limits for *K. pneumoniae* ATCC® 700603 with ceftazidime-avibactam when testing in HTM are 0.25/4–1/4 μg/mL. *K. pneumoniae* ATCC® 700603 should be tested against ceftazidime-avibactam and ceftazidime alone to confirm the activity of avibactam in the combination and to ensure that the plasmid encoding the β-lactamase has not been lost in this strain. The acceptable range for ceftazidime alone is > 16 μg/mL.
- e. Either H. influenzae ATCC® 49247 or 49766 may be used for routine QC testing.
- f. QC limits for *E. coli* ATCC® 25922 with ciprofloxacin, nalidixic acid, minocycline, and sulfisoxazole when tested in CAMHB with 2.5% to 5% LHB incubated either in ambient air or 5% CO₂ (when testing *N. meningitidis*) are the same as those listed in Table 5A-1.
- g. QC ranges reflect MICs obtained when CAMHB is supplemented with 0.002% polysorbate-80.
- h. QC ranges reflect MICs obtained when Mueller-Hinton broth is supplemented with calcium to a final concentration of 50 μg/mL. Agar dilution has not been validated for daptomycin.
- i. For broth microdilution testing of omadacycline and tigecycline, when MIC panels are prepared, the medium must be prepared fresh on the day of use. The medium must be no more than 12 hours old at the time the panels are made; however, the panels may then be frozen for later use.

NOTE 1: For four-dilution ranges, results at the extremes of the acceptable ranges should be suspect. Verify validity with data from other QC strains.

NOTE 2: Information in boldface type is new or modified since the previous edition.

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Table 5C. MIC QC Ranges for Neisseria gonorrhoeae (Agar Dilution Method) MIC QC Ranges, µg/mL Neisseria gonorrhoeae ATCC®a 49226 **Antimicrobial Agent** Azithromycin 0.25-1 0.008-0.03 Cefdinir 0.016-0.06 Cefepime Cefetamet 0.016-0.25 Cefixime 0.004-0.03 Cefmetazole 0.5-2 Cefotaxime 0.016-0.06 Cefotetan 0.5-2 0.5-2 Cefoxitin Cefpodoxime 0.03 - 0.12Ceftazidime 0.03-0.12 0.008-0.03 Ceftizoxime Ceftriaxone 0.004-0.016 Cefuroxime 0.25 - 10.001-0.008 Ciprofloxacin Enoxacin 0.016-0.06 Fleroxacin 0.008-0.03 0.002-0.016 Gatifloxacin 0.25-1 Gepotidacin Grepafloxacin 0.004-0.03 0.008-0.03 Lomefloxacin 0.008-0.03 Moxifloxacin Ofloxacin 0.004-0.016 Penicillin 0.25-1 Solithromycin 0.03-0.25 Sparfloxacin 0.004-0.016 8-32 Spectinomycin Tetracycline 0.25 - 1Trospectomycin 1–4 0.004-0.016 Trovafloxacin

0.06-0.5

Table 5C. (Continued)

Testing Conditions for Clinical Isolates and Performance of QC

Organism	Neisseria gonorrhoeae
Medium	Agar dilution: GC agar base and 1% defined growth supplement. The use of a cysteine-free supplement is necessary for agar dilution tests with carbapenems and clavulanate. Cysteine-containing defined growth supplements do not significantly alter dilution test results with other drugs.
Inoculum	Colony suspension, equivalent to a 0.5 McFarland standard
Incubation characteristics	36°C ± 1°C (do not exceed 37°C); 5% CO ₂ ; 20–24 hours

Abbreviations: ATCC®, American Type Culture Collection; MIC, minimal inhibitory concentration; QC; quality control.

Footnote

a. $ATCC^{\otimes}$ is a registered trademark of the American Type Culture Collection.

NOTE: Information in boldface type is new or modified since the previous edition.

Table 5D. MIC QC Ranges for Anaerobes (Agar Dilution Method)

	MIC QC Ranges, μg/mL						
Antimicrobial Agent	Bacteroides fragilis ATCC®a 25285	Bacteroides thetaiotaomicron ATCC® 29741	Clostridioides (formerly Clostridium) difficile ATCC® 700057	Eggerthella lenta (formerly Eubacterium lentum) ATCC® 43055 ^b			
Amoxicillin-clavulanate (2:1)	0.25/0.125-1/0.5	0.5/0.25–2/1	0.25/0.125-1/0.5	_			
Ampicillin	16–64	16–64	1–4	_			
Ampicillin-sulbactam (2:1)	0.5/0.25–2/1	0.5/0.25–2/1	0.5/0.25-4/2	0.25/0.125–2/1			
Cadazolid	_	_	0.12-0.5	_			
Cefmetazole	8–32	32–128	_	4–16			
Cefoperazone	32–128	32–128	-	32–128			
Cefotaxime	8–32	16–64	_	64–256			
Cefotetan	4–16	32–128	-	32–128			
Cefoxitin	4–16	8–32	_	4–16			
Ceftaroline	4–32	16–128	2–16	8–32			
Ceftaroline-avibactam	0.12/4-0.5/4	4/4–16/4	0.5/4-4/4	4/4–16/4			
Ceftizoxime	_	4–16	_	16–64			
Ceftolozane-tazobactam	0.12/4-1/4	16/4–128/4	_	_			
Ceftriaxone	32–128	64–256	_	_			
Chloramphenicol	2–8	4–16	_	_			
Clinafloxacin	0.03-0.125	0.06-0.5	_	0.03-0.125			
Clindamycin	0.5–2	2–8	2–8	0.06-0.25			
Doripenem	_	_	0.5–4	_			
Eravacycline	0.06-0.25	0.12–1	0.06–0.25	_			
Ertapenem	0.06–0.25	0.25–1	-	0.5–2			
Faropenem	0.03-0.25	0.12–1		1–4			
Fidaxomicin	-	-	0.06-0.25	_			
Finafloxacin	0.12-0.5	1–4	1–4	0.12–0.5			
Garenoxacin	0.06-0.5	0.25–1	0.5–2	1–4			
_	0.03-0.125	0.25-1	0.5–2	0.125-0.5			
mipenem							
mipenem-relebactam	0.03/4-0.25/4	0.06/4-0.5/4		0.12/4-1/4			
Linezolid	2–8	2–8	1–4	0.5–2			
Meropenem	0.03-0.25	0.125-0.5	0.5–4	0.125–1			
Metronidazole	0.25–1	0.5–2	0.125-0.5	-			
Moxifloxacin	0.125–0.5	1–4	1–4	0.125–0.5			
Nitazoxanide	 0.25–2	0.5–4	0.06–0.5 0.25–2	0.25–2			
Omadacycline Penicillin	0.25–2 8–32	8–32	0.25–2 1–4	0.25-2			
Piperacillin	2–8	8–32	4–16	8–32			
Piperacillin-tazobactam	0.125/4-0.5/4	4/4–16/4	4/4–16/4	4/4–16/4			

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Table 5D. (Continued)

Antimicrobial Agent	Bacteroides fragilis ATCC®a 25285	Bacteroides thetaiotaomicron ATCC® 29741	Clostridioides (formerly Clostridium) difficile ATCC® 700057	Eggerthella lenta (formerly Eubacterium lentum) ATCC® 43055 ^b
Ramoplanin	_	_	0.125–0.5	_
Razupenem	0.016–0.12	0.06-0.25	0.06-0.25	0.06-0.5
Ridinilazole	_	_	0.06-0.25	_
Rifaximin	_	_	0.004-0.016	-
Secnidazole	0.25–1	0.5–2	0.06–0.5	0.25–2
Sulopenem	-	0.06-0.5	1–4	0.5–2
Surotomycin ^c	-	-	0.12–1	2–8
Tetracycline	0.125–0.5	8–32	-	-
Ticarcillin	16–64	16–64	16–64	16–64
Ticarcillin-clavulanate	_	0.5/2–2/2	16/2–64/2	16/2–64/2
Tigecycline	0.12–1	0.5–2	0.125–1	0.06-0.5
Tinidazole	_	-	0.125–0.5	-
Tizoxanide	_	_	0.06–0.5	-
Vancomycin		_	0.5–4	_

Abbreviations: ATCC®, American Type Culture Collection; MIC, minimal inhibitory concentration; QC, quality control.

Footnotes

- a. ATCC® is a registered trademark of the American Type Culture Collection.
- b. MIC variability with some agents has been reported with *Eggerthella lenta* (formerly *E. lentum*) ATCC® 43055; therefore, QC ranges have not been established for all antimicrobial agents with this organism.
- c. QC ranges reflect MICs obtained when media are supplemented with calcium to a final concentration of 50 µg/mL.

Table 5E. MIC QC Ranges for Anaerobes (Broth Microdilution Method)

	MIC QC Ranges, µg/mL							
Antimicrobial Agent	Bacteroides fragilis ATCC ^{®a} 25285	Bacteroides thetaiotaomicron ATCC® 29741	Clostridioides (formerly Clostridium) difficile ATCC® 700057	Eggerthella lenta (formerly Eubacterium lentum) ATCC® 43055 ⁵				
Amoxicillin-clavulanate (2:1)	0.25/0.125-1/0.5	0.25/0.125-1/0.5	_	_				
Ampicillin-sulbactam (2:1)	0.5/0.25–2/1	0.5/0.25–2/1	_	0.5/0.25-2/1				
Cadazolid	_	_	0.06-0.25	_				
Cefotetan	1–8	16–128	_	16–64				
Cefoxitin	2–8	8–64	_	2–16				
Ceftaroline	2–16	8–64	0.5–4	_				
Ceftaroline-avibactam	0.06/4-0.5/4	2/4-8/4	0.25/4-1/4	4/4–16/4				
Ceftizoxime	_	-	_	8–32				
Ceftolozane-tazobactam	0.12/4-1/4	16/4–64/4	-	_				
Chloramphenicol	4–16	8–32	_	4–16				
Clindamycin	0.5–2	2–8	_	0.06-0.25				
Doripenem	0.12-0.5	0.12–1	_	_				
Doxycycline	_	2–8	_	2–16				
Eravacycline	0.016-0.12	0.06-0.25	0.016-0.06	_				
Ertapenem	0.06–0.5	0.5–2	_	0.5–4				
Faropenem	0.016-0.06	0.12–1	_	0.5–2				
Garenoxacin	0.06-0.25	0.25–2	_	0.5–2				
Imipenem	0.03-0.25	0.25–1	_	0.25–2				
Imipenem-relebactam	0.03/4-0.125/4	_	_	_				
Linezolid	2–8	2–8	_	0.5–2				
Meropenem	0.03-0.25	0.06-0.5	_	0.125–1				
Metronidazole	0.25–2	0.5–4	_	0.125–0.5				
Moxifloxacin	0.12-0.5	1.0–8	_	0.12-0.5				
Omadacycline ^c	0.12–1	0.25–1	0.06-0.25	0.06–5				
Penicillin	8–32	8–32	_	_				
Piperacillin	4–16	8–64	_	8–32				
Piperacillin-tazobactam	0.03/4-0.25/4	2/4-16/4	_	8/4-32/4				
Razupenem	0.03-0.25	0.12-0.5	0.06-0.5	0.12-0.5				
Ridinilazole	_		0.12-0.5	-				
Sulopenem	-	0.03-0.25	0.5–2	0.25–1				
Surotomycin ^d	_	_	0.12–1	1–4				
Ticarcillin-clavulanate	0.06/2-0.5/2	0.5/2-2/2	_	8/2-32/2				
Tigecycline ^c	0.06-0.5	0.25–1	0.03-0.12	_				

Abbreviations: ATCC®, American Type Culture Collection; MIC, minimal inhibitory concentration; QC, quality control.

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Table 5E. (Continued)

Footnotes

- a. ATCC® is a registered trademark of the American Type Culture Collection.
- b. MIC variability with some agents has been reported with *Eggerthella lenta* (formerly *E. lentum*) ATCC® 43055; therefore, QC ranges have not been established for all antimicrobial agents with this organism.
- c. For broth microdilution testing of omadacycline and tigecycline, when MIC panels are prepared, the medium must be prepared fresh on the day of use. The medium must be no greater than 12 hours old at the time the panels are made; however, the panels may then be frozen for later use.
- d. QC ranges reflect MICs obtained when broth is supplemented with calcium to a final concentration of 50 μg/mL.

NOTE: For four-dilution ranges, results at the extremes of the acceptable range(s) should be suspect. Verify validity with data from other QC strains.

Table 5F. MIC Reference Guide to QC Frequency

This table summarizes the suggested QC frequency when modifications are made to antimicrobial susceptibility test systems (refer to CLSI documents EP23¹ and M52²). It applies only to antimicrobial agents for which satisfactory results have been obtained with either the 15-replicate (3- × 5-day) plan or 20 or 30 consecutive test day plan. Otherwise QC is required each test day.

	Recommended QC Frequency			
Test Modification	1 Day	5 Days	15-Replicate Plan or 20- or 30-Day Plan	Comments
MIC test(s)				
Use new shipment or lot number.	X			
Expand dilution range.	Х			Example: Convert from breakpoint to expanded range MIC panels.
Reduce dilution range.	Х			Example: Convert from expanded dilution range to breakpoint panels.
Use new method (same company).			Х	Examples: Convert from overnight to rapid MIC test.
Use new manufacturer of MIC test.			X	In addition, perform in-house verification studies. In addition, perform in-house verification studies.
Use new manufacturer of broth or agar.		X	^	in addition, perform in-nouse verification studies.
Addition of new antimicrobial agent to existing system		, , , , , , , , , , , , , , , , , , ,	X	In addition, perform in-house verification studies.
Inoculum preparation				
Convert inoculum preparation/standardization to use of a device that has its own QC protocol.		х		Example: Convert from visual adjustment of turbidity to use of a photometric device for which a QC procedure is provided.
Convert inoculum preparation/standardization to a method that depends on user technique.			Х	Example: Convert from visual adjustment of turbidity to another method that is not based on a photometric device.
Instrument/software				
Software update that affects AST results		х		Monitor all drugs, not just those implicated in software modification.
Repair of instrument that affects AST results Abbreviations: AST, antimicrobial susceptibility to	Х			Depending on extent of repair (eg, critical component such as the photographic device), additional testing may be appropriate (eg, 5 days).

Abbreviations: AST, antimicrobial susceptibility testing; MIC, minimal inhibitory concentration; QC, quality control.

Table 5F. (Continued)

NOTE 1: QC can be performed before or concurrent with testing patient isolates. Patient results can be reported for that day if QC results are within the acceptable limits.

NOTE 2: Manufacturers of commercial or in-house-prepared tests should follow their own internal procedures and applicable regulations.

NOTE 3: Acceptable MIC QC limits for US Food and Drug Administration—cleared antimicrobial susceptibility tests may differ slightly from acceptable CLSI QC limits. Users of each device should use the manufacturer's procedures and QC limits as indicated in the instructions for use.

NOTE 4: For troubleshooting out-of-range results, refer to M07,³ Subchapter 4.8 and M100 Table 5G. Additional information is available in Appendix C (eg, organism characteristics, QC testing recommendations).

NOTE 5: Broth, saline, and/or water used to prepare an inoculum does not need routine QC.

References for Table 5F

- 1 CLSI. Laboratory Quality Control Based on Risk Management; Approved Guideline. CLSI document EP23-A™. Wayne, PA: Clinical and Laboratory Standards Institute; 2011.
- ² CLSI. *Verification of Commercial Microbial Identification and Antimicrobial Susceptibility Testing Systems*. 1st ed. CLSI guideline M52. Wayne, PA: Clinical and Laboratory Standards Institute; 2015.
- ³ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

Table 5G. MIC Troubleshooting Guide

This table provides guidance for troubleshooting and corrective action for out-of-range QC, primarily using CAMHB for broth microdilution. Refer to M07,¹ Chapter 4, for additional information. Out-of-range QC tests are often the result of contamination or the use of an incorrect QC strain; corrective action should first include repeating the test with a pure culture of a freshly subcultured QC strain. If the issue is unresolved, this troubleshooting guide should be consulted regarding additional suggestions for troubleshooting out-of-range QC results and unusual clinical isolate results. In addition, see general corrective action outlined in M07¹ and notify manufacturers of potential product problems.

General Comment

(1) QC organism maintenance: Avoid repeated subcultures. Retrieve new QC strain from stock (refer to M07,¹ Subchapter 4.4). If using lyophilized strains, follow the maintenance recommendations of the manufacturer.

Antimicrobial Agent	QC Strain	Observation	Probable Cause	Comments/Suggested Actions
β-LACTAMS				
β-lactam combination agents	A. baumannii ATCC®a 13304 E. coli ATCC® 35218 E. coli ATCC® 13353 K. pneumoniae ATCC® 700603 K. pneumoniae ATCC® BAA-1705™	MIC too low or susceptible for single β-lactam agent; in range for combination β-lactam agent	Spontaneous loss of the plasmid encoding the β-lactamase	Obtain new frozen or lyophilized stock culture. Use other routine QC strain (if available). These strains should be stored at −60°C or below, and frequent subcultures should be avoided. NOTE: K. pneumoniae ATCC® BAA-2814™ is stable and does not require QC integrity check.
β-lactam combination agents	A. baumannii ATCC® 13304 E. coli ATCC® 35218 E. coli ATCC® 13353 K. pneumoniae ATCC® 700603 K. pneumoniae ATCC® BAA-1705™ K. pneumoniae ATCC® BAA-2814™	MIC too high or resistant for both the single β-lactam agent and the combination β-lactam agent	Antimicrobial agent is degrading.	Use alternative lot of test materials. Check storage and package integrity. Imipenem and clavulanate are especially labile.
Carbenicillin	P. aeruginosa ATCC® 27853	MIC too high	QC strain develops resistance after repeated subculture.	See general comment (1) on QC organism maintenance.
Cefotaxime-clavulanate Ceftazidime-clavulanate	K. pneumoniae ATCC® 700603	Negative ESBL test	Spontaneous loss of the plasmid encoding the β-lactamase	See general comment (1) on QC organism maintenance.
Carbapenems	P. aeruginosa ATCC® 27853	MIC too high	Zn++ concentration in media is too high.	Use alternative lot.
Carbapenems	P. aeruginosa ATCC® 27853	MIC too high	Antimicrobial agent is degrading.	Use alternative lot. Check storage conditions and package integrity. Repeated imipenem QC results at the upper end of QC range with <i>P. aeruginosa</i> ATCC® 27853 may indicate deterioration of the drug.
Penicillin	S. aureus ATCC® 29213	MIC too high	QC strain is a β-lactamase producer; overinoculation may yield increased MICs.	Repeat with a carefully adjusted inoculum.

Table 5G. (Continued)

Any			
Anv			
,	MIC too low	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO ₂ incubation, which lowers pH.
Any	MIC too high	pH of media too high	Acceptable pH range = 7.2–7.4
Any	MIC initially acceptable, but increases to possibly be out of range over time	Imipenem, cefaclor, and clavulanate are especially labile. Antimicrobial agents are degrading.	Use alternative lot. Check storage and package integrity.
		, 5	
Any	MIC too high	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO₂ incubation, which lowers pH.
Any	MIC too low	pH of media too high	Acceptable pH range = 7.2-7.4
P. aeruginosa ATCC® 27853	MIC too low	Ca++ and/or Mg++ content too low	Acceptable range = Ca++ 20–25 mg/L Mg++ 10–12.5 mg/L
P. aeruginosa ATCC® 27853	MIC too low	Ca++ and/or Mg++ content too low	Acceptable range = Ca++ 20–25 mg/L Mg++ 10–12.5 mg/L
S. aureus ATCC® 29213 E. faecalis ATCC® 29212	MIC too high	Lack of polysorbate-80 in the media	Add polysorbate-80 to CAMHB to final concentration of 0.002% (v/v). See M07, ¹ Subchapter 3.5.1 and Appendix A.
S. aureus ATCC® 29213 E. faecalis ATCC® 29212 S. pneumoniae ATCC® 49619	MIC too high	Trailing end point	Read at first well where the trailing begins; tiny buttons of growth should be ignored. See general comment (2) in Table 2G.
S. aureus ATCC® 29213	MIC too high	Trailing end point	S. aureus ATCC [®] 25923 may be used as a supplementa QC strain for these drugs. This strain exhibits less trailing and MIC end points are easier to interpret.
S. aureus ATCC® 29213 E. faecalis ATCC® 29212	MIC too high	Lack of polysorbate-80 in the solvent and diluent	Dissolve antimicrobial powder and prepare dilutions in water containing a final concentration of 0.002% polysorbate-80 (v/v).
S. aureus ATCC® 29213 E. faecalis ATCC® 29212	MIC too high	Use of tissue-culture treated microdilution trays	Only use untreated microdilution trays for this antimicrobial agent. ²
S. aureus ATCC® 29213 E. faecalis ATCC® 29212	MIC too high	pH of media too low	Acceptable pH range = 7.2–7.4 Avoid CO₂ incubation, which lowers pH.
S. aureus ATCC® 29213 E. faecalis ATCC® 29212	MIC too low	pH of media too high	Acceptable pH range = 7.2–7.4
S. aureus ATCC® 29213 E. faecalis ATCC® 29212	MICs too low	Ca++ content too low	Acceptable Ca++ content 50 μg/mL in CAMHB
	Any Any P. aeruginosa ATCC® 27853 P. aeruginosa ATCC® 27853 S. aureus ATCC® 29213 E. faecalis ATCC® 29212 S. aureus ATCC® 29212 S. pneumoniae ATCC® 49619 S. aureus ATCC® 29213 E. faecalis ATCC® 29213 E. faecalis ATCC® 29212 S. aureus ATCC® 29213 E. faecalis ATCC® 29212 S. aureus ATCC® 29212 S. aureus ATCC® 29213 E. faecalis ATCC® 29213 E. faecalis ATCC® 29212 S. aureus ATCC® 29213 E. faecalis ATCC® 29213 E. faecalis ATCC® 29213	Any MIC initially acceptable, but increases to possibly be out of range over time MIC too high Any MIC too low P. aeruginosa ATCC®	Any MIC too high pH of media too low Any MIC too low pH of media too high P. aeruginosa ATCC® MIC too low low 27853 MIC too low 27853 MIC too high Lack of polysorbate-80 in the media E. aureus ATCC® 29213 MIC too high S. aureus ATCC® 29212 MIC too high Lack of polysorbate-80 in the solvent and diluent S. aureus ATCC® 29213 MIC too high Lack of polysorbate-80 in the solvent and diluent S. aureus ATCC® 29213 MIC too high Lack of polysorbate-80 in the media S. aureus ATCC® 29213 MIC too high Lack of polysorbate-80 in the media S. aureus ATCC® 29213 MIC too high Lack of polysorbate-80 in the solvent and diluent S. aureus ATCC® 29213 MIC too high Lack of polysorbate-80 in the solvent and diluent S. aureus ATCC® 29213 MIC too high Lack of polysorbate-80 in the solvent and diluent S. aureus ATCC® 29213 MIC too high Deciding MIC too high Decidin

Table 5G. (Continued) **Antimicrobial Agent** QC Strain Observation **Probable Cause Comments/Suggested Actions** NON-β-LACTAMS (Continued) Tetracyclines Any MIC too low pH of media too low Acceptable pH range = 7.2-7.4 Acceptable pH range = 7.2-7.4 Tetracyclines Any MIC too high pH of media too high Avoid CO₂ incubation, which lowers pH. Tetracyclines MIC too high Ca++ and/or Mg++ content too Acceptable range = Ca++20-25 mg/L Mg++10-12.5Any MIC too low Ca++ and/or Mg++ content too Acceptable range = Ca++ 20-25 mg/L Mg++ 10-12.5 Tetracyclines Any CAMHB has not been freshly Reference panels must be used or frozen within Omadacycline MIC too high Any Tigecycline prepared. 12 hours of CAMHB preparation. **ALL AGENTS** E. coli ATCC® 35218 Various MIC too low Spontaneous loss of the See general comment (1) on QC organism plasmid encoding the maintenance. K. pneumoniae ATCC® β-lactamase 700603 Various Any One QC result is out of N/A If antimicrobial agent is not normally reported, no repeat is necessary if adequate controls are in place to prevent range, but the reporting of the out-of-range antimicrobial agent. antimicrobial agent is not an agent reported for patient results (eg, not on hospital formulary). Repeat using McFarland 0.5 turbidity standard or Various Any Many MICs too low Inoculum too light; error in inoculum preparation standardizing device. Check expiration date and proper storage if using barium sulfate or latex standards. Check steps in inoculum preparation and inoculation procedure. Perform colony count check of growth control well immediately after inoculation and before incubation (E. coli ATCC® 25922 closely approximates 5×10⁵ CFU/mL; see M07, Subchapter 3.8). Many MICs too high or too CAMHB not optimal Use alternative lot. Various Any Many MICs too high or too Possible reading/transcription Various Any Recheck readings. low error Use alternative lot. Many MICs too high Repeat using McFarland 0.5 turbidity standard or Various Any Inoculum too heavy standardizing device. Check expiration date and proper storage if using barium sulfate or latex standards. Check steps in inoculum preparation and inoculation procedure. Perform colony count check of growth control well immediately after inoculation and before incubation (E. coli ATCC® 25922 closely approximates 5 ×10⁵ CFU/mL; see M07, 1 Subchapter 3.8).

