

# M07

## Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically

This standard covers reference methods for determining minimal inhibitory concentrations of aerobic bacteria by broth macrodilution, broth microdilution, and agar dilution.

A standard for global application developed through the Clinical and Laboratory Standards Institute consensus process.

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Clinical and Laboratory Standards Institute  
950 West Valley Road, Suite 2500  
Wayne, PA 19087 USA  
P: +1.610.688.0100  
F: +1.610.688.0700  
[www.clsi.org](http://www.clsi.org)  
[standard@cls.org](mailto:standard@cls.org)

## Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically

Melvin P. Weinstein, MD  
Jean B. Patel, PhD, D(ABMM)  
Carey-Ann Burnham, PhD, D(ABMM)  
Shelley Campeau, PhD, D(ABMM)  
Patricia S. Conville, MS, MT(ASCP)  
Christopher Doern, PhD, D(ABMM)  
George M. Eliopoulos, MD  
Marcelo F. Galas  
Romney M. Humphries, PhD, D(ABMM)  
Stephen G. Jenkins, PhD, D(ABMM), F(AAM)  
Susan M. Kircher, MS, MT(ASCP)  
James S. Lewis II, PharmD, FIDSA

Brandi Limbago, PhD  
Amy J. Mathers, MD, D(ABMM)  
Tony Mazzulli, MD, FACP, FRCP(C)  
Susan D. Munro, CLS, MT(ASCP)  
Margaret Ordoñez Smith de Danies, PhD  
Robin Patel, MD  
Sandra S. Richter, MD, D(ABMM), FCAP, FIDSA  
Michael Satlin, MD, MS  
Jana M. Swenson, MMSc  
Alexandra Wong, BS, MT(ASCP), SM  
Wayne F. Wang, MD, PhD  
Barbara L. Zimmer, PhD

### Abstract

Antimicrobial susceptibility testing is indicated for any organism that contributes to an infectious process warranting antimicrobial chemotherapy, if its susceptibility cannot be reliably predicted from knowledge of the organism's identity. Susceptibility tests are most often indicated when the causative organism is thought to belong to a species capable of exhibiting resistance to commonly used antimicrobial agents.

Various laboratory methods can be used to measure the *in vitro* susceptibility of bacteria to antimicrobial agents. Clinical and Laboratory Standards Institute standard M07—*Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically* describes standard broth dilution (macrodilution and microdilution [the microdilution method described in M07 is the same methodology outlined in ISO 20776-1<sup>1</sup>]) and agar dilution techniques, and it includes a series of procedures to standardize the way the tests are performed. The performance, applications, and limitations of the current CLSI-recommended methods are also described.

The supplemental information (M100<sup>2</sup> tables) used with this standard represents the most current information for drug selection, interpretation, and quality control using the procedures standardized in M07.

Clinical and Laboratory Standards Institute (CLSI). *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically*. 11th ed. CLSI standard M07 (ISBN 1-56238-836-3 [Print]; ISBN 1-56238-837-1 [Electronic]). Clinical and Laboratory Standards Institute, 950 West Valley Road, Suite 2500, Wayne, Pennsylvania 19087 USA, 2018.

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### **Suggested Citation**

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.

### **Previous Editions:**

June 1980, December 1982, June 1986, November 1988, April 1990, December 1993, January 1997, January 2000, January 2003, January 2006, January 2009, January 2012, January 2015

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## Foreword

The most current edition of CLSI document M100,<sup>2</sup> an annually published volume of tables, is made available with this standard to ensure users are aware of the latest recommendations related to the methods described in M07 and CLSI document M02.<sup>3</sup>

Many other editorial and procedural changes in this edition of M07 resulted from Subcommittee on Antimicrobial Susceptibility Testing meetings held since 2015. Specific changes to the tables are summarized at the beginning of M100.<sup>2</sup> The most important changes in M07 are summarized below.

## Overview of Changes

This standard replaces the previous edition of the approved standard, M07-A10, published in 2015. Several changes were made in this edition, including:

- **General:**
  - Harmonized language and information on drug selection and QC with CLSI document M02<sup>3</sup>
  - To harmonize with the International Organization for Standardization, the terms for the methods for inoculum preparation have been changed. “Growth method” has been changed to “broth culture method,” and “direct colony suspension method” has been changed to “colony suspension method” throughout the document
- **Subchapter 1.4.1, Definitions:**
  - Clarified definitions for breakpoint, interpretive category, susceptible, susceptible-dose dependent, intermediate, resistant, nonsusceptible, and quality control
  - Added definitions for minimal inhibitory concentration, routine test, supplemental test, surrogate agent test, CarbaNP test, and modified carbapenem inactivation method
- **Subchapter 1.4.2, Abbreviations and Acronyms:**
  - Deleted abbreviations for  $\beta$ -lactamase types
- **Subchapter 2.3, Antimicrobial Agent Classes:**
  - Clarified and updated antimicrobial agent classes
- **Subchapter 2.3.2.2, Folate Pathway Antagonists:**
  - Revised nomenclature from “folate pathway inhibitor” to “folate pathway antagonist”
- **Subchapter 3.9, Determining Broth Macro- or Microdilution End Points:**
  - Added photographs of growth control examples and for interpreting skipped wells
- **Subchapter 3.11, Table 1. Testing Considerations for Fastidious Organisms:**
  - Clarified source plate incubation times and inoculum broth for some fastidious organisms
- **Subchapter 3.12, Special Considerations for Detecting Resistance:**
  - Reorganized and streamlined
  - Moved Subchapters 3.12.4 (Inducible Clindamycin Resistance) and 3.12.6 ( $\beta$ -Lactamase Tests) to create a new subchapter, 3.13 (Supplemental [Not Routine] Tests)

- **Subchapter 3.12.1, Staphylococci:**
  - Added information for *Staphylococcus pseudintermedius* and *Staphylococcus schleiferi*
  - Reorganized and clarified information for staphylococci
  
- **Subchapter 3.12.4, Gram-Negative Bacilli:**
  - Expanded and clarified information on  $\beta$ -lactamases
  
  - Added footnote to Table 4, Enzyme Classifications for  $\beta$ -Lactamases, to clarify the difference between cephalosporin subclasses and generations
  
  - Updated nomenclature for *Enterobacter aerogenes* to *Klebsiella* (formerly *Enterobacter aerogenes*<sup>4</sup>)
  
- **Subchapter 3.13.1, Inducible Clindamycin Resistance:**
  - Consolidated information from former Subchapter 3.13.1.8
  
- **Subchapter 4.3, Selecting Strains for Quality Control:**
  - Clarified the example in the third paragraph
  
- **Appendixes:**
  - Reorganized to reflect the order in which they are referenced in the main text, as follows:
    - **Appendix A. Preparation of Supplements, Media, and Reagents** (formerly Appendix B)