Antimicrobial Agent	QC Strain	Observation	Probable Cause	Comments/Suggested Actions
ALL AGENTS (Continue	d)			
Various	Any	Skipped wells	Contamination. Improper inoculation of panel or inadequate mixing of inoculum. Actual concentration of drug in wells inaccurate. Volume of broth in wells inaccurate.	Repeat QC test. Use alternative lot.
Various	Any	QC results from one strain are out of range, but other QC strains are in range with the same antimicrobial agent.	One QC organism may be a better indicator of a QC problem (eg, <i>P. aeruginosa</i> ATCC® 27853 is a better indicator of imipenem deterioration than <i>E. coli</i> ATCC® 25922).	Determine if the in-range QC strain has an on-scale end point for the agent in question. Retest this strain to confirm reproducibility of acceptable results. Evaluate with alternative strains with known MICs. Initiate corrective action with problem QC strain/antimicrobial agent(s).
Various	Any	QC results from two strains are out of range with the same antimicrobial agent.	Indicates a problem with the antimicrobial agent. May be a systemic problem.	Initiate corrective action.
Various	Any	QC results from one strain are out of range, but the antimicrobial agent is not an agent reported for patient results (eg, not on hospital formulary).		If antimicrobial agent is not normally reported, no repeat is necessary if adequate controls are in place to prevent reporting of the out-of-range antimicrobial agent. Carefully check antimicrobial agents of the same class for similar trend toward out-of-control results. If the antimicrobial agent in question is consistently out of control, contact the manufacturer.
Various	E. coli ATCC® 35218 K. pneumoniae ATCC® 700603	MIC too low	Spontaneous loss of the plasmid encoding the β-lactamase	See general comment (1) on QC organism maintenance.

Abbreviations: ATCC®, American Type Culture Collection; CAMHB, cation-adjusted Mueller-Hinton broth; CFU, colony-forming unit(s); ESBL, extended-spectrum β-lactamase; MIC, minimal inhibitory concentration; N/A, not applicable; pH, negative logarithm of hydrogen ion concentration; QC, quality control.

Footnote

a. ATCC® is a trademark of the American Type Culture Collection.

References for Table 5G

- ¹ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ² Arhin FF, Sarmiento I, Belley A, et al. Effect of polysorbate 80 on oritavancin binding to plastic surfaces: implications for susceptibility testing. *Antimicrob Agents Chemother*. 2008;52(5):1597-1603.

Table 6A. Solvents and Diluents for Preparing Stock Solutions of Antimicrobial Agents^a

Antimicrobial Agent	Solvent ^b	Diluent ^b
	Unless otherwise stated, use a minimum amount of the listed solvent to solubilize the antimicrobial powder.	Finish diluting the final stock solution as stated below.
Amikacin	Water	Water
Amoxicillin	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L
Ampicillin	Phosphate buffer, pH 8, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L
Avibactam	Water	Water
Azithromycin	95% ethanol or glacial acetic acid ^{a,c}	Broth media
Azlocillin	Water	Water
Aztreonam	Saturated solution sodium bicarbonate	Water
Besifloxacin	Methanol	Water
Biapenem	Saline ^d	Saline ^d
Cadazolid	DMSO ^a	Water or broth
Carbenicillin	Water	Water
Cefaclor	Water	Water
Cefadroxil	Phosphate buffer, pH 6, 0.1 mol/L	Water
Cefamandole	Water	Water
Cefazolin	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L
Cefdinir	Phosphate buffer, pH 6, 0.1 mol/L	Water
Cefditoren	Phosphate buffer, pH 6, 0.1 mol/L	Water
Cefepime	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L or water
Cefetamet	Phosphate buffer, pH 6, 0.1 mol/L	Water
Cefiderocol	Saline ^d	Saline ^d
Cefixime	Phosphate buffer, pH 7, 0.1 mol/L	Phosphate buffer, pH 7, 0.1 mol/L
Cefmetazole	Water	Water
Cefonicid	Water	Water
Cefoperazone	Water	Water
Cefotaxime	Water	Water
Cefotetan	DMSO ^a	Water
Cefoxitin	Water	Water
Cefpodoxime	0.10% (11.9 mmol/L) aqueous sodium bicarbonate	Water
Cefprozil	Water	Water
Ceftaroline	DMSO ^a to 30% of total volume	Saline ^d
Ceftazidime	Sodium carbonate ^e	Water
Ceftibuten	1/10 volume of DMSO ^a	Water
Ceftizoxime	Water	Water
Ceftobiprole	DMSO plus glacial acetic acid ^{a,f}	Water, vortex vigorously

Table 6A. (Continued)

Antimicrobial Agent	Solvent ^b	Diluent ^b		
	Unless otherwise stated, use a minimum amount of the listed solvent to solubilize the antimicrobial powder.	Finish diluting the final stock solution as stated below.		
Ceftolozane	Water or saline ^d	Water or saline ^d		
Ceftriaxone	Water	Water		
Cefuroxime	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L		
Cephalexin	Phosphate buffer, pH 6, 0.1 mol/L	Water		
Cephalothin	Phosphate buffer, pH 6, 0.1 mol/L	Water		
Cephapirin	Phosphate buffer, pH 6, 0.1 mol/L	Water		
Cephradine	Phosphate buffer, pH 6, 0.1 mol/L	Water		
Chloramphenicol	95% ethanol	Water		
Cinoxacin	1/2 volume of water, then add 1 mol/L NaOH dropwise to dissolve	Water		
Ciprofloxacin	Water	Water		
Clarithromycin	Methanol ^a or glacial acetic acid ^{a,c}	Phosphate buffer, pH 6.5, 0.1 mol/L		
Clavulanate	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L		
Clinafloxacin	Water	Water		
Clindamycin	Water	Water		
Colistin ^g	Water	Water		
Dalbavancin	DMSO ^a	DMSO ^{a,h}		
Daptomycin	Water	Water		
Delafloxacin	1/2 volume of water, then 0.1 mol/L NaOH dropwise to dissolve	Water		
Dirithromycin	Glacial acetic acid ^c	Water		
Doripenem	Saline ^d	Saline ^d		
Doxycycline	Water	Water		
Durlobactam	Water	Water		
Enoxacin	1/2 volume of water, then 0.1 mol/L NaOH dropwise to dissolve	Water		
Enmetazobactam	Water	Water		
Eravacycline	Water	Water		
Ertapenem	Phosphate buffer, pH 7.2, 0.01 mol/L	Phosphate buffer, pH 7.2, 0.01 mol/L		
Erythromycin	95% ethanol or glacial acetic acid ^{a,c}	Water		
Exebacase	Supplied as a frozen stock in a buffer containing 20 mM L-histidine and 5% D-sorbitol, pH 7	CAMHB supplemented with 25% horse serum plus 0.5 mM DL-dithiothreitol (pH 7.2–7.4)		
Faropenem	Water	Water		
Fidaxomicin	DMSO ^a	Water		
Finafloxacin	Water	Water		
Fleroxacin	1/2 volume of water, then 0.1 mol/L NaOH dropwise to dissolve	Water		
Fosfomycin	Water	Water		
Fusidic acid	Water	Water		
Garenoxacin	Water (with stirring)	Water		

Table 6A. (Continued)

Antimicrobial Agent	Solvent ^b	Diluent ^b
	Unless otherwise stated, use a minimum amount of the listed solvent to solubilize the antimicrobial powder.	Finish diluting the final stock solution as stated below.
Gatifloxacin	Water (with stirring)	Water
Gemifloxacin	Water	Water
Gentamicin	Water	Water
Gepotidacin	DMSOa	Water
Iclaprim	DMSOa	Water
Imipenem	Phosphate buffer, pH 7.2, 0.01 mol/L	Phosphate buffer, pH 7.2, 0.01 mol/L
Kanamycin	Water	Water
Lefamulin	Water	Water
Levofloxacin	1/2 volume of water, then 0.1 mol/L NaOH dropwise to dissolve	Water
Levonadifloxacin	27.5 μg/mL solution of L-arginine in water	Water
Linezolid	Water	Water
Lomefloxacin	Water	Water
Loracarbef	Water	Water
Mecillinam	Water	Water
Meropenem	Water	Water
Meropenem-vaborbactam	DMSO ^a	Water
Metronidazole	DMSO ^a	Water
Minocycline	Water	Water
Moxalactam (diammonium salt) ⁱ	0.04 mol/L HCI (let sit for 1.5 to 2 hours)	Phosphate buffer, pH 6, 0.1 mol/L
Moxifloxacin	Water	Water
Mupirocin	Water	Water
Nacubactam	Water	Water
Nafcillin	Water	Water
Nafithromycin	½ volume of water, then glacial acetic acid dropwise to dissolve (acetic acid not to exceed 2.5 μL/mL)	Water
Nalidixic acid	1/2 volume of water, then add 1 mol/L NaOH dropwise to dissolve	
Netilmicin	Water	Water
Nitazoxanide	DMSO ^{a,j}	DMSO ^{a,j}
Nitrofurantoin ^k	Phosphate buffer, pH 8, 0.1 mol/L	Phosphate buffer, pH 8, 0.1 mol/L

Table 6A. (Continued)

Antimicrobial Agent	Solvent ^b	Diluent ^b
	Unless otherwise stated, use a minimum amount of the listed solvent to solubilize the antimicrobial powder.	Finish diluting the final stock solution as stated below.
Norfloxacin	1/2 volume of water, then 0.1 mol/L NaOH dropwise to dissolve	Water
Ofloxacin	1/2 volume of water, then 0.1 mol/L NaOH dropwise to dissolve	Water
Omadacycline	Water	Water
Oritavancin	0.002% polysorbate-80 in water ^l	0.002% polysorbate-80 in water ^l
Oxacillin	Water	Water
Ozenoxacin	10% volume of water, then 1M NaOH (8% of final volume)	Water
Penicillin	Water	Water
Pexiganan	Water	Water
Piperacillin	Water	Water
Plazomicin	Water	Water
Polymyxin B	Water	Water
Quinupristin-dalfopristin	Water	Water
Ramoplanin	Water	Water
Razupenem	Phosphate buffer, pH 7.2, 0.01 mol/L	Phosphate buffer, pH 7.2, 0.01 mol/L
Relebactam	Water	Water
Ridinilazole	DMSO ^a	DMSO ^a
Rifampin	Methanol ^a (maximum concentration = 640 μg/mL)	Water (with stirring)
Rifaximin	Methanol ^a	0.1 M phosphate buffer, pH 7.4 + 0.45% sodium dodecyl sulfate
Secnidazole	DMSO ^a	Water
Solithromycin	Glacial acetic acid ^c	Water
Sparfloxacin	Water	Water
Spectinomycin	Water	Water
Streptomycin	Water	Water
Sulbactam	Water	Water
Sulfonamides	1/2 volume hot water and minimal amount of 2.5 mol/L NaOH to dissolve	Water
Sulopenem ^m	0.01 M phosphate buffer, pH 7.2, vortex to dissolve	0.01 M phosphate buffer, pH 7.2
Surotomycin	Water	Water
Taniborbactam	Water	Water
Tazobactam	Water	Water
Tebipenem	Water	Water
Tedizolid	DMSO ^a	DMSO ^{a,n}
Teicoplanin	Water	Water
Telavancin	DMSO ^a	DMSO ^{a,h}
Telithromycin	Glacial acetic acid ^{a,c}	Water

Table 6A
Solvents and Diluents

Table 6A. (Continued)

Antimicrobial Agent	Solvent ^b	Diluent ^b
	Unless otherwise stated, use a minimum amount of the listed solvent to solubilize the antimicrobial powder.	Finish diluting the final stock solution as stated below.
Tetracycline	Water	Water
Ticarcillin	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L
Ticarcillin-clavulanate	Phosphate buffer, pH 6, 0.1 mol/L	Phosphate buffer, pH 6, 0.1 mol/L
Tigecycline	Water	Water
Tinidazole	DMSO ^{a,j}	Water
Tizoxanide	DMSO ^{a,j}	DMSO ^{a,j}
Tobramycin	Water	Water
Trimethoprim	0.05 mol/L lactic ^a or hydrochloric ^a acid, 10% of final volume	Water (may need heat)
Trimethoprim (if lactate)	Water	Water
Trospectomycin	Water	Water
Ulifloxacin (prulifloxacin)	DMSO ^a	Water
Vaborbactam	90% DMSO ^a /10% water	Water
Vancomycin	Water	Water
Zidebactam	Water	Water
Zoliflodacin	DMSO	Water

Abbreviations: **CAMHB**, **cation-adjusted Mueller-Hinton broth**; DMSO, dimethyl sulfoxide; pH, negative logarithm of hydrogen ion concentration.

Footnotes

- a. Consult the safety data sheets before working with any antimicrobial reference standard powder, solvent, or diluent. Some of the compounds (eg, solvents such as DMSO, methanol) are more toxic than others and may necessitate handling in a chemical fume hood.
- b. Although these solvents and diluents are recommended, users should always confirm with the manufacturer.
- c. For glacial acetic acid, use 1/2 volume of water, then add glacial acetic acid dropwise until dissolved, not to exceed 2.5 µL/mL.
- d. Saline a solution of 0.85% to 0.9% NaCl (w/v).
- e. Anhydrous sodium carbonate is used at a weight of exactly 10% of the ceftazidime to be used. The sodium carbonate is dissolved in solution in most of the necessary water. The antimicrobial agent is dissolved in this sodium carbonate solution, and water is added to the desired volume. The solution is to be used as soon as possible, but it can be stored up to six hours at no more than 25°C.
- f. For each 1.5 mg of ceftobiprole, add 110 μL of a 10:1 mixture of DMSO and glacial acetic acid. Vortex vigorously for one minute, then intermittently for 15 minutes. Dilute to 1 mL with distilled water.
- g. The formulation of colistin reference standard powder used in antimicrobial susceptibility tests is colistin sulfate and not colistin methane sulfonate (sulfomethate).

Table 6A. (Continued)

- h. Starting stock solutions of dalbavancin and telavancin should be prepared at concentrations no higher than 1600 µg/mL. Intermediate 100× concentrations should then be diluted in DMSO. Final 1:100 dilutions should then be made directly into CAMHB supplemented with 0.002% (v/v) polysorbate-80, so the final concentration of DMSO in the wells is no greater than 1%. See also Table 8B.
- i. The diammonium salt of moxalactam is very stable, but it is almost pure R isomer. Moxalactam for clinical use is a 1:1 mixture of R and S isomers. Therefore, the salt is dissolved in 0.04 mol/L HCl and allowed to react for 1.5 to 2 hours to convert it to equal parts of both isomers
- j. Final concentration of DMSO should not exceed 1%. This may be accomplished as follows: 1) prepare the stock solution at 10 times higher concentration than planned stock solution (ie, prepare at 12 800 μg/mL, rather than 1280 μg/mL); 2) add 1.8 mL sterile water to each agar deep; 3) add 0.2 mL of each antibiotic dilution to each agar deep
- k. Alternatively, nitrofurantoin is dissolved in DMSO.
- I. Starting stock solutions of oritavancin should be prepared at concentrations no higher than 1600 μg/mL in 0.002% polysorbate-80 in water. Intermediate 100× oritavancin concentrations should then be prepared in 0.002% polysorbate-80 in water. Final 1:100 dilutions should be made directly into CAMHB supplemented with 0.002% polysorbate-80, so the final concentration of polysorbate-80 in the wells is 0.002%.
- m. Must be made fresh on the day of use.
- n. Starting stock solutions of tedizolid should be prepared at concentrations no higher than 1600 μg/mL. Intermediate 100× concentrations should be diluted in DMSO. Final 1:100 dilutions should be made directly into CAMHB, so that the final concentration of DMSO in the wells is no greater than 1%. Also see Table 8B.

NOTE: Information in boldface type is new or modified since the previous edition.

Table 6B. Preparing Stock Solutions for Antimicrobial Agents Provided With Activity Expressed as Units

Antimicrobial	Pure Agent		
Agent	(Reference)	Calculation for µg/mg	Example
Potassium	0.625 µg/unit ¹	Multiply the activity expressed in units/mg by 0.625	Activity units/mg • 0.625 μg/unit = Activity μg/mg
Penicillin G		μg/unit.	
0 "			(eg, 1592 units/mg • 0.625 μg/unit = 995 μg/mg)
Sodium Penicillin G	0.6 μg/unit ¹	Multiply the activity expressed in units/mg by 0.6 μg/unit.	Activity units/mg • 0.6 μg/unit = Activity μg/mg
			(eg, 1477 units/mg • 0.6 μg/unit = 886.2 μg/mg)
Polymyxin B	10 000 units/mg=	Multiply the activity expressed in units/mg by 0.1 μg/unit.	Activity units/mg • 0.1 μg/unit = Activity μg/mg
	10 units/μg=		(eg, 8120 units/mg • 0.1 μg/unit = 812 μg/mg)
	0.1 µg/unit²	Divide the activity expressed in units/mg by 10 units/µg.	Activity units/mg/10 units/μg = Activity μg/mg
			(eg, 8120 units/mg/10 units/mg=812 μg/mg)
Colistin sulfate ^a	30 000 units/mg=	Multiply the activity expressed in units/mg by 0.03333 µg/unit.	Activity units/mg • 0.03333 μg/unit = Activity μg/mg
	30 units/µg=		(eg, 20 277 units/mg • 0.03333 μg/unit = 676 μg/mg)
		Divide the activity expressed in units/mg by 30 units/mg.	Activity units/mg/30 units/μg = Activity μg/mg
	0.03333 µg/unit ²		
			(eg, 20 277 units/mg/30 units/μg=676 μg/mg)
Streptomycin	785 units/mg ³	Divide the number of units given for the powder by 785. This gives the percent purity of the powder. Multiply the	([Potency units/mg]/[785 units/mg]) • (850 μg/mg) = Potency μg/mg
		percent purity by 850, which is the amount in the purest form of streptomycin. This result equals the activity factor	(eg, [751 units/mg/785 units/mg] • 850 μg/mg = 813 μg/mg)
		in μg/mg.	If powder contains 2.8% water:
			813 • (1 – 0.028) = potency
			813 • 0.972 = 790 μg/mg

Footnote

a. Do not use colistin methanesulfonate for in vitro antimicrobial susceptibility tests.

References for Table 6B

- Geddes AM, Gould IM. Benzylpenicillin (penicillin G). In: Grayson ML, ed. *Kucers' The Use of Antibiotics: A Clinical Review of Antibacterial, Antifungal, Antiparasitic and Antiviral Drugs*. 6th ed. Boca Raton, FL: CRC Press, Taylor & Francis Group; 2010:5-58.
- Polymyxins. In: Kucers A, Crowe SM, Grayson ML, Hoy JF, eds. *The Use of Antibiotics: A Clinical Review of Antibacterial, Antifungal, Antiparasitic and Antiviral Drugs.* 5th ed. Oxford, UK: Butterworth-Heinemann; 1997:667-675.
- United States Department of Agriculture, Food Safety and Inspection Service, Office of Public Health Science, Laboratory QA/QC Division. *Bioassay for the detection, identification and quantitation of antimicrobial residues in meat and poultry tissue.* Microbiology Laboratory Guidebook (MLG) 34.03; 2011.

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Table 6C. Preparing Solutions and Media Containing Combinations of Antimicrobial Agents

Antimicrobial Agent	Combination Tested	Preparation	Example
Amikacin- fosfomycin	5:2 ratio (amikacin:fosfomycin)	Prepare 10× starting concentration as 5:2 ratio and dilute as needed. NOTE: Media should be supplemented with 25 µg/mL glucose-6-phosphate.	Example
Amoxicillin- clavulanate	2:1 ratio (amoxicillin:clavulanate)	Prepare 10× starting concentration as 2:1 ratio and dilute as needed.	For a starting concentration of 128/64 in the panel, prepare a 10× stock concentration of 2560 μg/mL for amoxicillin and 1280 μg/mL for clavulanate. Then combine equal amounts of each to the first dilution tube, which will then contain 1280/640 μg/mL of the combination. Dilute 1:10 with broth to achieve the final concentration in microdilution wells.
Ampicillin- sulbactam	2:1 ratio (ampicillin:sulbactam)	Same as amoxicillin-clavulanate.	
Aztreonam- avibactam	Fixed concentration of avibactam at 4 µg/mL	Prepare 10× starting concentration of aztreonam at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of avibactam 80 µg/mL to each of the diluted tubes.	For a starting concentration of 128/4 in the panel, prepare a 10× stock concentration of aztreonam at 2560 μg/mL and dilute by serial twofold increments down to the final concentration needed in the panel. Prepare a stock concentration of avibactam at 80 μg/mL. Then add an equal volume of the avibactam 80 μg/mL solution to each diluted tube of aztreonam. For example, 5 mL of 2560 μg/mL aztreonam+5 mL of 80 μg/mL avibactam=10 mL of 1280/40 μg/mL aztreonam-avibactam. Dilute 1:10 with broth to achieve the final concentration in microdilution wells.
Cefepime- enmetazobactam	Fixed concentration of enmetazobactam at 8 mg/L	Prepare 10× starting concentration of cefepime at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of enmetazobactam 160 µg/mL to each of the diluted tubes.	For a starting concentration of 128/8 in the panel, prepare a 10× stock concentration of cefepime at 2560 μg/mL and dilute by serial twofold increments down to the final concentration needed in the panel. Prepare a stock concentration of enmetazobactam at 160 μg/mL. Then add an equal volume of the enmetazobactam 160 μg/mL solution to each diluted tube of cefepime. For example, 5 mL of 2560 μg/mL cefepime+5 mL of 160 μg/mL enmetazobactam = 10 mL of 1280/80 μg/mL cefepime-enmetazobactam. Dilute 1:10 with broth to achieve the final concentration in the microdilution wells.
Cefepime- taniborbactam	Fixed concentration of taniborbactam at 4 μg/mL	Prepare 10x starting concentration of cefepime at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of taniborbactam 80 μg/mL to each of the diluted tubes.	For a starting concentration of 128/4 in the panel, prepare a 10x stock concentration of cefepime at 2560 µg/mL and dilute by serial twofold increments down to the final concentration needed in the panel. Prepare a stock concentration of taniborbactam at 80 µg/mL. Then add an equal volume of the taniborbactam 80 µg/mL solution to each diluted tube of cefepime. For example, 5 mL of 2560 µg/mL cefepime + 5 mL of 80 µg/mL taniborbactam = 10 mL of 1280/40 µg/mL cefepime-taniborbactam. Dilute 1:10 with broth to achieve the final concentration in microdilution wells.

Table 6C. (Continued)

Antimicrobial	Combination Tasted	Propagation	Evample
Agent Cefepime-	Fixed concentration of	Preparation Prepare 10× starting concentration of	Example For a starting concentration of 128/8 in the panel, prepare a 10×
tazobactam	tazobactam at 8 μg/mL	cefepime at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of tazobactam 160 µg/mL to each of the diluted tubes.	stock concentration of cefepime at 2560 µg/mL and dilute by serial twofold increments down to the final concentration needed in the panel. Prepare a stock concentration of tazobactam at 160 µg/mL. Then add an equal volume of the tazobactam 160 µg/mL solution to each diluted tube of cefepime. For example, 5 mL of 2560 µg/mL cefepime+5 mL of 160 µg/mL tazobactam=10 mL of 1280/80 µg/mL cefepime-tazobactam. Dilute 1:10 with broth to achieve the final concentration in the microdilution wells.
Cefepime- zidebactam	1:1 ratio (cefepime:zidebactam)	Prepare 10× starting concentration as 1:1 ratio and dilute as needed.	For a starting concentration of 128/128 in the panel, prepare a 20× stock concentration of 2560 μg/mL for cefepime and 2560 μg/mL for zidebactam. Then combine equal amounts of each to the first dilution tube, which will then contain 1280/1280 μg/mL of the combination. Prepare twofold serial dilutions and dilute each 1:10 with broth to achieve the final concentration in the microdilution wells.
Ceftaroline- avibactam	Fixed concentration of avibactam at 4 µg/mL	Same as aztreonam-avibactam.	
Ceftazidime- avibactam	Fixed concentration of avibactam at 4 µg/mL	Same as aztreonam-avibactam.	
Ceftolozane- tazobactam	Fixed concentration of tazobactam at 4 μg/mL	Same as aztreonam-avibactam.	
Imipenem- relebactam	Fixed concentration of relebactam at 4 µg/mL	Same as aztreonam-avibactam.	
Meropenem- nacubactam	1:1 ratio (meropenem:nacubactam)	Prepare 10× starting concentration as 1:1 ratio and dilute as needed.	For a starting concentration of 128/128 in the panel, prepare a 20× stock concentration of 2560 µg/mL for meropenem and 2560 µg/mL for nacubactam. Combine equal amounts of each to the first dilution tube, which will then contain 1280/1280 µg/mL of the combination. Prepare 2-fold serial dilutions and dilute each 1:10 with broth to achieve the final concentration in the microdilution wells.
Meropenem- vaborbactam	Fixed concentration of vaborbactam at 8 µg/mL	Prepare 10× starting concentration of meropenem at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of vaborbactam 160 µg/mL to each of the diluted tubes.	For a starting concentration of 64/8 µg/mL in the panel, prepare a 10× stock concentration of meropenem at 1280 µg/mL and dilute by serial twofold increments down to the final concentration needed in the panel. Prepare a stock concentration of vaborbactam at 160 µg/mL. Then add an equal volume of the vaborbactam 160 µg/mL solution to each diluted tube of meropenem. For example, 5 mL of 1280 µg/mL meropenem + 5 mL of 160 µg/mL vaborbactam = 10 mL of 640/80 µg/mL meropenem-vaborbactam. Dilute 1:10 with broth to achieve the final concentration in the microdilution wells.

Table 6C. (Continued)

Antimicrobial Agent	Combination Tested	Preparation	Example
Piperacillin- tazobactam	Fixed concentration of tazobactam at 4 µg/mL	Same as aztreonam-avibactam.	
Sulbactam- durlobactam	Fixed concentration of durlobactam at 4 ug/mL	Prepare 10× starting concentration of sulbactam at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of durlobactam 80 µg/mL to each of the diluted tubes.	For a starting concentration of 128/4 in the panel, prepare a 10× stock concentration of sulbactam at 2560 µg/mL and dilute by serial twofold increments down to the final concentration needed. Prepare a stock concentration of durlobactam at 80 µg/mL. Then add an equal volume of the durlobactam 80 µg/mL solution to each diluted tube of sulbactam. For example, 5 mL of 2560 µg/mL sulbactam+5 mL of 80 µg/mL clavulanate = 10 mL of 1280/40 µg/mL sulbactam-durlobactam. Dilute 1:10 with broth to achieve the final concentration in microdilution wells.
Ticarcillin- clavulanate	Fixed concentration of clavulanate at 2 μg/mL	Prepare 10× starting concentration of ticarcillin at twice the concentration needed and dilute as usual using serial twofold dilutions. Add an equal volume of clavulanate 40 µg/mL to each of the diluted tubes.	For a starting concentration of 128/2 in the panel, prepare a 10× stock concentration of ticarcillin at 2560 µg/mL and dilute by serial twofold increments down to the final concentration needed. Prepare a stock concentration of clavulanate at 40 µg/mL. Then add an equal volume of the clavulanate 40 µg/mL solution to each diluted tube of ticarcillin. For example, 5 mL of 2560 µg/mL ticarcillin+5 mL of 40 µg/mL clavulanate = 10 mL of 1280/20 µg/mL ticarcillin-clavulanate. Dilute 1:10 with broth to achieve the final concentration in microdilution wells.
Trimethoprim- sulfa- methoxazole	1:19 ratio (trimethoprim:sulfamethoxazole)	Prepare a 10× starting concentration of trimethoprim at 1600 μg/mL (or at 1280 μg/mL that will need dilution to 160 μg/mL). Prepare a 10× starting concentration of sulfamethoxazole at a log ₂ multiple of 1520 μg/mL (eg, 1520, 3040, or 6080 μg/mL) depending on the starting concentration needed.	For a starting concentration of 8/152 in the panel, prepare a $10\times$ concentration of trimethoprim at $160~\mu g/mL$. Prepare a $10\times$ starting concentration of sulfamethoxazole at $3040~\mu g/mL$. Add an equal volume of the $160~\mu g/mL$ trimethoprim and the $3040~\mu g/mL$ sulfamethoxazole to the first dilution tube, and then dilute by serial twofold dilutions as usual. For example, $5~mL$ of $160~\mu g/mL$ trimethoprim and $5~mL$ of $3040~\mu g/mL$ sulfamethoxazole = $10~mL$ of $80/1520$ trimethoprimsulfamethoxazole. Dilute $1:10~mL$ broth to achieve the final concentration in microdilution wells.
Quinupristin- dalfopristin	Preparation usually not necessary, because drug powder is received as combination.		

Table 6C. (Continued)

NOTE 1: To prepare intermediate dilutions of antimicrobial agents, a convenient formula to use is $C_1 \cdot V_1 = C_2 \cdot V_2$, where C_1 is the concentration of stock solution of the antimicrobial agent (usually 1280 μ g/mL or greater); V_1 is the unknown volume that will be needed to make the intermediate concentration; C_2 is the intermediate concentration needed; and V_2 is the volume of the intermediate stock solution needed. For example, to prepare 20 mL of a 40 μ g/mL solution from a 1280 μ g/mL stock solution:

C₁•V₁=C₂•V₂

1280
$$\mu$$
g/mL•V₁=40 μ g/mL•20 mL

V₁= $\frac{40 \mu$ g/mL•20 mL}
1280 μ g/mL

Therefore, add 0.625 mL of the 1280 µg/mL stock solution to 19.375 mL of diluent (usually water) for a final volume of 20 mL of a 40 µg/mL solution.

NOTE 2: Information in boldface type is new or modified since the previous edition.

Table 7. Preparing Dilutions of Antimicrobial Agents to Be Used in Agar Dilution Susceptibility Tests

Antimicrobial Solution										
Step	Concentration, μg/mL	Source	Volume, mL	+	Diluent, mL	=	Intermediate Concentration, µg/mL	=	Final Concentration at 1:10 Dilution in Agar, μg/mL	Log ₂
	5120	Stock	_		_		5120		512	9
1	5120	Stock	2		2		2560		256	8
2	5120	Stock	1		3		1280		128	7
3	5120	Stock	1		7		640		64	6
4	640	Step 3	2		2		320		32	5
5	640	Step 3	1		3		160		16	4
6	640	Step 3	1		7		80		8	3
7	80	Step 6	2		2		40		4	2
8	80	Step 6	1		3		20		2	1
9	80	Step 6	1		7		10		1	0
10	10	Step 9	2		2		5		0.5	-1
11	10	Step 9	1		3		2.5		0.25	-2
12	10	Step 9	1		7		1.25		0.125	-3

NOTE: This table is modified from Ericsson HM, Sherris JC. Antibiotic sensitivity testing: report of an international collaborative study. *Acta Pathol Microbiol Scand B Microbiol Immunol.* 1971;217(suppl):1+.

When serial twofold dilution minimal inhibitory concentrations are being prepared and tested, the actual dilution scheme is:

128, 64, 32, 16, 8, 4, 2, 1, 0.5, 0.25, 0.125, 0.0625, 0.03125, 0.015625, 0.0078125, 0.0039063, 0.0019531 µg/mL, etc.

For convenience only, and not because these are the actual concentrations tested, it was decided to use the following values in these tables:

 $128, 64, 32, 16, 8, 4, 2, 1, 0.5, 0.25, 0.12, 0.06, 0.03, 0.016, 0.008, 0.004, 0.002 \mug/mL, etc.$

The values that appear in the tables are equivalent to the actual values tested, eg, 0.12 μg/mL = 0.125 μg/mL, 0.016 μg/mL = 0.015625 μg/mL.

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Table 8A. Preparing Dilutions of Antimicrobial Agents to Be Used in Broth Dilution Susceptibility Tests

	Antin	nicrobial Solution						
Step	Concentration, ^a μg/mL	Source	Volume, ^a mL	+	CAMHB ^b Volume, ^c mL	"	Final Concentration, µg/mL	Log₂
1	5120	Stock	1		9		512	9
2	512	Step 1	1		1		256	8
3	512	Step 1	1		3		128	7
4	512	Step 1	1		7		64	6
5	64	Step 4	1		1		32	5
6	64	Step 4	1		3		16	4
7	64	Step 4	1		7		8	3
8	8	Step 7	1		1		4	2
9	8	Step 7	1		3		2	1
10	8	Step 7	1		7		1	0
11	1	Step 10	1		1		0.5	-1
12	1	Step 10	1		3		0.25	-2
13	1	Step 10	1		7		0.125	-3

Abbreviation: CAMHB, cation-adjusted Mueller-Hinton broth.

Footnotes

- a. See Table 7 for the actual dilution scheme when serial twofold dilution minimal inhibitory concentrations are being prepared and tested.
- b. Adjustment with cations, if necessary, occurs before this step.
- c. The volumes selected can be any multiple of these figures, depending on the number of tests to be performed.

NOTE: This table is modified from Ericsson HM, Sherris JC. Antibiotic sensitivity testing: report of an international collaborative study. *Acta Pathol Microbiol Scand B Microbiol Immunol*. 1971;217(suppl):1:+.

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Table 8B. Preparing Dilutions of Water-Insoluble Antimicrobial Agents to Be Used in Broth Dilution Susceptibility Tests

Antimicrobial Solution										
Step	Concentration, μg/mL	Source	Volume, mL	+	Solvent, mL (eg, DMSO)	=	Intermediate Concentration, µg/mL	=	Final Concentration at 1:100, μg/mL	Log₂
1	1600	Stock					1600		16	4
2	1600	Stock	0.5		0.5		800		8.0	3
3	1600	Stock	0.5		1.5		400		4.0	2
4	1600	Stock	0.5		3.5		200		2.0	1
5	200	Step 4	0.5		0.5		100		1.0	0
6	200	Step 4	0.5		1.5		50		0.5	-1
7	200	Step 4	0.5		3.5		25		0.25	-2
8	25	Step 7	0.5		0.5		12.5		0.125	-3
9	25	Step 7	0.5		1.5		6.25		0.0625	-4
10	25	Step 7	0.5		3.5		3.1		0.03	-5
11	3.1	Step 10	0.5		0.5		1.6		0.015	-6
12	3.1	Step 10	0.5		1.5		0.8		0.008	-7
13	3.1	Step 10	0.5		3.5		0.4		0.004	-8
14	0.4	Step 13	0.5		0.5		0.2		0.002	-9

Abbreviation: DMSO, dimethyl sulfoxide.

For Use With M07—MIC Testing

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M100, 30th ed.

Appendix A. Suggestions for Confirming Antimicrobial Susceptibility Test Results and Organism Identification for Agents Approved by the US Food and Drug Administration for Clinical Use

the US Food and	Drug Administration for Clin	icai use			
				ignificance of Resistar	
			Take Foll	owing Confirmation of	f Results ^a
			Category I	Category II	Category III
			Not reported or only rarely reported to date	Uncommon in most institutions	May be common but generally considered of epidemiological concern
				Action Steps:	
			Confirm ID and susceptibility. ^a Report to infection prevention. Check with public health department to determine appropriate reporting and isolate referral	Confirm ID and susceptibility if uncommon in the institution. ^a Check with infection prevention in the facility to determine if special reporting procedures or	Confirm ID and susceptibility if uncommon in the institution. Check with infection prevention in the facility to determine if special reporting procedures or
			procedures. • Save isolate.	additional actions are needed. Check with public	additional action are needed.
			NOTE: It may be appropriate to notify infection prevention of preliminary findings	health department to determine appropriate reporting and	
Organism or		Antimicrobial Agent(s) and	before confirmation	isolate referral	
Organism Group	Antimicrobial Class/Subclass	Resistance Phenotype Detected ^a	of results.	procedures.	
Any	β-lactam combination agents	Ceftazidime-avibactam – R		X	
Enterobacterales		Meropenem-vaborbactam – I or R		X	
	Carbapenems	Any carbapenem – I or R ^b		X	
	Aminoglycosides	Amikacin, gentamicin, and tobramycin – R			Х
		Plazomicin – R (except P. mirabilis)	X		
	Lipopeptides	Colistin/Polymyxin B – R	Х		

Appendix A. (Continued)

				ignificance of Resistal owing Confirmation o	
			Category I	Category II	Category III
Organism or Organism Group	Antimicrobial Class/Subclass	Antimicrobial Agent(s) and Resistance Phenotype Detected ^a	Not reported or only rarely reported to date	Uncommon in most institutions	May be common but generally considered of epidemiological concern
Escherichia coli, Klebsiella pneumoniae, K. oxytoca, and Proteus mirabilis	Cephems	Cephalosporin III/IV – I/SDD or R			Х
Salmonella and	Cephems	Cephalosporin III – I or R		X	
Shigella spp.c	Macrolides	Azithromycin – NWT or R		X	
	Fluoroquinolones	Any fluoroquinolone – I or R		X	
Acinetobacter	Carbapenems	Any carbapenem ^d – I or R			Х
baumannii complex	Lipopeptides	Colistin/polymyxin B – R	х		
Pseudomonas	β-lactam combination agents	Ceftolozane-tazobactam – I or R		X	
aeruginosa	Carbapenems	Any carbapenem ^d – I or R			X
	Aminoglycosides	Amikacin and gentamicin and tobramycin – R			X
	Lipopeptides	Colistin/polymyxin B – R	Х		
Stenotrophomonas maltophilia	Folate pathway antagonists	Trimethoprim-sulfamethoxazole – I or R			Х

Х

Aminoglycosides

M100, 30th ed

Appendix A. (Continued) Occurrence and Significance of Resistance and Actions to Take Following Confirmation of Results^a Category I Category II Category III May be common but generally Not reported or considered of Antimicrobial Agent(s) and Organism or only rarely Uncommon in epidemiological Resistance Phenotype Detected^a **Organism Group Antimicrobial Class/Subclass** reported to date most institutions concern Haemophilus **Penicillins** Ampicillin – R and β-lactamase negative Х influenzae Amoxicillin-clavulanate - R Χ **β-lactam combination agents** Cephems Cephalosporin III/IV - NS Χ Ceftaroline - NS Carbapenems **Any** carbapenem – NS Χ

Fluoroquinolones Any fluoroquinolone – NS Χ Cephems Cephalosporin III/IV - NS Χ Neisseria gonorrhoeae Macrolides Azithromycin – NS Χ Fluoroquinolones Ciprofloxacin – I or R Χ **Glycopeptides** Vancomycin – R Х Enterococcus spp. Lipoglycopeptides Dalbavancin - NS Χ (Vancomycin-susceptible Oritavancin – NS E. faecalis only) Telavancin - NS Lipopeptides Daptomycin - SDD, I, or R Χ Oxazolidinones Linezolid – R Χ Tedizolid - NS

Gentamicin high level - R

Streptomycin high level - R

Appendix A. (Continued)

				Significance of Resistal Ilowing Confirmation o	
			Category I	Category II	Category III
Organism or Organism Group	Antimicrobial Class/Subclass	Antimicrobial Agent(s) and Resistance Phenotype Detected ^a	Not reported or only rarely reported to date	Uncommon in most institutions	May be common but generally considered of epidemiological concern
Staphylococcus aureus	Penicillinase-stable penicillins	Oxacillin – R			Х
	Cephems	Ceftaroline – SDD or R		X	
	Glycopeptides	Vancomycin – I ^e		Х	
		Vancomycin – R	Х		
	Lipoglycopeptides	Dalbavancin – NS Oritavancin – NS Telavancin – NS	X		
	Lipopeptides	Daptomycin – NS		Х	
	Streptogramins	Quinupristin-dalfopristin (MSSA only) – I or R		Х	
	Oxazolidinones	Linezolid – R Tedizolid – I or R		Х	
Staphylococcus	Glycopeptides	Vancomycin – I or R ^f		X	
spp. other than	Lipopeptides	Daptomycin – NS		X	
S. aureus	Oxazolidinones	Linezolid – R		X	

M100, 30th ed

Appendix A. (Continued) Occurrence and Significance of Resistance and Actions to Take Following Confirmation of Results^a Category I Category II Category III May be common but generally Not reported or considered of Antimicrobial Agent(s) and Organism or only rarely Uncommon in most epidemiological **Organism Group Antimicrobial Class/Subclass** Resistance Phenotype Detected^a reported to date institutions concern Streptococcus Penicillins Amoxicillin or penicillin (nonmeningitis) -Χ pneumoniae R Cephalosporin III/IV (nonmeningitis) - R Χ Cephems Ceftaroline (nonmeningitis) - NS Х Any carbapenem - I, R, or NS Х Carbapenems Vancomycin – NS **Glycopeptides** Χ **Any** fluoroquinolone – I or R **Fluoroquinolones** Χ Streptogramins Quinupristin-dalfopristin - I or R Х Rifampin – I or R Χ **Ansamycins** Oxazolidinones Linezolid - NS Х Ampicillin or penicillin - NS Χ Streptococcus. **Penicillins** β-hemolytic group Cephalosporin III/IV - NS Х Cephems Ceftaroline – NS Any carbapenem - NS Carbapenems Х Χ **Glycopeptides** Vancomycin – NS Dalbavancin – NS Χ Lipoglycopeptides Oritavancin - NS Χ Χ Telavancin – NS Lipopeptides Daptomycin - NS Χ Streptogramins Quinupristin-dalfopristin (S. pyogenes Х only) – I or R Oxazolidinones Linezolid - NS X

Tedizolid - NS

Appendix A. (Continued)

				Significance of Resista Howing Confirmation of	
			Category I	Category II	Category III
Organism or Organism Group	Antimicrobial Class/Subclass	Antimicrobial Agent(s) and Resistance Phenotype Detected ^a	Not reported or only rarely reported to date	Uncommon in most institutions	May be common but generally considered of epidemiological concern
Streptococcus,	Carbapenems	Any carbapenem – NS	X		
viridans group	Glycopeptides	Vancomycin – NS	X		
	Lipoglycopeptides	Dalbavancin (S. anginosus group only) – NS	Х		
		Oritavancin – NS Telavancin – NS	X X		
	Streptogramins	Quinupristin-dalfopristin – I or R	X		
	Oxazolidinones	Linezolid – NS Tedizolid – NS	X X		
Neisseria	Penicillins	Ampicillin or penicillin – I		X	
meningitidis		Ampicillin or penicillin – R	X		
	Cephems	Cephalosporin III- NS	X		
	Carbapenems	Meropenem – NS	X		
	Macrolides	Azithromycin – NS		Х	
	Tetracyclines	Minocycline – NS		Х	
	Fluoroquinolones	Any fluoroquinolone – I or R		Х	
	Phenicols	Chloramphenicol – I or R		Х	
	Ansamycins	Rifampin – I or R		X	
Bacteroides spp.	Carbapenems	Any carbapenem – I or R		Х	·
and <i>Parabacteroides</i> spp.	Nitroimidazoles	Metronidazole – I or R		Х	

Abbreviations: I, intermediate; ID, identification; mCIM, modified carbapenem inactivation method; MIC, minimal inhibitory concentration; MSSA, methicillin (oxacillin)-susceptible *Staphylococcus aureus;* NS, nonsusceptible; NWT, non-wild-type; R, resistant; SDD, susceptible-dose dependent.

Appendix A. (Continued)

Footnotes

- a. Ensure antimicrobial susceptibility test results and organism identification are accurate and reproducible. Consider the following steps:
 - 1. Check for transcription errors, contamination, or defective panel, plate, or card.
 - 2. Check previous reports on the patient to determine if the isolate was encountered and confirmed earlier.
 - 3. Repeat organism identification and antimicrobial susceptibility tests with initial method to ensure they reproduce. For category I and II, the laboratory may elect to skip step 3 and go to steps 4 and 5. For category III, repeat and/or confirmatory testing may not be needed if resistance is common in the institution.
 - 4. Confirm organism identification with second method performed in-house or at a referral laboratory.
 - 5. Confirm antimicrobial susceptibility test results with second method (eg, in-house or referral laboratory). The second method might be a CLSI reference method (eg, broth microdilution, agar dilution, or disk diffusion) or a US Food and Drug Administration—cleared commercial test.
- b. Imipenem MICs for *Proteus* spp., *Providencia* spp., and *Morganella morganii* tend to be higher (eg, MI–Cs in the intermediate or resistant category than those with meropenem or doripenem MICs. **MICs for these agents may be elevated due to mechanisms other than carbapenemases among these organisms.** A phenotypic test such as mCIM or CarbaNP may be used to identify carbapenemase-producing isolates (see Tables 3A and 3B).
- c. When submitting the report to a public health department, include antimicrobial susceptibility test results for *Salmonella* spp. that are intermediate or resistant to third-generation cephalosporins (cephalosporin III) and/or intermediate or resistant to fluoroquinolone or resistant to nalidixic acid.
- d. Excludes organisms with intrinsic resistance to listed agents as described in Appendix B.
- e. S. aureus isolates demonstrating vancomycin MICs 4 μg/mL may represent testing variation and need not be reported or submitted to public health department; S. aureus isolates demonstrating MICs > 4 μg/mL should be reported to the local public health department.
- f. There are some Staphylococcus spp. other than S. aureus for which vancomycin MICs may test within the intermediate range (MIC 8–16 μg/mL). In contrast, vancomycin-resistant Staphylococcus spp. (MIC ≥ 32 μg/mL) are rare.

NOTE 1: NS: A category used for isolates for which only a susceptible interpretive criterion has been designated because of the absence or rare occurrence of resistant strains. Isolates that have MICs above or zone diameters below the value indicated for the susceptible breakpoint should be reported as nonsusceptible.

NOTE 2: An isolate that is interpreted as nonsusceptible does not necessarily mean that the isolate has a resistance mechanism. It is possible that isolates with MICs above the susceptible breakpoint that lack resistance mechanisms may be encountered within the wild-type distribution subsequent to the time the susceptible-only breakpoint is set.

NOTE 3: For strains yielding results in the "nonsusceptible" category, organism identification and antimicrobial susceptibility test results should be confirmed (see footnote "a").

NOTE 4: Information in boldface type is new or modified since the previous edition.

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Appendix B Intrinsic Resistance

Appendix B. Intrinsic Resistance

Intrinsic resistance is defined as inherent or innate (not acquired) antimicrobial resistance, which is reflected in wild-type antimicrobial patterns of all or almost all representatives of a species. Intrinsic resistance is so common that susceptibility testing is unnecessary. For example, *Citrobacter* spp. are intrinsically resistant to ampicillin.

These tables can be helpful in at least three ways: 1) they provide a way to evaluate the accuracy of testing methods; 2) they aid in the recognition of common phenotypes; and 3) they can assist with verification of cumulative antimicrobial susceptibility test data. In the tables, an "R" occurring with an antimicrobial agent/organism combination means that strains should test resistant. A small percentage (1% to 3%) may appear susceptible due to method variation, mutation, or low levels of resistance expression.

Each laboratory should decide which agents to test and report in consultation with institutional leaders representing infectious diseases practitioners, the pharmacy and therapeutics and infection **prevention** committees of the medical staff, and the antimicrobial stewardship team. If tested, the result for an antimicrobial agent/organism combination listed as having intrinsic resistance should be reported as resistant. Consideration may be given to adding comments regarding intrinsic resistance of agents not tested. See Appendix A, footnote "a."

Appendix B. (Continued)

B1. Enterobacterales

Antimicrobial Agent Organism	Ampicillin	Amoxicillin- clavulanate	Ampicillin- sulbactam	Ticarcillin	Cephalosporins I: Cefazolin, Cephalothin	Cephamycins: Cefoxitin, Cefotetan	Cephalosporin II: Cefuroxime	Imipenem	Tetracyclines	Tigecycline	Nitrofurantoin	Polymyxin B Colistin	Aminoglycosides
Citrobacter freundii	R	R	R		R	R	R						
Citrobacter koseri, Citrobacter amalonaticus group ^a	R			R									
Enterobacter cloacae complexb	R	R	R		R	R							
Escherichia coli	There is	s no intrin	sic resista	ance to β-	lactams in	this organi	sm.						
Escherichia hermannii	R			R									
Hafnia alvei	R	R	R		R	R							
Klebsiella (formerly Enterobacter) aerogenes	R	R	R		R	R							
Klebsiella pneumoniae, Klebsiella oxytoca, Klebsiella variicola	R			R									
Morganella morganii	R	R			R		R	С		R	R	R	
Proteus mirabilis		s no intrin organism.	sic resista	ance to pe	enicillins an	d cephalos	porins	С	R	R	R	R	
Proteus penneri	R				R		R	С	R	R	R	R	
Proteus vulgaris	R				R		R	С	R	R	R	R	
Providencia rettgeri	R	R			R			С	R	R	R	R	
Providencia stuartii	R	R			R			С	R	R	R	R	d
Raoultella spp.e	R			R									

Appendix B. (Continued)

B1. Enterobacterales (Continued)

Antimicrobial Agent Organism	Ampicillin	Amoxicillin- clavulanate	Ampicillin- sulbactam	Ticarcillin	Cephalosporins I: Cefazolin, Cephalothin	Cephamycins: Cefoxitin, Cefotetan	Cephalosporin II: Cefuroxime	Imipenem	Tetracyclines	Tigecycline	Nitrofurantoin	Polymyxin B Colistin	Aminoglycosides
Salmonella and Shigella spp.					lactams in	these orga	nisms;						
	refer to	WARNIN	G below t	for reporti	ng.								
Serratia marcescens	R	R	R		R	R	R		•		R	R	
Yersinia enterocolitica	R	R		R	R								

Abbreviation: R, resistant.

WARNING: For Salmonella spp. and Shigella spp., aminoglycosides, first- and second-generation cephalosporins, and cephamycins may appear active in vitro but are not effective clinically and should not be reported as susceptible.

Footnotes

- a. Citrobacter amalonaticus group includes C. amalonaticus, C. farmeri, and C. sedlakii.
- b. E. cloacae complex includes Enterobacter asburiae, Enterobacter cloacae, and Enterobacter hormaechei. Other members of the complex include Enterobacter kobei and Enterobacter ludwigii, for which antimicrobial susceptibility testing data are not available.
- c. *Proteus* spp., *Providencia* spp., and *Morganella* spp. may have elevated minimal inhibitory concentrations to imipenem by mechanisms other than by production of carbapenemases. Isolates that test as susceptible should be reported as susceptible.
- d. P. stuartii should be considered resistant to gentamicin, netilmicin, and tobramycin but not intrinsically resistant to amikacin.
- e. Raoultella spp. includes R. ornithinolytica, R. terrigena, and R. planticola.

NOTE 1: Cephalosporins III, cefepime, aztreonam, ticarcillin-clavulanate, piperacillin-tazobactam, and the carbapenems are not listed, because there is no intrinsic resistance in **Enterobacterales**.

NOTE 2: Enterobacterales are also intrinsically resistant to clindamycin, daptomycin, fusidic acid, glycopeptides (vancomycin), lipoglycopeptides (oritavancin, teicoplanin, telavancin), linezolid, tedizolid, quinupristin-dalfopristin, rifampin, and macrolides (erythromycin, clarithromycin, and azithromycin). However, there are some exceptions with macrolides (eg, *Salmonella* and *Shigella* spp. with azithromycin).

NOTE 3: Information in boldface type is new or modified since the previous edition.

Appendix B. (Continued)

B2. Non-Enterobacterales

Antimicrobial Agent Organism	Ampicillin, Amoxicillin	Piperacillin	Ticarcillin	Ampicillin-sulbactam	Amoxicillin- clavulanate	Piperacillin-tazobactam	Cefotaxime	Ceftriaxone	Ceftazidime	Cefepime	Aztreonam	Imipenem	Meropenem	Ertapenem	Polymyxin B Colistin	Aminoglycosides	Tetracyclines/ Tigecycline	Trimethoprim	Trimethoprim- sulfamethoxazole	Chloramphenicol	Fosfomycin
Acinetobacter baumannii/ Acinetobacter calcoaceticus complex	R				R						R			R				R		R	R
Burkholderia cepacia complex ^a	R	R	R	R	R	а	а	а		а	а	а		R	R	а		а			R
Pseudomonas aeruginosa	R			R	R		R	R						R			R	R	R	R	
Stenotrophomonas maltophilia	R	R	R	R	R	R	R	R			R	R	R	R		R	b	R			R

Abbreviation: MIC, minimal inhibitory concentration; R, resistant.

Footnotes

- a. B. cepacia complex isolates have chromosomal genes that require mutational changes before leading to resistance. It is not known how often these mutations occur during growth. Intrinsic resistance implies the presence of resistance mechanisms in natural or wild-type strains that result in phenotypic resistance for all or nearly all strains. Environmental B. cepacia complex strains lacking mutations do not express resistance mechanisms, resulting in low MICs to many antimicrobial agents, whereas clinical strains that express resistance genes, such as those from cystic fibrosis patients, have high MIC values to these same antimicrobial agents. There is insufficient clinical evidence to confirm whether strains that test susceptible in vitro, despite the presence of resistance mechanisms, will respond in vivo. Therefore, intrinsic resistance to the footnoted antibiotics (listed as resistant in previous editions of M100) cannot be confirmed
- b. S. maltophilia is intrinsically resistant to tetracycline but not to doxycycline, minocycline, or tigecycline.

NOTE 1: These nonfermentative gram-negative bacteria are also intrinsically resistant to penicillin (ie, benzylpenicillin), cephalosporins I (cephalosporins I (cephalosporin II (cefuroxime), cephamycins (cefoxitin, cefotetan), clindamycin, daptomycin, fusidic acid, glycopeptides (vancomycin), linezolid, macrolides (erythromycin, azithromycin, clarithromycin), quinupristin-dalfopristin, and rifampin.

NOTE 2: Information in boldface type is new or modified since the previous edition.

Appendix B. (Continued)

B3. Staphylococci

Antimicrobial Agent Organism	Novobiocin	Fosfomycin	Fusidic Acid
S. aureus S. lugdunensis	There	is no intrinsic resistance in these spe	ecies.
S. epidermidis			
S. haemolyticus			
S. saprophyticus	R	R	R
S. capitis		R	
S. cohnii	R		
S. xylosus	R		

Abbreviations: MRS, methicillin (oxacillin) resistant staphylococci; R, resistant.

NOTE 1: These gram-positive bacteria are also intrinsically resistant to aztreonam, polymyxin B/colistin, and nalidixic acid.

NOTE 2: MRS, as defined by cefoxitin or oxacillin testing, as appropriate to the species, are considered resistant to other β -lactam agents, ie, penicillins, β -lactam combination agents, cephems with the exception of ceftaroline, and carbapenems. This is because most cases of documented MRS infections have responded poorly to β -lactam therapy, or because convincing clinical data that document clinical efficacy for those agents have not been presented.

NOTE 3: Information in boldface type is new or modified since the previous edition.

Appendix B. (Continued)

B4. Enterococcus spp.

Antimicrobial Agent Organism	Cephalosporins	Vancomycin	Teicoplanin	Aminoglycosides	Clindamycin	Quinupristin-dalfopristin	Trimethoprim	Trimethoprim-sulfamethoxazole	Fusidic Acid
E. faecalis	Rª			Rª	Rª	R	R	Rª	R
E. faecium	Rª			Rª	Rª		R	Rª	R
E. gallinarum/E. casseliflavus	Rª	R		Rª	Rª	R	R	Rª	R

Abbreviation: R, resistant.

a. **Warning:** For *Enterococcus* spp., cephalosporins, aminoglycosides (except for high-level resistance testing), clindamycin, and trimethoprim-sulfamethoxazole may appear active *in vitro* but are not effective clinically and should not be reported as susceptible.

NOTE: These gram-positive bacteria are also intrinsically resistant to aztreonam, polymyxin B/colistin, and nalidixic acid.

Appendix B. (Continued)

B5. Anaerobic Gram-Positive Bacilli

Antimicrobial Agent Organism	Vancomycin	Aminoglycosides
Clostridium and Clostridioides spp.		R
Clostridium innocuum	R	R

Abbreviation: R, resistant.

B6. Anaerobic Gram-Negative Bacilli

Antimicrobial Agent Organism	Aminoglycosides	Penicillin	Ampicillin	Quinolones
Bacteroides spp.	R	R	R	
Fusobacterium canifelinum	R			R

Abbreviation: R, resistant.

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Appendix C. QC Strains for Antimicrobial Susceptibility Tests

	Organism	Disk Diffusion			
QC Strains	Characteristics	Tests	MIC Tests	Other Tests	Comments
Acinetobacter baumannii NCTC 13304 ^{a,b}	OXA-27 (carbapenemase)	β-lactam combination agents	β-lactam combination agents		
Bacteroides fragilis ATCC®c 25285	β-lactamase positive		All anaerobes		
Bacteroides thetaiotaomicron ATCC® 29741	β-lactamase positive		All anaerobes		
Clostridioides (formerly Clostridium) difficile ATCC® 700057	β-lactamase negative		Gram-positive anaerobes		
Eggerthella lenta (formerly Eubacterium lentum) ATCC® 43055			All anaerobes		Growth on Brucella medium not optimal No longer required when establishing new QC ranges due to organism variability
Enterococcus faecalis ATCC [®] 29212			Nonfastidious gram- positive bacteria	Vancomycin agar HLAR tests High-level mupirocin resistance MIC test	 Assess suitability of medium for sulfonamide or trimethoprim MIC and disk diffusion tests.^d Assess suitability of cation content in each batch/lot of MHB for daptomycin broth microdilution. Agar dilution has not been validated for daptomycin.
E. faecalis ATCC® 33186					Alternative to <i>E. faecalis</i> ATCC® 29212 to assess suitability of MHA for sulfonamide or trimethoprim disk diffusion tests.d
E. faecalis ATCC® 51299	 vanB (vancomycin resistant) Resistant to high-level aminoglycosides 			Vancomycin agarHLAR tests	

Appendix C. (Continued)

QC Strains	Organism Characteristics	Disk Diffusion Tests	MIC Tests	Other Tests	Comments
Escherichia coli ATCC® 25922	β-lactamase negative	Nonfastidious gram-negative bacteria Neisseria meningitidis	Nonfastidious gram- negative bacteria N. meningitidis		
E. coli ATCC® 35218 ^{a,b,1,2}	• TEM-1	 β-lactam combination agents 	β-lactam combination agents		
E. coli NCTC 13353 ^{a,b,3}	• CTX-M-15 (ESBL)	β-lactam combination agents	β-lactam combination agents		
<i>E. coli</i> AR Bank #0349 ⁴	• MCR-1			Colistin broth disk elution Colistin agar test	
Haemophilus influenzae ATCC® 10211					Assess each batch/lot of HTM for growth capabilities.
H. influenzae ATCC® 49247	• BLNAR	H. influenzaeHaemophilus parainfluenzae	H. influenzae H. parainfluenzae		
H. influenzae ATCC® 49766	Ampicillin susceptible	H. influenzae H. parainfluenzae	H. influenzae H. parainfluenzae		More reproducible than H. influenzae ATCC® 49247 with selected β-lactam agents
Klebsiella pneumoniae ATCC® 700603 ^{a,b}	 SHV-18 (ESBL)^{1,2} OXA-2 Mutations in OMPK35 and OMPK37 	β-lactam combination agents	β-lactam combination agents	ESBL tests	
K. pneumoniae ATCC [®] BAA- 1705™ ^{a,b}	KPC-2 (carbapenemase) TEM SHV	β-lactam combination agents	β-lactam combination agents	Carbapenemase tests	
K. pneumoniae ATCC [®] BAA- 1706™	Resistant to carbapenems by noncarbapenemase mechanism			Carbapenemase tests	
K. pneumoniae ATCC® BAA- 2146™	• NDM			Carbapenemase tests	

M100, 30th ed.

	Organism	Disk Diffusion			
QC Strains	Characteristics	Tests	MIC Tests	Other Tests	Comments
K. pneumoniae ATCC [®] BAA- 2814 ^{™a,b} – previously B21(KP1074)	KPC-3 (carbapenemase)SHV-11TEM-1	β-lactam combination agents	β-lactam combination agents		Higher MIC (see Table 5A-2) and better indicator of antimicrobial agent stability than K. pneumoniae BAA-1705™
Neisseria gonorrhoeae ATCC® 49226	• CMRNG	N. gonorrhoeae	N. gonorrhoeae		
Pseudomonas aeruginosa ATCC® 27853e	Inducible AmpC β-lactamase	Nonfastidious gram-negative bacteria	Nonfastidious gram- negative bacteria		Assess suitability of cation content in each batch/lot of CAMHB.
Staphylococcus aureus ATCC® 25923	 β-lactamase negative mecA negative mupA negative 	Nonfastidious gram-positive bacteria		High-level mupirocin resistance disk diffusion test ICR disk diffusion test (D-zone test)	Little value in MIC testing due to its extreme susceptibility to most drugs
S. aureus ATCC® 29213	 Weak β-lactamase– producing strain mecA negative mupA negative 		Nonfastidious gram-positive bacteria	 Oxacillin salt agar High-level mupirocin resistance MIC test ICR MIC test Penicillin zone-edge test 	Assess suitability of cation content in each batch/lot of MHB for daptomycin broth microdilution.
S. aureus ATCC® 43300	mecA positive	Cefoxitin disk diffusion testing	Cefoxitin MIC testing	Oxacillin salt agar	
S. aureus ATCC® BAA-976™	msrA-mediated macrolide-only resistance			ICR MIC test and disk approximation test (D-zone test)	
S. aureus ATCC [®] BAA-977™	Inducible <i>erm</i> A-mediated macrolide resistance			ICR MIC test and disk approximation test (D-zone test)	

Appendix C. (Continued)

QC Strains	Organism Characteristics	Disk Diffusion Tests	MIC Tests	Other Tests	Comments
S. aureus ATCC [®] BAA-1708™	mupA-mediated high- level mupirocin resistance			High-level mupirocin resistance test	
Streptococcus pneumoniae ATCC® 49619	Penicillin intermediate by altered penicillin-binding protein	S. pneumoniae Streptococcus spp. N. meningitidis	S. pneumoniaeStreptococcus spp.N. meningitidis	ICR MIC test	

Abbreviations: ATCC[®], American Type Culture Collection; BLNAR, β-lactamase negative, ampicillin-resistant; CAMHB, cation-adjusted Mueller-Hinton broth; CMRNG, chromosomally mediated penicillin-resistant *Neisseria gonorrhoeae;* ESBL, extended-spectrum β-lactamase; HLAR, high-level aminoglycoside resistance; HTM, *Haemophilus* test medium; **ICR, inducible clindamycin resistance;** MHA, Mueller-Hinton agar; MHB, Mueller-Hinton broth; MIC, minimal inhibitory concentration; NCTC, National Collection of Type Cultures; QC, quality control.

Footnotes

- a. Careful attention to organism maintenance (eg, minimal subcultures) and storage (eg, -60°C or below) is especially important for these QC strains because spontaneous loss of the plasmid encoding the β-lactamase has been documented. If stored at temperatures above -60°C or if repeatedly subcultured, these strains may lose their resistance characteristics and QC results may be outside the acceptable ranges.
- b. To confirm the integrity of the QC strain, test one of the single β-lactam agents highlighted in orange in Tables 4A-2 and 5A-2 by either a disk diffusion or MIC test when the strain is first subcultured from a frozen or lyophilized stock culture. In-range results for the single agent indicate the QC strain is reliable for QC of β-lactam combination agents. It is not necessary to check the QC strain again with a single agent until a new frozen or lyophilized stock culture is put into use.
- c. ATCC® is a registered trademark of the American Type Culture Collection. Per ATCC® convention, the trademark symbol is used after "BAA" in each catalog number, in conjunction with the registered ATCC® name.
- d. Disk diffusion and MIC end points should be easy to read as 80% or greater reduction in growth if the medium has acceptable levels of thymidine.
- e. May develop resistance to β-lactam antimicrobial agents after repeated subcultures. Minimize this risk by subculturing from a frozen or lyophilized stock culture at least monthly or whenever the strain demonstrates results outside the acceptable range.

M100, 30th ed

Appendix C. (Continued)

NOTE 1: Routine QC strains listed in Tables 2A through 2J (in "Routine QC Recommendations" boxes at the top of each page) are tested regularly (ie, daily or weekly) to ensure the test system is working and produces results that fall within specified ranges listed in M100. The routine QC strains recommended in this document should be included if a laboratory performs CLSI reference disk diffusion or MIC testing as described herein. For commercial test systems, manufacturer's recommendations should be followed for all QC procedures. Other QC strains are used to assess particular characteristics of a test or test system in select situations or may represent alternative QC strains. For example, *H. influenzae* ATCC® 10211 is more fastidious than *H. influenzae* ATCC® 49247 or *H. influenzae* ATCC® 49766 and is used to ensure HTM can adequately support the growth of patient isolates of *H. influenzae* and *H. parainfluenzae*. QC strains may possess susceptibility or resistance characteristics specific for one or more special tests listed in M02⁵ and M07.⁶ They can be used to assess a new test, for training new personnel, and for competence assessment, and it is not necessary to include them in routine daily or weekly antimicrobial susceptibility testing QC programs.

NOTE 2: Information in boldface type is new or modified since the previous edition.

References for Appendix C

- Rasheed JK, Anderson GJ, Yigit H, et al. Characterization of the extended-spectrum beta-lactamase reference strain, *Klebsiella pneumoniae* K6 (ATCC[®] 700603), which produces the novel enzyme SHV-18. *Antimicrob Agents Chemother*. 2000;44(9):2382-2388.
- Queenan AM, Foleno B, Gownley C, Wira E, Bush K. Effects of inoculum and beta-lactamase activity in AmpC- and extended-spectrum beta-lactamase (ESBL)-producing *Escherichia coli* and *Klebsiella pneumoniae* clinical isolates tested by using NCCLS ESBL methodology. *J Clin Microbiol*. 2004;42(1):269-275.
- Woodford N, Ward ME, Kaufmann ME, et al. Community and hospital spread of *Escherichia coli* producing CTX-M extended-spectrum beta-lactamases in the UK. *J Antimicrob Chemother*. 2004;54(4):735-743.
- Centers for Disease Control and Prevention. CDC & FDA Antibiotic Resistance Isolate Bank. https://wwwn.cdc.gov/arisolatebank/. Accessed 3 December 2019.
- ⁵ CLSI. *Performance Standards for Antimicrobial Disk Susceptibility Tests*. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- 6 CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.

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Appendix D. Anaerobe Cumulative Antibiogram¹

NOTE: Isolates collected from selected US hospitals from 1 January 2013 to 31 December 2016.^a

D1. Bacteroides spp. and Parabacteroides spp.

Anaerobic Organisms	Number of Strains	Amnicillin-		Number of Strains	ili o cari	tazobactam	Number of Strains	19170	CEIOXIIII	Number of Strains		Ertapenem	Number of Strains		Imipenem	Number of Strains		Meropenem
Percent susceptible (%S) and percent resistant (%R) ^b		%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 ^c	84°	16 ^c	49	100	0	236	95	1
B. vulgatus	20°	45°	15°	168	92	0	153	73	14	_	_	_	35	97	0	171	96	4
B. uniformis	19°	84°	0c	78	96	0	72	85	10	_	_	_	19°	100°	0°	93	100	0
Parabacteroides distasonis	27°	59°	19°	92	95	1	82	29	43	_	_	_	26°	100°	0	119	97	2

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Appendix D. (Continued)

D1. Bacteroides spp. and Parabacteroides spp. (Continued)

Anaerobic Organisms	Number of Strains		Clindamycin	Number of Strains	Moxifloxacin		Number of Strains	Metronidazole	
Percent susceptible (%S) and percent resistant (%R) ^b		%S	%R		%S	%R		%S	%R
Breakpoints, μg/mL		≤2	≥8		≤2	≥8		≤8	≥32
B. fragilis	1013	26	22	256	61	32	1140	100	0
B. thetaiotaomicron	328	28	49	70	54	36	322	100	0
B. ovatus	207	46	51	59	41	25	236	100	0
B. vulgatus	171	53	46	29°	31°	45°	186	100	0
B. uniformis	87	45	48	25°	48°	40°	89	100	0
Parabacteroides distasonis	108	43	44	37	62	35	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 M39¹ recommendation of 30 isolates.

Reference for D1

¹ 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.

Appendix D. (Continued)

NOTE: Isolates collected from selected US hospitals from 1 January 2013 to 31 December 2016.^a

D2. Anaerobic Organisms Other Than Bacteroides spp. and Parabacteroides spp.

D2. Anaerobic Organis	ilis Othe	i illafi	Dacter VIDE	so spp. ai	iu Paiab	acteroide	s spp.								
Anaerobic Organisms	Number of Strains	s:11:5:20 A	Ampicinin- sulbactam	Number of Strains	Piperacillin-	tazobactam	Number of Strains	lmipenem		Number of Strains	•	мегорепет	Number of Strains	:	Penicillin
Percent susceptible (%S) and percent resistant (%R) ^b		%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 ^c	97°	3 ^c	63	100	0	29 ^c	100	0	92	98	0	63	100	0
Fusobacterium spp.	20°	100 ^c	0°	55	96	2	75	95	4	20°	100 ^c	0°	_d	_d	_d
Anaerobic gram- positive cocci ^e	_d	_d	_d	1853	99	1	134	99	0	1647	100	0	1647	100	0
Cutibacterium (formerly Propionibacterium) acnes ^f	_d	_d	_d	18°	100°	Oc	17°	94 ^c	O _q	_d	_d	_d	_d	_d	_d
Clostridium perfringens	15 ^c	100°	0	410	100	0	23 ^c	100°	0°	417	100	0	402	90	4
Clostridioides (formerly Clostridium) difficile ⁹	76	99	0	542	93	0	480	69	4	609	99	0	533	6	37
Other Clostridium spp.	_d	_d	_d	439	94	1	71	99	0	390	100	0	390	69	13

Appendix D. (Continued)

D2. Anaerobic Organisms Other Than Bacteroides spp. and Parabacteroides spp. (Continued)

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

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Appendix D. (Continued)

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 M39¹ 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

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.

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M100, 30th ed

Appendix E. Dosage Regimens Used to Establish Susceptible or Susceptible-Dose Dependent Breakpoints

The evolving science of pharmacokinetics-pharmacodynamics has become increasingly important in recent years in determining minimal inhibitory concentration (MIC) breakpoints. Recently approved susceptible or susceptible-dose dependent (SDD) breakpoints for a number of agents have been based on a specific dosage regimen(s); these dosage regimens are listed in the table below. Proper application of the breakpoints necessitates drug exposure at the site of infection that corresponds to or exceeds the expected systemic drug exposure at the dose listed in adult patients with normal renal function. This information should be shared with pharmacists, infectious diseases staff, and others making dosing recommendations for the institution.

	Breakpoints and Interpretive Categories								
		Susceptible		SDD					
Antimicrobial Agent	MIC	Dose	MIC	Dose					
Table 2A. Enterobacterales									
Azithromycin (Salmonella enterica	≤16 µg/mL	500 mg administered daily	N/A						
ser. Typhi)									
Aztreonam	≤4 µg/mL	1 g administered every 8 h	N/A						
Cefazolin	≤2 µg/mL	2 g administered every 8 h	N/A						
Ceftaroline	≤0.5 µg/mL	600 mg administered every 12 h	N/A						
Cefepime	≤2 µg/mL	1 g administered every 12 h	4 μg/mL	1 g administered every 8 h or					
				2 g administered every 12 h					
			8 μg/mL	2 g administered every 8 h					
			or	(Because it is not possible to correlate specific zone					
			zone diameter: 19–24 mm	diameters with specific MICs, an isolate with a zone diameter in the SDD range should be treated as if it might					
Cefiderocol	≤4 µg/mL	2 g every 8 h administered over 3 h	N/A	be an MIC of 8 μg/mL.)					
Cefotaxime	≤1 µg/mL	1 g administered every 8 h	N/A						
Ceftriaxone	≤1 μg/mL	1 g administered every 6 h	N/A						
Cefoxitin	≤8 µg/mL	8 g per day (eg, 2 g administered every 6 h)	N/A						
Cefuroxime	≤8 µg/mL	1.5 q administered every 8 h	N/A						
Ceftazidime	≤4 µg/mL	1 g administered every 8 h	N/A						
Ceftazidime-avibactam	≤8/4 μg/mL	2.5 g (2 g ceftazidime + 0.5 g avibactam) every 8 h	N/A						
Condition dybaotam	= 0/+ µg/IIIL	administered over 2 h	14/74						
Ceftizoxime	≤1 µg/mL	1 g administered every 12 h	N/A						
Ceftolozane-tazobactam	≤ 2/4 µg/mL	1.5 g administered every 8 h	N/A						
Ciprofloxacin	≤0.25 µg/mL	400 mg IV or 500 mg orally administered every 12 h	N/A						
Colistin or polymyxin B	≤2 μg/mL ^a	See International Consensus Guidelines ¹ for dosage recommendations.	N/A						
Doripenem	≤1 µg/mL	500 mg administered every 8 h	N/A						
Ertapenem	≤0.5 µg/mL	1 g administered every 24 h	N/A						
Imipenem	≤1 µg/mL	500 mg administered every 6 h or 1 g every 8 h	N/A						
Levofloxacin	≤0.5 µg/mL	750 mg administered every 24 h	N/A						
Meropenem	≤1 µg/mL	1 g administered every 8 h	N/A						
Meropenem-vaborbactam	≤4/8 μg/mL	4 g (2 g meropenem+2 g vaborbactam) every 8 h administered over 3 h	N/A						

Appendix E. (Continued)

· · · · · · · · · · · · · · · · · · ·	Breakpoints and Interpretive Categories									
		Susceptible		SDD						
Antimicrobial Agent	MIC	Dose	MIC	Dose						
Table 2B-1. Pseudomonas aeru	uginosa									
Aztreonam	≤8 µg/mL	1 g administered every 6 h or 2 g every 8 h	N/A							
Cefepime	≤8 µg/mL	1 g administered every 8 h or 2 g every 12 h	N/A							
Cefiderocol	≤4 µg/mL	2 g every 8 h administered over 3 h	N/A							
Ceftazidime	≤8 µg/mL	1 g administered every 6 h or 2 g every 8 h	N/A							
Ceftazidime-avibactam	≥8/4 µg/mL	2.5 g (2 g ceftazidime+0.5 g avibactam) administered every 8 h over 2 h	N/A							
Ciprofloxacin	≤0.5 µg/mL	400 mg IV administered every 8h	N/A							
Colistin or polymyxin B	≤2 µg/mLª	See International Consensus Guidelines ¹ for dosage recommendations	N/A							
Doripenem	≤2 µg/mL	500 mg administered every 8 h	N/A							
Imipenem	≤2 µg/mL	1 g administered every 8 h or 500 mg every 6 h	N/A							
Levofloxacin	≤1 µg/mL	750 mg administered every 24 h	N/A							
Meropenem	≤2 µg/mL	1 g administered every 8 h	N/A							
Piperacillin	≤16 µg/mL	3 g administered every 6 h	N/A							
Piperacillin-tazobactam	≤16/4 µg/mL	3 g administered every 6 h	N/A							
Ticarcillin	≤16 µg/mL	3 g administered every 6 h	N/A							
Ticarcillin-clavulanate ≤16/2 μg/mL		3 g administered every 6 h	N/A							
Table 2B-2. Acinetobacter spp.										
Cefiderocol	≤4 µg/mL	2 g every 8 h administered over 3 h	N/A							
Colistin or polymyxin B	≤2 µg/mLª	See International Consensus Guidelines ¹ for dosage recommendations	N/A							
Doripenem	≤2 µg/mL	500 mg administered every 8 h	N/A							
Imipenem	≤2 µg/mL	500 mg administered every 6 h	N/A							
Meropenem	≤2 µg/mL	1 g administered every 8 h or 500 mg every 6 h	N/A							
Table 2B-4. Stenotrophomonas	s maltophilia									
Cefiderocol	≤4 µg/mL	2 g every 8 h administered over 3 h	N/A							
Table 2C. Staphylococcus spp										
Ceftaroline (S. aureus only)	≤1 µg/mL	600 mg administered every 12 h	2–4 μg/mL	600 mg every 8 h administere over 2 h NOTE: For <i>S. aureus</i> only.						
Dalbavancin	≤0.25 µg/mL	1500 mg (single dose) IV administered over 30 minutes or 1000 mg (two does) followed one week later by 500 mg IV administered over 30 minutes (adult patients with creatinine clearance ≥ 30 mL/minute)	N/A							
Oritavancin	≤0.12 µg/mL	1200 mg single IV dose	N/A							
Tedizolid	≤0.5 µg/mL	200 mg administered every 24 h	N/A							
Telavancin	≤0.12 µg/mL	10 mg/kg administered every 24 h	N/A							

Appendix E. (Continued)

	Breakpoints and Interpretive Categories								
		Susceptible		SDD					
Antimicrobial Agent	MIC	Dose	MIC Dose						
Table 2D. Enterococcus spp.									
Dalbavancin	≤0.25 µg/mL	1500 mg (single dose) IV administered over 30 minutes or 1000 mg (two does) followed one week later by 500 mg IV administered over 30 minutes (adult patients with creatinine clearance ≥ 30 mL/minute).	N/A						
Daptomycin E. faecium only	N/A	N/A	≤4 μg/mL	8–12 mg/kg administered every 24 h					
Daptomycin Enterococcus spp. other than E. faecium	≤2 μg/mL	6 mg/kg administered every 24 h	N/A						
Oritavancin	≤0.12 µg/mL	1200 mg single IV dose	N/A						
Tedizolid	≤0.5 µg/mL	200 mg administered every 24 h	N/A						
Telavancin	≤0.25 µg/mL	10 mg/kg administered every 24 h	N/A						
Table 2E. Haemophilus influenza	e and Haemophilus								
Ceftaroline	≤0.5 µg/mL	600 mg administered every 12 h	N/A						
Table 2G. Streptococcus pneumo	oniae	· · · ·							
Ceftaroline (nonmeningitis)	≤0.5 µg/mL	600 mg administered every 12 h	N/A						
Penicillin (nonmeningitis)	≤2 µg/mL	2 million units administered every 4 h (12 million units per day)	N/A						
Penicillin parenteral (meningitis)	≤0.06 µg/mL	3 million units administered every 4 h	N/A						
Table 2H-1. Streptococcus spp. ß	B-Hemolytic Group								
Ceftaroline	≤0.5 µg/mL	600 mg administered every 12 h	N/A						
Dalbavancin	≤0.25 µg/mL	1500 mg (single dose) IV administered over 30 minutes or 1000 mg (two doses) followed one week later by 500 mg IV administered over 30 minutes (adult patients with creatinine clearance ≥ 30 mL/minute).	N/A						
Oritavancin	≤0.25 µg/mL	1200 mg single IV dose	N/A						
Tedizolid	≤0.25 µg/mL	200 mg administered every 24 h	N/A						
Telavancin	≤0.12 µg/mL	10 mg/kg administered every 24 h	N/A						
Table 2H-2. Streptococcus spp. V	/iridans Group								
Dalbavancin	≤0.25 µg/mL	1500 mg (single dose) IV administered over 30 minutes or 1000 mg (two doses) followed one week later by 500 mg IV administered over 30 minutes (adult patients with creatinine clearance ≥ 30 mL/minute).	N/A						
Oritavancin	≤0.25 µg/mL	1200 mg single IV dose	N/A						
Tedizolid	≤0.5 µg/mL	200 mg administered every 24 h	N/A						
Telavancin	≤0.06 µg/mL	10 mg/kg administered every 24 h	N/A						

Abbreviations: IV, intravenous; MIC, minimal inhibitory concentration; N/A, not applicable; SDD, susceptible-dose dependent.

Appendix E. (Continued)

Footnote

a. MIC ≤2 µg/mL for colistin and polymyxin B corresponds to intermediate category.

NOTE: Information in boldface type is new or modified since the previous edition.

Reference for Appendix E

Tsuji BT, Pogue JM, Zavaxcki AP, et al. International consensus guidelines for the optimal use of the polymyxins: endorsed by the American College of Clinical Pharmacy (ACCP), European Society of Clinical Microbiology and Infectious Diseases (ESCMID), Infectious Diseases Society of America (IDSA), International Society for Anti-Infective Pharmacology (ISAP), Society of Critical Care Medicine (SCCM), and Society of Infectious Diseases Pharmacists (SIDP). *Pharmacotherapy*. 2019;39(1):10-39.

Appendix F
Susceptible-Dose Dependent
Interpretive Category

Appendix F. Susceptible-Dose Dependent Interpretive Category

Abbreviations for Appendix F

AST antimicrobial susceptibility testing
FDA US Food and Drug Administration
MIC minimal inhibitory concentration
QC quality control

SDD susceptible-dose dependent

Susceptible-dose dependent (SDD) is recommended instead of "intermediate" for several drug and organism combinations for which there are multiple approved or routinely used dosing options:

Enterobacterales: cefepimeStaphylococcus aureus: ceftaroline

Enterococcus faecium: daptomycin

SDD highlights the option of using higher doses or alternative dosing regimens by which to achieve a higher dose exposure for the treatment of infections caused by isolates when the minimal inhibitory concentration (MIC) or the zone diameter is in the SDD range.

What does SDD mean?

SDD is a category defined by a breakpoint that implies that susceptibility of an isolate depends on the dosing regimen that is used in the patient. To achieve levels that are likely to be clinically effective against isolates for which the susceptibility testing results (either MICs or zone diameters) are in the SDD category, it is necessary to use a dosing regimen (ie, higher doses, more frequent doses, or both) that results in higher drug exposure than that achieved with the dose that was used to establish the susceptible breakpoint. Consideration should be given to the maximum, literature-supported dosage regimens, because higher exposure gives the highest probability of adequate coverage of an SDD isolate. Appendix E lists the doses used when establishing SDD categories. The drug label should be consulted for recommended doses and adjustment for organ function.

NOTE: The concept of SDD has been included within the intermediate category definition for antimicrobial agents. However, this is often overlooked or not understood by clinicians and microbiologists when an intermediate result is reported. The SDD category may be assigned when doses well above those used to calculate the susceptible breakpoint are supported by the literature, widely used clinically, and/or approved and for which sufficient data to justify the designation exist and have been reviewed. When the intermediate category is used, its definition remains unchanged.

Why is SDD being used now?

- There is a growing need to refine susceptibility reporting to maximize clinicians' use of available drugs.
- Intermediate too often means "resistant" to clinicians because they do not appreciate the full definition of "intermediate."
- SDD is more specific and conveys what we know—a higher dose can be considered for isolates with MICs (or zones **of inhibition**) that fall in this interpretive category.

Appendix F. (Continued)

- SDD is already well established for use in antifungal susceptibility testing.
- Antibiotic stewardship programs, which emphasize dosage regimen and duration of therapy options, are increasing awareness of appropriate use of antibiotics. Personnel from these programs should be able to describe the significance to clinicians of an SDD result.

How should this change be implemented?

- Meet with the appropriate practitioners at your institution (eg, members of the antimicrobial stewardship team, infectious diseases staff, pathology group, pharmacy) to explain SDD and determine a plan for implementation, if appropriate.
- Talk to the manufacturer of your antimicrobial susceptibility testing (AST) device to determine how to implement reporting SDD on your device.
 - NOTE: Because the US Food and Drug Administration (FDA) does not yet recognize the SDD interpretive category and commercial manufacturers must
 use FDA breakpoints, the manufacturer cannot adopt the CLSI SDD breakpoints. However, for most systems, you can manually change the breakpoints
 and implement, following a verification study.
- Work with your laboratory information system staff to report "SDD" or dose ("D") when MICs or zone **diameters** are in the SDD range. Some laboratory information systems may handle only a single character and use of "D" for "dose" may be appropriate. Ideally, this could be translated to SDD on the final patient report. Regardless of approach, make certain that SDD will be transmitted to the hospital information system and appropriately displayed on reports viewed by clinicians.
- Distribute user-specific educational materials to laboratory staff and clinicians receiving AST results from your laboratory. Examples of these materials can be
 found on the CLSI Subcommittee on Antimicrobial Susceptibility Testing webpage at www.clsi.org.

Additional Questions and Answers:

- 1. Q: Does CLSI recommend a comment to be reported with the new SDD breakpoints?
 - A: If a laboratory chooses to report a comment explaining the SDD range, CLSI recommends the following: "The interpretive criterion for susceptible is based on a dosage regimen of [dose] (refer to Appendix E). The interpretive criterion for SDD is based on dosage regimens that result in higher antimicrobial exposure, either higher doses or more frequent doses, or both."
- 2. Q: Will all intermediate ranges become SDD?
 - A: No, the SDD category will be implemented for drug and organism combinations only when there is sufficient evidence to suggest alternative approved dosage regimens may be appropriate for organisms that have MICs or zone diameters between the susceptible and resistant categories.
- 3. Q: Will SDD be applied to other antimicrobial agents?
 - A: CLSI will examine the SDD category possibility for additional drug and organism combinations for which multiple dosing options exist and have been well studied.

Appendix F. (Continued)

- 4. Q: How do we perform a verification study before implementing the new breakpoints on our AST device?
 - A: Guidelines for performance of such a verification study are available (see CLSI document M521).2
- 5. Q: Does SDD apply to all patients and specimen types (eg, pediatric, geriatric, immunosuppressed)?
 - A: Yes, in terms of laboratory reporting. Clinicians must decide how to use an SDD result for a specific patient while considering all other clinical and physiological parameters for that patient.
- 6. Q: Is any special QC needed once the SDD breakpoints are implemented?
 - A: No, currently recommended routine QC is sufficient.
- 7. Q: Will it be necessary to report SDD on proficiency testing survey samples?
 - A: Sponsors of proficiency testing surveys are aware of the difficulties encountered by laboratories in implementing newer CLSI breakpoints. It is highly unlikely that there will be a mandate to report SDD in the near future, but it would be best to check with your proficiency testing survey provider.
- 8. Q: If we can implement the revised breakpoints but cannot facilitate reporting of SDD, can we report "intermediate" instead of SDD?
 - A: A decision related to this question should be made following consultation with your laboratory director, antibiotic stewardship team (if available), infectious diseases practitioners, pharmacists, and infection prevention practitioners.
- 9. Q: If we can implement the revised breakpoints but cannot facilitate reporting of SDD, can we report an MIC or zone diameter without an interpretation?
 - A: A zone diameter should never be reported without an interpretation because there is a high risk of misinterpretation of this value, which poses patient safety issues. There is a lesser danger of reporting an MIC without an interpretation, but this should not be done without an accompanying qualifying comment. See answer to question 8, above.
- 10. Q: What does the dosing information that is given with breakpoints mean?
 - A: The evolving science of pharmacokinetics-pharmacodynamics has become increasingly important in recent years in determining MIC breakpoints. Recently approved susceptible or SDD breakpoints for a number of agents have been based on a specific dosage regimen(s); these dosage regimens are listed in Appendix E. Proper application of the breakpoints necessitates drug exposure at the site of infection that corresponds to or exceeds the expected systemic drug exposure, at the dose listed, in adult patients with normal renal function. This information should be shared with pharmacists, infectious diseases staff, and others making dosing recommendations for the institution.

Appendix F. (Continued)

NOTE: Information in boldface type is new or modified since the previous edition.

References for Appendix F

- ¹ CLSI. *Verification of Commercial Microbial Identification and Antimicrobial Susceptibility Testing Systems*. 1st ed. CLSI guideline M52. Wayne, PA: Clinical and Laboratory Standards Institute; 2015.
- Patel J, Sharp S, Novak-Weekley S. Verification of antimicrobial susceptibility testing methods: a practical approach. *Clin Microbiol Newslett.* 2013;35(13):103-109.

Appendix G
Epidemiological Cutoff Values

Appendix G. Epidemiological Cutoff Values

Abbreviations for Appendix G

ECV epidemiological cutoff value MIC minimal inhibitory concentration

NWT non-wild-type WT wild-type

G1 Defining Epidemiological Cutoff Values

G1.1 Definitions

epidemiological cutoff value (ECV) – the minimal inhibitory concentration (MIC) or zone diameter value that separates microbial populations into those with and without phenotypically detectable resistance (non-wild-type [NWT] or wild-type [WT], respectively). The ECV defines the highest MIC or smallest zone diameter for the WT population of isolates.

EXAMPLE:

	ECVs						
Interpretive Category	MIC, μg/mL	Zone Diameter, mm					
Wild-type	≤4	≥20					
Non-wild-type	≥8	≤19					

- wild-type (WT) an interpretive category defined by an ECV that describes the microbial population with no phenotypically detectable mechanisms of resistance or reduced susceptibility for the antimicrobial (antifungal) agent being evaluated.
- non-wild-type (NWT) an interpretive category defined by an ECV that describes the microbial population with phenotypically detectable mechanisms of resistance and reduced susceptibility for the antimicrobial (antifungal) agent being evaluated.

G1.2 Epidemiological Cutoff Values vs Clinical Breakpoints

ECVs are based on *in vitro* data only, using MIC or zone diameter distributions. ECVs are <u>not</u> clinical breakpoints, and the clinical relevance of ECVs for a particular patient has not yet been identified or approved by CLSI or any regulatory agency.

By contrast, clinical breakpoints are established using MIC distributions, pharmacokinetic-pharmacodynamic data, and clinical outcome data, when available (as described in CLSI document M23¹).

"Caution": Zone diameter (disk diffusion) and MIC values for which ECVs are defined are not to be interpreted or reported as susceptible, intermediate, or resistant, but rather as WT or NWT. The ECVs should not be used as clinical breakpoints.

Appendix G. (Continued)

G1.3 Establishing Epidemiological Cutoff Values

ECVs are determined by collecting and merging MIC distribution data obtained by testing **microbes** from a variety of sources and then applying statistical techniques for estimating the MIC at the upper end of the WT distribution. Subsequently, corresponding zone diameter data from disk diffusion testing are examined and a disk diffusion ECV is determined, when appropriate. To ensure reliability, ECVs are estimated while accounting for both biological (strain-to-strain) variation and MIC/disk assay variation within and between laboratories. They are based on the assumption that the WT distribution of a particular antimicrobial agent/organism combination does not vary geographically or over time.

Several conditions must be fulfilled to generate reliable ECVs. The most important are:

- An ECV can be determined only within a single species for a single agent because of the genetic diversity between species within a genus.
- All MIC values included in the dataset must have been determined using a standard reference method (eg, the CLSI MIC broth dilution method as described in M07,² which is also the method outlined in an international reference standard³). Similarly, the standard reference disk diffusion method as described in M02⁴ must be used when zone diameter ECVs are defined.
- Data must be sourced from at least three separate laboratories and at least 100 unique isolates must be included in the merged dataset.
- MIC values contributed from an individual laboratory dataset should be "on scale" (ie, the MIC is not below the lowest or above the highest concentration tested), whenever possible. This is particularly important for MICs of the presumptive WT strains. Before merging data from individual laboratories, the MIC distribution from each laboratory must be inspected, and if the lowest concentration tested is also the mode, the data must be excluded.
 - Once acceptable data are merged, there are several methods that can be used to estimate the ECV.
 - Visual inspection is the simplest method and is generally acceptable for MIC distributions when there is clear separation of WT and NWT strains.
 When there is obvious overlap between WT and NWT strains, visual inspection is too subjective to set a reliable ECV.
 - Statistical methods are preferred because they remove potential observer bias from the estimation. The two most widely referenced statistical methods are those described by Turnidge et al.⁵ and by Kronvall.⁶
 - Establishment of ECVs from MIC distributions may be supplemented with molecular tests for known resistance genes. The detection of a resistance gene per se in strains with MICs at or below the ECV does not necessarily contradict the choice of ECV, unless it can be accompanied by evidence that the gene is being expressed. In such cases, the ECV may need to be reassessed.

G1.4 Epidemiological Cutoff Value Use by the Medical Microbiology Laboratory

The need for testing and interpreting drug and organism combinations with an ECV but no clinical breakpoint must be discussed with appropriate clinical specialists (eg, antibiotic stewardship, infectious diseases, and pharmacy). While ECVs do not predict clinical outcome, laboratories may consider noting WT or NWT MIC (or zone diameter) interpretations on laboratory reports. Many physicians may choose not to consider using antimicrobial agents with an NWT interpretation, if other therapeutic options are available. However, it is critical that laboratories refrain from reporting report WT as susceptible, or NWT as resistant, as there are insufficient clinical data to support this practice. ECVs may be used to signal the emergence of resistance, although this application for ECVs is best suited to public health laboratories and surveillance studies.

Appendix G. (Continued)

References for G1

- ¹ CLSI. Development of In Vitro Susceptibility Testing Criteria and Quality Control Parameters. 5th ed. CLSI guideline M23. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- CLSI. Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ³ ISO. Clinical laboratory testing and in vitro diagnostic test systems Susceptibility testing of infectious agents and evaluation of performance of antimicrobial susceptibility test devices Part 1: Reference method for testing the in vitro activity of antimicrobial agents against rapidly growing aerobic bacteria involved in infectious diseases. ISO 20776-1. Geneva, Switzerland: International Organization for Standardization; 2006.
- 4 CLSI. Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed. CLSI standard M02. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- ⁵ Turnidge J, Kahlmeter G, Kronvall G. Statistical characterisation of bacterial wild-type MIC value distributions and the determination of epidemiological cut-off values. *Clin Microbiol Infect.* 2006;12(5):418-425.
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Appendix G. (Continued)

G2 Epidemiological Cutoff Value Tables

"Caution": Zone diameter (disk diffusion) and MIC values for which ECVs are defined are not to be interpreted or reported as susceptible, intermediate, or resistant, but rather as WT or NWT. The ECVs should not be used as clinical breakpoints.

ECVs listed in Tables G1 and G2 are applicable only to the species indicated. Currently, there are insufficient data to support their use with other species.

Table G1. ECV for Enterobacterales

Antimicrobial	Disk		neter ECV, nm	MIC ECV, μg/mL		
Agent	Content	WT	NWT	WT	NWT	Comments
Azithromycin ¹⁻⁵	15 µg	≥16	≤15	≤8	≥16	For use with Shigella flexneri.
						See Table 2A for azithromycin and Salmonella spp.
	_	-	_	≤16	≥32	For use with Shigella sonnei.
		L				<u> </u>

Abbreviations: ECV, epidemiological cutoff value; MIC, minimal inhibitory concentration; NWT, non-wild-type; WT, wild-type.

NOTE: Information in boldface type is new or modified since the previous edition.

References for Table G1

- ¹ Klontz KC, Singh N. Treatment of drug-resistant Shigella infections. Expert Rev Anti Infect Ther. 2015;13(1):69-80.
- ² Baker KS, Dallman TJ, Ashton PM, et al. Intercontinental dissemination of azithromycin-resistant shigellosis through sexual transmission: a cross-sectional study. *Lancet Infect Dis.* 2015;15(8):913-921.
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- ⁴ Valcanis M, Brown JD, Hazelton B, et al. Outbreak of locally acquired azithromycin-resistant *Shigella flexneri* infection in men who have sex with men. *Pathology*. 2015;47(1):87-88.
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Appendix G Epidemiological Cutoff Values

Appendix G. (Continued)

Table G2. ECVs for Specific Anaerobic Species

	MIC ECV, μg/mL		
Antimicrobial Agent	WT	NWT	Comments
Vancomycin	≤2	≥4	For use with <i>Cutibacterium</i> (formerly <i>Propionibacterium</i>) acnes ¹⁻⁴ and <i>Clostridioides</i> (formerly <i>Clostridium</i>) <i>difficile</i> . ⁵⁻⁷

Abbreviations: ECV, epidemiological cutoff value; MIC, minimal inhibitory concentration; NWT, non-wild-type; WT, wild-type.

References for Table G2

- ¹ Citron DM, Kwok YY, Appleman MD. In vitro activity of oritavancin (LY333328), vancomycin, clindamycin, and metronidazole against *Clostridium perfringens, Propionibacterium acnes,* and anaerobic Gram-positive cocci. *Anaerobe.* 2005;11(1-2):93-95.
- ² Goldstein EJ, Citron DM, Merriam CV, Warren YA, Tyrrell KL, Fernandez HT. In vitro activities of the new semisynthetic glycopeptide telavancin (TD-6424), vancomycin, daptomycin, linezolid, and four comparator agents against anaerobic gram-positive species and *Corynebacterium* spp. *Antimicrob Agents Chemother*. 2004;48(6):2149-2152.
- Oprica C, Nord CE; ESCMID Study Group on Antimicrobial Resistance in Anaerobic Bacteria. European surveillance study on the antibiotic susceptibility of *Propionibacterium acnes. Clin Microbiol Infect.* 2005;11(3):204-213.
- ⁴ Tyrrell KL, Citron DM, Warren YA, Fernandez HT, Merriam CV, Goldstein EJ. In vitro activities of daptomycin, vancomycin, and penicillin against *Clostridium difficile*, *C. perfringens, Finegoldia magna*, and *Propionibacterium acnes*. *Antimicrob Agents Chemother*. 2006;50(8):2728-2731.
- Snydman DR, McDermott LA, Jacobus NV, et al. U.S.-based National Sentinel Surveillance Study for the epidemiology of *Clostridium difficile*-associated diarrheal isolates and their susceptibility to fidaxomicin. *Antimicrob Agents Chemother*. 2015;59(10):6437-6443.
- Goldstein EJ, Citron DM, Tyrrell KL, Merriam CV. Comparative in vitro activities of SMT19969, a new antimicrobial agent, against *Clostridium difficile* and 350 gram-positive and gram-negative aerobic and anaerobic intestinal flora isolates. *Antimicrob Agents Chemother*. 2013;57(10):4872-4876.
- Goldstein EJ, Babakhani F, Citron DM. Antimicrobial activities of fidazomicin. *Clin Infect Dis.* 2012;55 Suppl 2:S143-8.

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Appendix H. Using Molecular Assays for Resistance Detection

Abbreviations for Appendix H

AST antimicrobial susceptibility testing
ESBL extended-spectrum β-lactamase
MIC minimal inhibitory concentration

MRSA methicillin (oxacillin)-resistant Staphylococcus aureus

A not applicable

PBP2a penicillin-binding protein 2a
VRE vancomycin-resistant enterococci

Antimicrobial resistance and susceptibility are complex, and current *in vitro* methods have been developed to predict a microorganism's response to antibacterial therapy *in vivo*. Standardized phenotypic methods have evolved over many decades, but faster and potentially more reliable nucleic acid— and protein-based methods have been recently developed to detect antimicrobial resistance. The current challenge for medical laboratories is to integrate molecular assays for antimicrobial resistance determinants with conventional antimicrobial susceptibility testing (AST) procedures, sometimes despite an incomplete understanding of test limitations.

The tables in this appendix provide a practical approach for testing and reporting results among medical laboratories that routinely use molecular techniques (with or without a phenotypic test) for detecting antimicrobial resistance. Antimicrobial resistance is genetically complex and based on available data. Molecular methods are often used as a screening tool (eg, methicillin (oxacillin)-resistant Staphylococcus aureus [MRSA] from nasal swabs) or as a rapid adjunct to traditional phenotypic methods (eg, KPC from instrument-flagged blood culture bottles). Interpretation necessitates critical thinking and an understanding of the dynamics between detecting "resistance" determinants and testing phenotypic "susceptibility." Detecting a resistance marker does not necessarily predict therapeutic failure of antimicrobial agents. The gene may be nonfunctional or expressed at clinically insignificant levels. Conversely, the absence of the genetic marker does not necessarily indicate susceptibility, because technical issues may interfere with detection (eg, inhibition of amplification, emergence of genetic variants). In some cases, a molecular approach may be superior to traditional phenotypic methods, such as in the case of low *in vitro* expression, heteroresistance, or poor growth masking higher minimal inhibitory concentrations (MICs). Overall, laboratorians should attempt to apply a consistent approach to molecular-based methods and aim to resolve discordant results with repeat or supplementary testing, by referral to a reference laboratory or by reporting both results in accordance with institutional policies.

As understanding of the molecular mechanisms of antimicrobial resistance continues to develop, more sophisticated approaches to molecular detection of antimicrobial resistance in the medical microbiology laboratory will undoubtedly emerge. The following tables will be updated as needed to ensure the provision of relevant guidance as methods evolve.

Appendix H. (Continued)

Table H1. Strategies for Reporting Methicillin (Oxacillin) Results When Using Molecular and Phenotypic AST Methods for S. aureus

				Resu		10 Phenotypic AST Methods	101 01 001 000	
Indication	Target(s)	Method	Specimen Type	Genotype or Predicted Phenotype	Observed Colony Phenotype (if tested)	Suggestions for Resolution	Consider reporting as ^a :	Comments ^b
Detecting methicillin	PBP2a	Latex agglutination, immuno-	Colony	PBP2a positive	Cefoxitin R	N/A	Methicillin (oxacillin) R	1
(oxacillin)		chromatography		PBP2a negative	Cefoxitin S	N/A	Methicillin (oxacillin) S	1
resistance in S. aureus	tance in			PBP2a positive	Cefoxitin S	Confirm isolate identification, repeat latex agglutination and AST, and consider <i>mecA</i> colony NAAT, if available.	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	1–2
				PBP2a negative	Cefoxitin R	Confirm isolate identification, repeat latex agglutination and AST, and consider <i>mecA</i> colony NAAT, if available.	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	1
	mecA	NAAT, microarray hybridization, ISH		mecA detected	Cefoxitin R	N/A	If tested, report phenotypic result as found (methicillin [oxacillin] R) and consider reporting molecular result per institutional protocol.	3–6
				mecA not detected	Cefoxitin S	N/A	If tested, report phenotypic result as found (methicillin [oxacillin] S) and consider reporting molecular result per institutional protocol.	3–6
			mecA detected	Cefoxitin S	Confirm isolate identification, repeat AST, and repeat or perform <i>mecA</i> colony NAAT, if available. If mixed specimen, test isolates individually.	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	2–5, 8–9	
				mecA not detected	Cefoxitin R	Confirm isolate identification, repeat AST, and repeat or perform <i>mecA</i> colony NAAT, if available. If mixed specimen, test isolates individually.	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	3, 7

Appendix H. (Continued)

Table H1. (Continued)

				Resul	ts			
Indication	Target(s)	Method	Specimen Type	Genotype or Predicted Phenotype	Observed Colony Phenotype (if tested)	Suggestions for Resolution	Consider reporting as ^a :	Comments ^b
Detecting methicillin (oxacillin) resistance in S. aureus (Continued)	SCC <i>mec-orfX</i> functional regions <u>only</u>	NAAT	Blood culture broth, surveillance specimen	SCC <i>mec</i> detected	Cefoxitin R	N/A	If tested, report phenotypic result as found (methicillin [oxacillin] R) and consider reporting molecular result per institutional protocol.	3–6
				SCCmec not detected	Cefoxitin S	N/A	If tested, report phenotypic result as found (methicillin [oxacillin] S) and consider reporting molecular result per institutional protocol.	3–6
				SCC <i>mec</i> detected	Cefoxitin S	Confirm isolate identification, repeat AST and consider <i>mecA</i> colony NAAT, if available. If mixed culture, test isolates individually.	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	2, 10
				SCCmec not detected	Cefoxitin R	Confirm isolate identification, repeat AST and consider <i>mecA</i> colony NAAT, if available. If mixed culture, test isolates individually.	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	7, 12

Appendix H. (Continued)

Table H1. (Continued)

				Re	sults			
Indication	Target(s)	Method	Specimen Type	Genotype or Predicted Phenotype	Observed Colony Phenotype (if tested)	Suggestions for Resolution	Consider reporting as ^a :	Comments ^b
Detection of methicillin resistance in S. aureus (Continued)	SCCmec- orfX junctional regions and mecA	NAAT	Blood culture broth, surveillance specimen	SCCmec AND mecA or other target detected	Cefoxitin R	N/A	If tested, report phenotypic result as found (methicillin [oxacillin] R) and consider reporting molecular result per institutional protocol.	3–6
	and/or other targets			SCCmec AND mecA or other target not detected	Cefoxitin S	N/A	If tested, report phenotypic result as found (methicillin [oxacillin] S) and consider reporting molecular result per institutional protocol.	3–6
				SCCmec AND mecA or other target detected	Cefoxitin S	Confirm isolate identification, repeat AST and consider <i>mecA</i> colony NAAT if available. If mixed culture, test isolates individually	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	2
	107			SCCmec AND mecA or other target not detected	Cefoxitin R	Confirm isolate identification, repeat AST and consider <i>mecA</i> colony NAAT if available. If mixed culture, test isolates individually	If discrepancy is not resolved by suggested testing, report as methicillin (oxacillin) R.	3, 11

Abbreviations: AST, antimicrobial susceptibility testing; ISH, *in situ* hybridization; MSSA, methicillin **(oxacillin)**-susceptible *Staphylococcus aureus*; MRSA, methicillin **(oxacillin)**-resistant *S. aureus*; N/A, not applicable; NAAT, nucleic acid amplification test; PBP2a, penicillin-binding protein 2a; PCR, polymerase chain reaction; R, resistant; S, susceptible.

Appendix H. (Continued)

Table H1. (Continued)

Comments

- (1) False-positive and false-negative PBP2a latex bead agglutination results have been observed.
- (2) Rare mecA-positive S. aureus isolates will test susceptible to cefoxitin.^{2,3}
- (3) mecC or mecA variant gene-mediated methicillin (oxacillin) resistance may not be detected by mecA PCR.^{4,5}
- (4) The **simultaneous** presence of *mecA*-positive **Staphylococcus spp.** (other than **S.** aureus) and MSSA may result in false-positive MRSA molecular results.^{6,7}
- (5) Strains harboring unstable SCCmec insertions may lose mecA during culture.8
- (6) Compared with culture, the sensitivity of molecular methods may be higher, while the specificity may be lower.
- (7) Occasional false-negative mecA results have been reported for direct blood culture molecular assays.9
- (8) For ISH assays with a cefoxitin induction step, false-positive mecA results should be rare. 10
- (9) In polymicrobial cultures, the presence of mecA cannot be attributed to a specific isolate.
- (10) Strains harboring an SCCmec remnant lacking the mecA gene (mecA dropout) or mutant mecA allele may test positive in assays that target only SCCmecorfX junctional regions. Laboratories using molecular tests that detect only SCCmec-orfX junctional region targets may consider adding a disclaimer to the report stating the proportion of false-positive results related to mecA dropouts observed in isolates from the patient population served.¹¹
- (11) Multiple SCCmec types exist; depending on the design of the assay, some SCCmec variants may not be detected. 12

Footnotes

- a. Isolates that test as methicillin resistant are also oxacillin resistant, and the term "methicillin R" is synonymous with "oxacillin R."
- b. In addition to the specific possibilities listed in the comments, genotype and/or phenotype discrepancies could arise as a consequence of suboptimal sampling, mixed cultures, emergence of new genotypes or mutations, and/or wild-type reversions of resistance targets.

NOTE: Information in boldface type is new or modified since the previous edition.

Appendix H. (Continued)

Table H1. (Continued)

References for Table H1

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M100, 30th ed.

Appendix H. (Continued)

Table H2. Strategies for Reporting Vancomycin Results When Using Molecular and Phenotypic Antimicrobial Susceptibility Testing Methods for

Enterococcus spp.

				Resu	ilts			
Indication	Target(s)	Method	Specimen Type	Genotype or Predicted Phenotype	Observed Phenotype (if tested)	Suggestions for Resolution	Report as:	Comments
Detection of vancomycin- resistant enterococci	vanA vanB	NAAT or array hybridization technology	Blood culture broth or surveillance cultures	vanA and/or vanB detected	Vancomycin R	N/A	Report phenotypic result as found (if available); consider reporting presence of molecular target per institutional protocol.	1–3
				vanA and/or vanB not detected	Vancomycin S	N/A	Report phenotypic result as found (if available); consider reporting presence of molecular target per institutional protocol.	
				vanA and/or vanB detected	Vancomycin S	Confirm isolate identification to species level (eg, Enterococcus faecalis) and repeat AST. If mixed culture, test isolates individually.	If discrepancy is not resolved by suggested testing, report as vancomycin R.	1–3
				vanA and/or vanB not detected	Vancomycin R	Confirm isolate identification to species level (eg, <i>E. faecalis</i>) and repeat AST. If mixed culture, test isolates individually.	If discrepancy is not resolved by suggested testing, report as vancomycin R.	4

Appendix H. (Continued)

Table H2. (Continued)

				Res	ults			
Indication	Target(s)	Method	Specimen Type	Genotype or Predicted Phenotype	Observed Phenotype (if tested)	Suggestions for Resolution	Report as:	Comments ^a
Detection of vancomycin- resistant enterococci (Continued)	vanA	NAAT	Surveillance cultures	vanA detected	Vancomycin R	N/A	Report phenotypic result as found (if available); consider reporting presence of molecular target per institutional protocol.	1–2
				vanA not detected	Vancomycin S	N/A	Report phenotypic result as found (if available); consider reporting presence of molecular target per institutional protocol.	5
				vanA detected	Vancomycin S	Confirm isolate identification to species level (eg, <i>E. faecalis</i>) and repeat AST. If mixed culture, test isolates individually.	If the discrepancy is not resolved by suggested testing, report as vancomycin R.	1–2
				vanA not detected	Vancomycin R	Confirm isolate identification to species level (eg, <i>E. faecalis</i>) and repeat AST. If mixed culture, test isolates individually.	If the discrepancy is not resolved by suggested testing, report as vancomycin R.	4–5

Abbreviations: AST, antimicrobial susceptibility testing; N/A, not applicable; NAAT, nucleic acid amplification test; R, resistance; S, susceptible; VRE, vancomycin-resistant enterococci.

Comments

- (1) vanA may be present in nonenterococcal species.1
- (2) Vancomycin-variable *Enterococcus faecium* isolates were recently revealed in Canada. They carry wild-type *vanA* but initially test as vancomycin susceptible with a culture-based method. They can convert to a resistant phenotype during vancomycin treatment.^{2,3}
- (3) The *vanB* gene has been found in several commensal nonenterococcal bacteria, which may lead to misclassification of vancomycin-susceptible enterococci as resistant in surveillance cultures containing mixed bacterial species.⁴

Appendix H. (Continued)

Table H2. (Continued)

- (4) Constitutive low-level vancomycin resistance can be detected phenotypically (2–32 µg/mL) from the presence of *vanC*, an intrinsic resistance characteristic of *Enterococcus gallinarum (vanC1)* and *Enterococcus casseliflavus (vanC2–C4)*.⁵
- (5) Targeting vanA only may miss regional vanB-carrying VRE.6

Footnote

a. In addition to the specific possibilities referenced in the comments, genotype and/or phenotype discrepancies could arise as a consequence of suboptimal sampling, mixed cultures, emergence of new genotypes, or mutations and/or wild-type reversions of resistance targets.

References for Table H2

- Patel R. Enterococcal-type glycopeptide resistance genes in non-enterococcal organisms. FEMS Microbiol Lett. 2000;185(1):1-7.
- Gagnon S, Lévesque S, Lefebvre B, Bourgault AM, Labbé AC, Roger M. vanA-containing Enterococcus faecium susceptible to vancomycin and teicoplanin because of major nucleotide deletions in Tn1546. J Antimicrob Chemother. 2011;66(12):2758–2762.
- Thaker MN, Kalan L, Waglechner N, et al. Vancomycin-variable enterococci can give rise to constitutive resistance during antibiotic therapy. *Antimicrob Agents Chemother*. 2015;59(3):1405-1410.
- ⁴ Ballard SA, Grabsch EA, Johnson PD, Grayson ML. Comparison of three PCR primer sets for identification of *vanB* gene carriage in feces and correlation with carriage of vancomycin-resistant enterococci: interference by *vanB*-containing anaerobic bacilli. *Antimicrob Agents Chemother*. 2005;49(1):77-81.
- ⁵ Courvalin P. Vancomycin resistance in gram-positive cocci. *Clin Infect Dis.* 2006;42(suppl):S25-S34.
- Nebreda T, Oteo J. Aldea C, et al. Hospital dissemination of a clonal complex 17 vanB2-containing Enterococcus faecium. J Antimicrob Chemother. 2007;59(4):806-807.

Appendix H. (Continued)

Table H3. Reporting Results From Extended-Spectrum β-Lactamase Resistance and Carbapenemase Molecular Tests for Enterobacterales

				F	Results			
Indication	Target(s)	Method	Specimen Type	Molecular Target Results	Observed Phenotype (if tested)	Suggestions for Resolution	Report as:	Comments ^a
Detection of ESBL resistance in Enterobacterales (in an isolate susceptible to all carbapenems)	ESBL type CTX-M, SHV, TEM	NAAT, microarray	Colony, blood culture	Detection of any ESBL target	R to all 3rd- and 4th- generation cephalosporins tested (eg, ceftriaxone R, cefotaxime R, ceftazidime R, cefepime R)	N/A	Report phenotypic results as found (if available); consider reporting presence of molecular target per institutional protocol.	1–12
				Detection of any ESBL target	S to all 3rd- and 4th- generation cephalosporins tested (eg, ceftriaxone S, cefotaxime S, ceftazidime S, cefepime S)	Repeat molecular and phenotypic tests. If blood culture, check for mixed culture. If mixed, test isolates individually and report phenotypic results as found.	If the discrepancy is not resolved, repeat AST should be performed using a reference method, and the conflicting genotypic and phenotypic testing results should both be reported.	1–12
				Detection of CTX-M ESBL target	Variable resistance to 3rd- and 4th- generation cephalosporins (eg, ceftriaxone R, cefotaxime R, ceftazidime R or S, cefepime R or S)	Expected phenotype for some CTX-M strains. Check cefepime using a reference method if S.	Report phenotypic results as found, including reference cefepime result; consider reporting presence of molecular target per institutional protocol.	1–12
				Detection of TEM or SHV ESBL target	Variable resistance to 3rd- and 4th- generation cephalosporins (eg, ceftriaxone R or S, cefotaxime R or S, ceftazidime R or S, cefepime R or S).	Expected phenotype for some <i>TEM/SHV</i> strains. Check cefepime using a reference method if S.	Report phenotypic results as found, including reference cefepime result; consider reporting presence of molecular target per institutional protocol.	1–12

Appendix H. (Continued)

Table H3. (Continued)

				R	esults			
Indication	Target(s)	Method	Specimen Type	Molecular Target Results	Observed Phenotype (if tested)	Suggestions for Resolution	Report as:	Comments ^a
Detection of ESBL resistance in Enterobacterales (in an isolate susceptible to all carbapenems) (Continued)				No detection of ESBL targets	Resistance to 3rd- generation cephalosporins and variable resistance to 4th-generation cephalosporins (eg, ceftriaxone R, cefotaxime R, ceftazidime R, cefepime R or S)	Likely non-tested broad spectrum β-lactamase (eg, AmpC, carbapenemase, or other ESBL); consider repeating molecular tests and checking cefepime using reference method if S.	Report phenotypic results as found, including reference cefepime result if tested.	1–12
Detection of carbapenem resistance in Enterobacterales	KPC, OXA- 48-like, VIM, NDM, or IMP	NAAT, microarray	Colony, blood culture	Detection of any tested carbapenemase target Detection of any tested carbapenemase target	Resistance to all carbapenems (eg, meropenem R, imipenem R, doripenem R) Susceptible to all carbapenems except ertapenem (variable) (eg, meropenem S, imipenem S, doripenem R or S)	Repeat molecular and phenotypic tests. If blood culture, check for mixed culture. If mixed, test isolates individually and report phenotypic results as found; consider a phenotypic test for carbapenemase activity (such as CarbaNP or mCIM).	Report phenotypic results as found (if available); consider reporting presence of molecular target per institutional protocol. If the discrepancy is not resolved, repeat AST should be performed using a reference method and the conflicting genotypic and phenotypic testing results should both be reported along with a comment advising caution; current clinical and laboratory evidence is insufficient to conclude whether carbapenem monotherapy of carbapenemase-carrying strains with an MIC in the S range will be effective, or whether the molecular assays are completely accurate.	1–4, 12–14

Appendix H. (Continued)

Table H3. (Continued)

				Res	sults			
			Specimen	Molecular	Observed Phenotype (if	Suggestions for		
Indication	Target(s)	Method	Туре	Target Results	tested)	Resolution	Report as:	Comments ^a
Detection of carbapenem resistance in Enterobacterales (Continued)	KPC, OXA-48- like, VIM, NDM, or IMP	NAAT, microarray	Colony, blood culture	No detection of tested carbapenemase targets	Susceptible to all carbapenems except ertapenem (eg, meropenem S, imipenem S, doripenem R)	Likely ESBL/AmpC and porin alteration, especially for Enterobacter spp.; consider a phenotypic test for carbapenemase activity (eg, CarbaNP or mCIM); carbapenemase unlikely if negative, although rare carbapenemases (eg, GES-types, are still possible).	If carbapenemase activity is detected, repeat AST should be performed using a reference method, and the conflicting genotypic and phenotypic testing results should both be reported along with a comment advising caution; current clinical and laboratory evidence is insufficient to conclude whether carbapenem monotherapy of carbapenemase-carrying strains with an MIC in the susceptible range will be effective or whether the molecular assays are completely accurate. Otherwise report phenotypic results as found.	1–4, 12–15

Appendix H. (Continued)

Table H3. (Continued)

				Re	esults			
Indication	Target(s)	Method	Specimen Type	Molecular Target Results	Observed Phenotype (if tested)	Suggestions for Resolution	Report as:	Comments ^a
Detection of carbapenem resistance in Enterobacterales (Continued)	KPC, OXA-48- like, VIM, NDM, or IMP	NAAT, microarray	Colony, blood culture	No detection of tested carbapenemase targets	Resistance to any carbapenems except ertapenem (eg, meropenem R, imipenem R, doripenem R, ertapenem R or S)	Possible other carbapenemase. If blood culture, check for mixed culture. If mixed, test isolates individually and report as found; consider repeating molecular and AST and performing a phenotypic test for carbapenemase activity (eg, CarbaNP or mCIM).	If carbapenemase activity is detected, repeat AST should be performed using a reference method, and the conflicting genotypic and phenotypic testing results should both be reported along with a comment advising caution; current clinical and laboratory evidence is insufficient to conclude whether carbapenem monotherapy of carbapenemase-carrying strains with an MIC in the S range will be effective or whether the molecular assays are completely accurate. Otherwise report phenotypic results as found.	1–4, 12–16

Abbreviations: AST, antimicrobial susceptibility testing; ESBL, extended-spectrum β-lactamase; mCIM, modified carbapenem inactivation method; MIC, minimal inhibitory concentration; N/A, not applicable; NAAT, nucleic acid amplification test; R, resistant; S, susceptible.

Comments

- (1) Multiple β-lactamases may be carried by individual bacterial isolates. Most carbapenemase-producing bacteria are resistant to 3rd- and 4th-generation cephalosporins, although bacteria with OXA-48 enzymes may not be unless they co-produce an ESBL or AmpC enzyme.
- (2) Molecular assays can detect the presence of specific β-lactamase genes but cannot exclude the presence of other β-lactamase genes or resistance mechanisms, or novel variants with changes in primer or probe annealing sites. Therefore, phenotypic resistance should always be reported.
- (3) Isolates with phenotypic susceptibility despite the presence of a resistance determinant may indicate the potential for resistance to emerge during therapy.

Appendix H. (Continued)

Table H3. (Continued)

- (4) These are provisional guidelines based on general principles; however, the performance characteristics of many individual research use-only assays are presently unknown.
- (5) Susceptibility of TEM/SHV-carrying strains to β-lactam combinations is variable.
- (6) Susceptibility of ESBL-carrying strains to cefepime is variable.
- (7) Susceptibility of ESBL-carrying strains to β-lactam combination agents is variable.
- (8) Some strains carrying CTX-M ESBLs remain susceptible to ceftazidime.
- (9) Some strains carrying TEM/SHV-derived ESBLs remain susceptible to cefotaxime and ceftriaxone.
- (10) Some molecular assays for AmpC may not reliably distinguish between chromosomal and plasmid-encoded genes in some bacterial species.
- (11) Most strains with derepressed AmpC expression remain susceptible to cefepime.
- (12) These recommendations are based on cephalosporin and carbapenem breakpoints in M100.
- (13) The susceptibility to other carbapenems of ertapenem-resistant strains with ESBL or AmpC enzymes and reduced porin expression that do not contain carbapenemase genes or express carbapenemase activity may be reported as measured in phenotypic susceptibility assays.
- (14) Rapid tests for carbapenemase activity (eq. CarbaNP) may not detect OXA-48-like and some other carbapenemases.
- (15) Caution is advised. Current clinical evidence is insufficient to conclude whether carbapenem monotherapy of carbapenemase-carrying strains with an MIC in the susceptible range will be effective.
- (16) Some isolates of **Enterobacterales**, in particular but not exclusively *Morganella* spp., *Proteus* spp., and *Providencia* spp., may exhibit intrinsic low-level resistance to imipenem on a non–carbapenemase-mediated basis.

Footnote

a. In addition to the specific possibilities listed in the comments, genotype and/or phenotype discrepancies could arise as a consequence of mixed cultures, emergence of new genotypes, or mutations and/or wild-type reversions of resistance targets.

NOTE: Information in boldface type is new or modified since the previous edition.

Appendix I. Cefiderocol Broth Preparation and Reading Broth Microdilution Minimal Inhibitory Concentration End Points

Abbreviations for Appendix I

CAMHB cation-adjusted Mueller-Hinton broth

ID-CAMHB iron-depleted cation-adjusted Mueller-Hinton broth negative logarithm of hydrogen ion concentration

I1. Supplements

I1.1 Calcium and Magnesium Stock Solutions

Refer to M07¹ for cation stock solution preparation.

I1.2 Zinc Stock Solution

The steps for preparing zinc stock solution are listed below.

Step	Action	Comment
1	Dissolve 0.29 g ZnSO ₄ · 7H ₂ O in 100 mL deionized water.	This solution contains 10 mg Zn ⁺⁺ /mL.
2	Sterilize the solution by membrane filtration.	
3	Store the solution at 15 to 25°C.	

12. Iron-depleted Cation-adjusted Mueller-Hinton Broth

The steps for preparing iron-depleted cation-adjusted Mueller-Hinton broth (ID-CAMHB) are listed below.²

Step	Action	Comment
1	Prepare the CAMHB.	Follow manufacturer's instructions.
2	Autoclave the media and let cool to room temperature.	
3	Add 100 g chelating resin to 1 L autoclaved CAMHB. ²	Removes cations in the medium- to low-level concentrations (range, 0–0.18 mg/L). ²
4	Stir the solution at room temperature for approximately 2 hours using a magnetic stir bar.	
5	Filter the solution using a 0.2-µm filter.	Removes the resin.
6	Check the pH to determine whether it is 7.3.	If the pH is above 7.3, adjust it using 0.1 M HCI, and if the pH is below 7.3,use 2.5 N NaOH.
7	Add the cation to achieve final concentrations in the following ranges: Ca ⁺⁺ 20–25 mg/L Mg ⁺⁺ 10–12.5 mg/L Zn ⁺⁺ 0.5–1.0 mg/L	The final concentration of Fe ⁺⁺ in ID-CAMHB prepared using this method is ≤0.03 mg/L. Refer to M07 ¹ and the table below for calculating the amount of Ca ⁺⁺ , Mg ⁺⁺ , and Zn ⁺⁺ needed.

Appendix I. (Continued)

I2. ID-CAMBH (Continued)

Step	Action	Comment
8	Check the pH to determine whether it is 7.2–7.4.	If the pH exceeds 7.4, adjust it using 0.1 M HCl. If the pH is below
		7.2, use 2.5 N NaOH.
9	Filter the final product using a 0.2-µm filter.	
10	Store the media at 4 to 8°C for up to 2 months.	

Abbreviations: CAMHB, cation-adjusted Mueller-Hinton broth; ID-CAMHB, iron-depleted cation-adjusted Mueller-Hinton broth.

Example for preparing CAMHB that contains below-detectable concentrations (< 0.0001 mg/L) of Zn⁺⁺ after chelation in step 3²:

Step	Action	Comment
1	Calculate the amount of Zn ⁺⁺ needed using this formula:	For Zn ⁺⁺ , the final amount needed is 0.5–1 mg/L.
	Final amount needed – amount in medium = amount to be added	1 mg/L - 0 mg/L = 1 mg/L
2	Add 0.1 mL Zn ⁺⁺ stock per L to obtain a concentration of 1 mg/L.	1 mg/mL · 0.1 mL = 0.1 mL
3	Proceed with steps 8 and 9 above.	

I3. Determining Broth Microdilution End Points

The steps for reading and interpreting broth microdilution end points for cefiderocol when tested with ID-CAMHB are listed below.

Step	Action	Comment
1	Read the MIC as the lowest concentration of antimicrobial agent that completely inhibits organism growth in the tubes or microdilution wells as detected by the unaided eye.	See step 2 for exceptions. Viewing devices intended to facilitate reading microdilution tests and recording results may be used as long as there is no compromise in the ability to discern growth in the wells.
2	Compare the amount of growth in the wells containing the cefiderocol with the amount of growth in the growth-control well containing ID-CAMHB (no antimicrobial agent).	For a test to be considered valid, acceptable growth (definite turbidity or button) must occur in the growth-control well (see Figure 1). Trailing may occur in some organisms (eg, <i>Acinetobacter</i> spp.) and should be ignored when a tiny button or light or faint turbidity relative to the growth control may be observed. Read the MIC as
3	Interpret the results.	the first well in which growth is significantly reduced (see Figure 2). Refer to the appropriate portion of Tables 2 for breakpoints.

Abbreviations: ID-CAMHB, iron-depleted cation-adjusted Mueller-Hinton broth; MIC, minimal inhibitory concentration.

M100, 30th ed

Appendix I. (Continued)

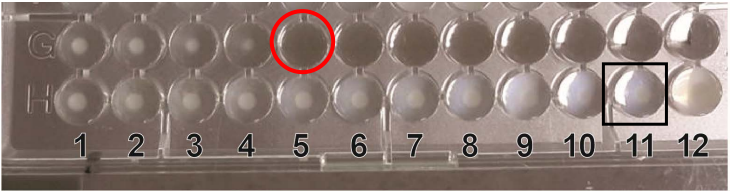


Figure 1. Cefiderocol Test With a Clear End Point. The cefiderocol concentrations in wells G1 to G12 are 0.03 to 64 µg/mL. Row G shows the cefiderocol MIC at 0.5 µg/mL in well G5 (red circle). The growth-control well is H11 (black box). (Courtesy of Meredith M. Hackel, International Health Management Associates. Used with permission.)



Figure 2. Cefiderocol Test With a Trailing End Point. The cefiderocol concentrations in wells A1 to A12 are 0.03 to 64 μg/mL. Row A shows the cefiderocol MIC at 0.25 μg/mL in well A4 (red circle). The growth control well is B11 (black box). (Courtesy of Meredith M. Hackel, International Health Management Associates. Used with permission.)

References for Appendix I

- ¹ CLSI. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07. Wayne, PA: Clinical and Laboratory Standards Institute; 2018.
- Hackel, MA, Tsuji M, Yamano Y, Echols R, Karlowsky JA, Sahm DF. Reproducibility of broth microdilution MICs for the novel siderophore cephalosporin, cefiderocol, determined using iron-depleted cation-adjusted Mueller-Hinton broth. *Diagn Microbiol Infect Dis.* 2019;94(4):321-325.

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Glossary I (Part 1). β-Lactams: Class and Subclass Designations and Generic Names

In the late 1990s, several authorities were consulted to construct the glossary. The intention was to include all agents that appeared in M100, along with related agents available for human use. Since that time, agents have been added to the glossary as they were introduced to CLSI, and they do not need to be FDA cleared to be included. It cannot be assumed that the list is exhaustive, and it should be noted that some agents are no longer available for human use.

Antimicrobial Class	Antimicro	bial Subclass(es)	Agent(s) Included; Generic Name(s)		
Penicillins	Penicillinase-labile	Penicillin	Penicillin		
	penicillins ^a	Aminopenicillins	Amoxicillin		
			Ampicillin		
		Carboxypenicillins	Carbenicillin		
			Ticarcillin		
		Ureidopenicillins	Azlocillin		
			Piperacillin		
	Penicillinase-stable		Cloxacillin		
	penicillins ^b		Dicloxacillin		
			Nafcillin		
			Oxacillin		
	Aminopenicillin		Mecillinam		
β-lactam combination agents			Amoxicillin-clavulanate		
			Ampicillin-sulbactam		
			Aztreonam-avibactam		
			Cefepime-enmetazobactam (4:1)		
			Cefepime-taniborbactam		
			Cefepime-tazobactam (1:1)		
			Cefepime-zidebactam		
			Ceftaroline-avibactam		
			Ceftazidime-avibactam		
			Ceftolozane-tazobactam		
			Imipenem-relebactam		
			Meropenem-nacubactam (1:1)		
			Meropenem-vaborbactam		
			Piperacillin-tazobactam		
			Sulbactam-durlobactam		
			Ticarcillin-clavulanate		
Cephems (parenteral)	Cephalosporins I ^c		Cefazolin		
	·		Cephalothin		
			Cephapirin		
			Cephradine		
	Cephalosporins II ^c		Cefamandole		
	·		Cefonicid		
			Cefuroxime (parenteral)		
	Cephalosporins III ^c		Cefoperazone		
	·		Cefotaxime		
			Ceftazidime		
			Ceftizoxime		
			Cetriaxone		

Glossary I (Part 1). (Continued)

Antimicrobial Class	Antimicrobial Subclass(es)	Agent(s) Included; Generic Name(s)
Cephems (parenteral) (Continued)	Cephalosporins IV ^c	Cefepime
, , , , ,		Cefpirome
	Cephalosporins with anti-MRSA activity	Ceftaroline
		Ceftobiprole
	Cephamycins	Cefmetazole
		Cefotetan
		Cefoxitin
	Oxacephem	Moxalactam
	Siderophore cephalosporin	Cefiderocol
Cephems (oral)	Cephalosporins	Cefaclor
. , ,		Cefadroxil
		Cefdinir
		Cefditoren
		Cefetamet
		Cefixime
		Cefpodoxime
		Cefprozil
		Ceftibuten
		Cefuroxime (oral)
		Cephalexin
		Cephradine
	Carbacephem	Loracarbef
Monobactams		Aztreonam
Penems	Carbapenems	Biapenem
		Doripenem
		Ertapenem
		Imipenem
		Meropenem
		Razupenem
		Tebipenem
	Penems	Faropenem
		Sulopenem

Abbreviations: MRSA, methicillin (oxacillin)-resistant Staphylococcus aureus; FDA, US Food and Drug Administration.

Footnotes

- a. Hydrolyzed by staphylococcal penicillinase.
- b. Not hydrolyzed by staphylococcal penicillinase.
- c. Cephalosporins I, II, III, and IV are sometimes referred to as first-, second-, third-, and fourth-generation cephalosporins, respectively. Cephalosporins III and IV are also referred to as "extended-spectrum cephalosporins." This does not imply activity against extended-spectrum β-lactamase–producing gram-negative bacteria.

NOTE: Information in boldface type is new or modified since the previous edition.

Glossary I (Part 2). Non-β-Lactams: Class and Subclass Designations and Generic Names

In the late 1990s, several authorities were consulted to construct the glossary. The intention was to include all agents that appeared in M100, along with related agents available for human use. Since that time, agents have been added to the glossary as they were introduced to CLSI, and they do not need to be FDA cleared to be included. It cannot be assumed that the list is exhaustive, and it should be noted that some agents are no longer available for human use.

Antimicrobial Class	Antimicrobial Subclass(es)	Agent(s) Included; Generic Name(s)
Aminocyclitols		Spectinomycin
Aminoglycosides		Amikacin
		Gentamicin
		Kanamycin
		Netilmicin
		Plazomicin
		Streptomycin
		Tobramycin
Aminoglycoside-fosfomycin		Amikacin-fosfomycin
Ansamycins	Rifamycins	Rifabutin
		Rifapentine
		Rifampin
		Rifaximin
Antistaphylococcal lysin		Exebacase
Folate pathway antagonists	Dihydrofolate reductase inhibitors	Iclaprim
		Sulfonamides
		Trimethoprim
		Trimethoprim-sulfamethoxazole
	Sulfonamides	Sulfamethoxazole
		Sulfisoxazole
	Combination	Trimethoprim-sulfamethoxazole
Fosfomycins		Fosfomycin
Glycopeptides	Glycopeptide	Vancomycin
	Lipoglycopeptides	Dalbavancin
		Oritavancin
		Teicoplanin
		Telavancin
	Lipoglycodepsipeptide	Ramoplanin
Lincosamides		Clindamycin
		Lincomycin
Lipopeptides		Daptomycin
		Surotomycin
	Polymyxins	Colistin
	,,	Polymyxin B
Macrocyclic lactone		Fidaxomicin
acrosjone isotorio		

Glossary I (Part 2). (Continued)

Antimicrobial Class	Antimicrobial Subclass(es)	Agent(s) Included; Generic Name(s)
Macrolides	•	Azithromycin
		Clarithromycin
		Dirithromycin
		Erythromycin
	Fluoroketolide	Solithromycin
	Ketolides	Nafithromycin
		Telithromycin
Nitroheterocyclics	Nitrofuran	Nitrofurantoin
· · · · · · · · · · · · · · · · · · ·	Nitroimidazoles	Metronidazole
	· · · · · · · · · · · · · · · · · · ·	Secnidazole
		Tinidazole
	Thiazolides	Nitazoxanide
	111111111111111111111111111111111111111	Tizoxanide
Oxazolidinones		Linezolid
O/ALSIIAIITOTIOS		Tedizolid
Peptide	Magainin	Pexiganan
Phenicols	Maganin	Chloramphenicol
THETHOUS		Thiamphenicol
Pleuromutilins		Lefamulin
1 leuromuliins		Retapamulin
Pseudomonic acid		Mupirocin
Quinolones		Cinoxacin
Quinolones		
		Garenoxacin Nalidixic acid
	Benzoquinolizine	Levonadifloxacin
		Besifloxacin
	Fluoroquinolones	
		Ciprofloxacin Clinafloxacin
		Delafloxacin
		Enoxacin
		Finafloxacin
		Fleroxacin
		Gatifloxacin
		Gemifloxacin
		Grepafloxacin Levofloxacin
		Lomefloxacin
		Moxifloxacin
		Norfloxacin
		Ofloxacin
		Ozenoxacin
		Pefloxacin
		Sparfloxacin
		Trovafloxacin
		Ulifloxacin (prulifloxacin)

Glossary I

Glossary I (Part 2). (Continued)

Antimicrobial Class	Antimicrobial Subclass(es)	Agent(s) Included; Generic Name(s)
Quinolonyl oxazolidinone		Cadazolid
Spiropyrimidinetrione		Zoliflodacin
Steroid	Fusidane	Fusidic acid
Streptogramins		Quinupristin-dalfopristin
Tetracyclines		Doxycycline
		Minocycline
		Tetracycline
	Fluorocycline	Eravacycline
	Glycylcycline	Tigecycline
	Aminomethylcycline	Omadacycline
Triazaacenaphthylene		Gepotidacin

Abbreviation: FDA, US Food and Drug Administration.

NOTE: Information in boldface type is new or modified since the previous edition.

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M100, 30th ed.

Glossary II. Antimicrobial Agent Abbreviation(s), Route(s) of Administration, and Drug Class

In the late 1990s, several authorities were consulted to construct the glossary. The intention was to include all agents that appeared in M100, along with related agents available for human use. Since that time, agents have been added to the glossary as they were introduced to CLSI, and they do not need to be FDA cleared to be included. It cannot be assumed that the list is exhaustive, and it should be noted that some agents are no longer available for human use.

		Route(s) of Administration ^b				
Antimicrobial Agent		PO	IM `	IV	Topical	Drug Class or Subclass
Amikacin	AN, AK, Ak, AMI, AMK		Х	Х		Aminoglycoside
Amikacin-fosfomycin	AKF	Xc				Aminoglycoside-fosfomycin
Amoxicillin	AMX, Amx, AMOX, AC, AML	Х				Penicillin
Amoxicillin-clavulanate	AMC, Amc, A/C, AUG, Aug, XL, AML	Х				β-lactam combination agent
Ampicillin	AM, Am, AMP	Х	Х	Х		Penicillin
Ampicillin-sulbactam	SAM, A/S, AMS, AB			Х		β-lactam combination agent
Azithromycin	AZM, Azi, AZI, AZ	Х		Х		Macrolide
Azlocillin	AZL, AZ, Az		Х	Х		Penicillin
Aztreonam	ATM, AZT, Azt, AT, AZM			Х		Monobactam
Aztreonam-avibactam	AZA			Х		β-lactam combination agent
Besifloxacin	BES				Х	Fluoroquinolone
Biapenem	BPM			Х		Carbapenem
Cadazolid	CDZ	Х				Quinolonyl oxazolidinone
Carbenicillin (indanyl salt)	CB, Cb, BAR, CAR	Х				Penicillin
Carbenicillin	СВ		Х	X		
Cefaclor	CEC, CCL, Cfr, FAC, CF	Х				Cephem
Cefadroxil	CFR, FAD	Х				Cephem
Cefamandole	MA, CM, Cfm, FAM		Х	Х		Cephem
Cefazolin	CZ, CFZ, Cfz, FAZ, KZ		Х	Х		Cephem
Cefdinir	CDR, Cdn, DIN, CD, CFD	Х				Cephem
Cefditoren	CDN, DIT, FD	Х				Cephem
Cefepime	FEP, Cpe, PM, CPM		X	X		Cephem
Cefepime-enmetazobactam	FPE			Х		β-lactam combination agent
Cefepime-taniborbactam	FTB			Х		β-lactam combination agent
Cefepime-tazobactam	FPT			Х		β-lactam combination agent
Cefepime-zidebactam	FPZ			Х		β-lactam combination agent
Cefetamet	CAT, FET	Х				Cephem
Cefiderocol	FDC			Х		Siderophore β-lactam
Cefixime	CFM, FIX, Cfe, IX	Х				Cephem
Cefmetazole	CMZ, CMZS, CMT, Cmz		Х	Х		Cephem
Cefonicid	CID, Cfc, FON, CPO		Х	Х		Cephem

Glossary II. (Continued)

ordery in (continuou)	t Abbreviation(s) ^a	Route(s) of Administration ^b				
Antimicrobial Agent		РО	IM)	IV	Topical	Drug Class or Subclass
Cefoperazone	CFP, Cfp, CPZ, PER, FOP, CP		Х	Х		Cephem
Cefotaxime	CTX, TAX, Cft, FOT, CT		X	Χ		Cephem
Cefotetan	CTT, CTN, Ctn, CTE, TANS, CN		X	Χ		Cephem
Cefoxitin	FOX, CX, Cfx, FX		Х	Х		Cephem
Cefpirome	CPO, CPR, CR		X	Χ		Cephem
Cefpodoxime	CPD, Cpd, POD, PX	X				Cephem
Cefprozil	CPR, CPZ, FP	X				Cephem
Ceftaroline	CPT, Cpt			Х		Cephem
Ceftaroline-avibactam	CPA			Х		β-lactam combination agent
Ceftazidime	CAZ, Caz, TAZ, TZ		Х	Х		Cephem
Ceftazidime-avibactam	CZA			X		β-lactam combination agent
Ceftibuten	CTB, TIB, CB	Х				Cephem
Ceftizoxime	ZOX, CZX, CZ, CZ, CTZ, TIZ		Х	Х		Cephem
Ceftobiprole	BPR			X		Cephem
Ceftolozane-tazobactam	C/T CXT			Х		β-lactam combination agent
Ceftriaxone	CRO, CTR, FRX, Cax, AXO, TX		Х	Х		Cephem
Cefuroxime (oral)	CXM, CFX,	Х				Cephem
Coran commo (cran)	ROX, Crm,					G 5 p. 1 s. 1.
Cefuroxime (parenteral)	FUR, XM		X	Χ		
Cephalexin	CN, LEX, CFL, CL	Х				Cephem
Cephalothin	CF, Cf, CR, CL, CEP, CE, KF			Х		Cephem
Cephapirin	CP, HAP		Х	Х		Cephem
Cephradine	RAD, CH, CED, CE	Х				Cephem
Chloramphenicol	C, CHL, CL	Х		Х		Phenicol
Cinoxacin	CIN, Cn	Х				Quinolone
Ciprofloxacin	CIP, Cp, CI	X		Х		Fluoroguinolone
Clarithromycin	CLR, CLM, CLA, Cla, CH	X				Macrolide
Clinafloxacin	CFN, CLX, LF	Х		Х		Fluoroquinolone
Clindamycin	CC, CM, CD, Cd, CLI, DA	Х	Х	Х		Lincosamide
Colistin	CL, CS, CT, CI, CO, COL			Х		Lipopeptide
Dalbavancin	DAL			X		Lipoglycopeptide
Daptomycin	DAP, Dap, DPC			X		Lipopeptide
Delafloxacin	DLX	Х		Х		Fluoroguinolone
Dicloxacillin	DX, DIC	X				Penicillin
Dirithromycin	DTM, DT, DIR	X				Macrolide
Doripenem	DOR, Dor			Х		Carbapenem
Doxycycline	DO, DOX, DC, DOXY, D, DX, Dox	Х		X		Tetracycline
Eravacycline	ERV	X		X		Fluorocycline
Ertapenem	ETP, Etp		Х	X		Carbapenem
Erythromycin	E, ERY, EM	Х		X		Macrolide
Exebacase	EXE	† ^ ·		X		Antistaphylococcal lysin
Faropenem	FAR, FARO, FPM, Faro	Х				Penem

M100, 30th ed.

Glossary II. (Continued)

			Route(s) o				
Antimicrobial Agent	Abbreviation(s) ^a	РО	IM	IV	Topical	Drug Class or Subclass	
Fidaxomicin	FDX	Х				Macrocyclic	
Finafloxacin	FIN	Х		X	Х	Fluoroquinolone	
Fleroxacin	FLE, Fle	X		X		Fluoroquinolone	
Fosfomycin	FOS, FF, FO, FM, Fos	X				Fosfomycin	
Fusidic acid	FA, FC, FUS, FD	Х		Х	Х	Steroidal	
Garenoxacin	GRN, Grn, GA	X		X		Quinolone	
Gatifloxacin	GAT, Gat	X		X		Fluoroquinolone	
Gemifloxacin	GEM, Gem	Х				Fluoroquinolone	
Gentamicin	GM, Gm, CN, GEN		Х	Х		Aminoglycoside	
Gentamicin synergy	GM500, HLG, Gms, GHLR, GMS						
Gepotidacin	GEP, GEN, CN, GN	Х		Х		Triazaacenaphthylene	
Grepafloxacin	GRX, Grx, GRE, GP	Х				Fluoroquinolone	
Iclaprim	ICL, IP			Х		Folate pathway antagonist	
Imipenem	IPM, IMI, Imp, IP			Х		Carbapenem	
Imipenem-relebactam	IMR			Х		β-lactam combination agents	
Kanamycin	K, KAN, HLK, KM		Х	Х		Aminoglycoside	
Lefamulin	LMU, LE	Х		X		Pleuromutilin	
Levofloxacin	LVX, Lvx, LEV, LEVO, LE	Х		Х		Fluoroquinolone	
Levonadifloxacin	LND			X		Benzoquinolizine	
Linezolid	LNZ, LZ, LZD, Lzd	Х		Х		Oxazolidinone	
Lomefloxacin	LOM, Lmf	Х				Fluoroquinolone	
Loracarbef	LOR, Lor, LO	Х				Cephem	
Mecillinam	MEC, Mec	Х				Penicillin	
Meropenem	MEM, Mer, MERO, MRP, MP			Х		Carbapenem	
Meropenem-nacubactam	MNC			Х		β-lactam combination agent	
Meropenem-vaborbactam	MEV			Х		β-lactam combination agent	
Metronidazole	MET, MTZ, MZ, MRD, MTR	Х		X		Nitroimidazole	
Minocycline	MI, MIN, Min, MN, MNO, MC, MH	X		X		Tetracycline	
Moxalactam	MOX, Mox		Х	X		Cephem	
Moxifloxacin	MXF, Mxf, MX	Х		X		Fluoroquinolone	
Mupirocin	MUP, MOP, MU, Mup				Х	Pseudomonic acid	
Nafcillin	NF, NAF, Naf		Х	Х		Penicillin	
Nafithromycin	ZMK	Х				Ketolide	
Nalidixic acid	NA, NAL	X				Quinolone	
Netilmicin	NET, Nt, NC		Х	Х		Aminoglycoside	
Nitazoxanide	NIT	X				Thiazolide	
Nitrofurantoin	FM, F/M, FD, Fd, FT, NIT, NI, F	X				Nitrofuran	
Norfloxacin	NOR, Nxn, NX	X				Fluoroquinolone	
Ofloxacin	OFL, OFX, Off, OF	X	Х	Х		Fluoroquinolone	
Omadacycline	OMC, OLE, OL	X	 ^	X		Tetracycline	
Omadacyonne	OIVIO, OLL, OL	^	I .	^	1	Tottadyonite	

Glossary II. (Continued)

			Route(s) of				
Antimicrobial Agent	Abbreviation(s) ^a	PO	IM	IV	Topical	Drug Class or Subclass	
Oritavancin	ORI			X		Lipoglycopeptide	
Oxacillin	OX, Ox, OXS, OXA	X	X	Х		Penicillin	
Ozenoxacin	OZN				X	Fluoroquinolone	
Pefloxacin	PEF, PF, Pef, PE					Fluoroquinolone	
Penicillin	P, PEN, PV, PG	X	X	X		Penicillin	
Pexiganan	PEX, P/N				X	Peptide	
Piperacillin	PIP, PI, PP, Pi, PRL		X	X		Penicillin	
Piperacillin-tazobactam	TZP, PTZ, P/T, PTc			X		β-lactam combination agent	
Plazomicin	PLZ			Х		Aminoglycoside	
Polymyxin B	PB, POL, PO			Х		Lipopeptide	
Quinupristin-dalfopristin	SYN, Syn, QDA, RP			Х		Streptogramin	
Razupenem	RZ, RZM			Х		Carbapenem	
Ramoplanin	RAM	Х				Lipoglycodepsipeptide	
Rifampin	RA, RIF, Rif, RI, RD	Х		X		Ansamycin	
Rifaximin	RFP	Х				Ansamycin	
Secnidazole	SEC	X				Nitroimidazole	
Solithromycin	SOL	X		Х	X	Fluoroketolide	
Sparfloxacin	SPX, Sfx, SPA, SO	X				Fluoroquinolone	
Spectinomycin	SPT, SPE, SC, SP		Х	Х		Aminocyclitol	
Streptomycin	STS, S, STR,		X	Х		Aminoglycoside	
	StS, SM,						
Streptomycin synergy	ST2000, HLS						
Sulbactam-durlobactam	SUD					β-lactam combination agent	
Sulfonamides	SF, G, SSS, S3	X		Х		Folate pathway antagonist	
	017 0111 0	.,				(some PO only)	
Sulopenem	SLP, SULO	X		Х		Penem	
Surotomycin	SUR	X				Lipopeptide	
Tebipenem	TBP	X				Carbapenem	
Tedizolid	TZD	Х		X		Oxazolidinone	
Teicoplanin	TEC, TPN, Tei, TEI, TP, TPL		X	Х		Lipoglycopeptide	
Telavancin	TLV			Х		Lipoglycopeptide	
Telithromycin	TEL	X				Ketolide	
Tetracycline	TE, Te, TET, TC	X		X		Tetracycline	
Ticarcillin	TIC, TC, TI, Ti		X	Х		Penicillin	
Ticarcillin-clavulanate	TIM, Tim, T/C, TCC, TLc			Х		β-lactam combination agent	
Tigecycline	TGC, Tgc			Х		Glycylcycline	
Tinoxanide	TIN	X				Thiazolide	
Tinidazole	TNZ	X				Nitroimidazoles	

Glossary II

Glossary II. (Continued)

			Route(s) of			
Antimicrobial Agent	Abbreviation(s) ^a	PO	IM	IV	Topical	Drug Class or Subclass
Tobramycin	TM, NN, TO, To, TOB		X	X		Aminoglycoside
Trimethoprim	TMP, T, TR, W	X				Folate pathway antagonist
Trimethoprim-sulfamethoxazole	SXT, SxT, T/S, TS, COT	X		X		Folate pathway antagonist
Trospectomycin	TBR		X	X		Aminocyclitol
Trovafloxacin	TVA, Tva, TRV, TV, TRO	X		X		Fluoroquinolone
Ulifloxacin (prulifloxacin)	PRU	X				Fluoroquinolone
Vancomycin	VA, Va, VAN	X		X		Glycopeptide
Zoliflodacin	ZFD	Х				Spiropyriminetrione

Abbreviations: FDA, US Food and Drug Administration; PO, oral; IM, intramuscular; IV, intravenous.

Footnotes

- a. Abbreviations assigned to one or more diagnostic products in the United States. If no diagnostic product is available, abbreviation is that of the manufacturer.
- b. As available in the United States.
- c. Amikacin-fosfomycin is aerosolized and inhaled.

NOTE: Information in boldface type is new or modified since the previous edition.

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M100, 30th ed.

Glossary III. List of Identical Abbreviations Used for More Than One Antimicrobial Agent in US Diagnostic Products

In the late 1990s, several authorities were consulted to construct the glossary. The intention was to include all agents that appeared in M100, along with related agents available for human use. Since that time, agents have been added to the glossary as they were introduced to CLSI, and they do not need to be FDA cleared to be included. It cannot be assumed that the list is exhaustive, and it should be noted that some agents are no longer available for human use.

Abbreviation	Antimicrobial Agents for Which Respective Abbreviation Is Used
AZ	Azithromycin, azlocillin
AZM	Azithromycin, aztreonam
CB, Cb	Ceftibuten, carbenicillin
CD, Cd	Clindamycin, cefdinir
CF, Cf	Cefaclor, cephalothin
CFM, Cfm	Cefixime, cefamandole
CFR, Cfr	Cefaclor, cefadroxil
CFX, Cfx	Cefoxitin, cefuroxime
CH	Clarithromycin, cephradine
CL	Cephalothin, chloramphenicol
CM	Clindamycin, cefamandole
CN, Cn	Cephalexin, cefotetan, cinoxacin, gentamicin
CP, Cp	Cephapirin, cefoperazone, ciprofloxacin
CPZ	Cefprozil, cefoperazone
CZ, Cz	Ceftizoxime, cefazolin
DX	Doxycycline, dicloxacillin
FO	Fleroxacin, fosfomycin
NIT	Nitazoxanide, nitrofurantoin
TC	Tetracycline, ticarcillin

Abbreviation: FDA, US Food and Drug Administration.

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M100, 30th ed. For Use With M02 and M07

The Quality Management System Approach

Clinical and Laboratory Standards Institute (CLSI) subscribes to a quality management system (QMS) approach in the development of standards and guidelines that facilitates project management, defines a document structure using a template, and provides a process to identify needed documents. The QMS approach applies a core set of "quality system essentials" (QSEs), basic to any organization, to all operations in any health care service's path of workflow (ie, operational aspects that define how a particular product or service is provided). The QSEs provide the framework for delivery of any type of product or service, serving as a manager's guide. The QSEs are:

•	Organization and Leadership	•	Supplier and Inventory	•	Information Management
•	Customer Focus		Management	•	Nonconforming Event Management
•	Facilities and Safety	•	Equipment Management	•	Assessments
	Management	•	Process Management	•	Continual Improvement
•	Personnel Management	•	Documents and Records		-
			Management		

M100 covers the QSE indicated by an "X." For a description of the other documents listed in the grid, please refer to the Related CLSI Reference Materials section.

Organization and Leadership	Customer Focus	Facilities and Safety Management	Personnel Management	Supplier and Inventory Management	Equipment Management	Process Management	Documents and Records Management	Information Management	Nonconforming Event Management	Assessments	Continual Improvement
						X EP23					
						M02					
						M07					
						M11					
						M23					
						M39					
						M45					
						M52					

Path of Workflow

A path of workflow is the description of the necessary processes to deliver the particular product or service that the organization or entity provides. A laboratory path of workflow consists of the sequential processes: preexamination, examination, and postexamination and their respective sequential subprocesses. All laboratories follow these processes to deliver their services, namely quality laboratory information.

M100 covers the medical laboratory path of workflow processes indicated by an "X." For a description of the other documents listed in the grid, please refer to the Related CLSI Reference Materials section.

Preexamination					Exami	Postexamination				
Examination ordering	Specimen collection	Specimen transport	Specimen receipt, accessioning, and processing	Examination method selection	Examination performance	Results review and follow-up	Laboratory results interpretation	Communication of alert values and issuance of preliminary reports	Release of final reports	Specimen management
				EP23 M07 M11	EP23 M02 M07 M11	X EP23 M02 M07 M11	M02 M07 M11 M39 M45		X	

For Use With M02 and M07 M100, 30th ed.

Related CLSI Reference Materials*

EP23TM Laboratory Quality Control Based on Risk Management. 1st ed., 2011. This document provides guidance based on risk management for laboratories to develop quality control plans tailored to the particular combination of measuring system, laboratory setting, and clinical application of the test.

M02 Performance Standards for Antimicrobial Disk Susceptibility Tests. 13th ed., 2018. This standard covers the current recommended methods for disk susceptibility testing and criteria for quality control testing.

M02QG M02 Disk Diffusion Reading Guide. 1st ed., 2018. The Disk Diffusion Reading Guide provides photographic examples of the proper method for reading disk diffusion susceptibility testing results.

M07 Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically. 11th ed., 2018. This standard covers reference methods for determining minimal inhibitory concentrations of aerobic bacteria by broth macrodilution, broth microdilution, and agar dilution.

M11 Methods for Antimicrobial Susceptibility Testing of Anaerobic Bacteria. 9th ed., 2018. This standard provides reference methods for determining minimal inhibitory concentrations of anaerobic bacteria by agar dilution and broth microdilution.

M23 Development of *In Vitro* Susceptibility Testing Criteria and Quality Control Parameters. 5th ed., 2018. This guideline discusses the necessary and recommended data for selecting appropriate breakpoints and quality control ranges for antimicrobial agents.

M39 Analysis and Presentation of Cumulative Antimicrobial Susceptibility Test Data. 4th ed., 2014. This document describes methods for recording and analysis of antimicrobial susceptibility test data, consisting of cumulative and ongoing summaries of susceptibility patterns of clinically significant microorganisms.

M45 Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently Isolated or Fastidious Bacteria. 3rd ed., 2016. This guideline informs clinical, public health, and research laboratories on susceptibility testing of infrequently isolated or fastidious bacteria that are not included in CLSI documents M02, M07, or M100. Antimicrobial agent selection, test interpretation, and quality control are addressed.

Werification of Commercial Microbial Identification and Antimicrobial Susceptibility Testing Systems.

1st ed., 2015. This guideline includes recommendations for verification of commercial US Food and Drug Administration—cleared microbial identification and antimicrobial susceptibility testing systems by clinical laboratory professionals to fulfill regulatory or quality assurance requirements for the use of these systems for diagnostic testing.

^{*} CLSI documents are continually reviewed and revised through the CLSI consensus process; therefore, readers should refer to the most current editions.

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NOTES

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PRINT ISBN 978-1-68440-066-9

ELECTRONIC ISBN 978-1-68440-067-6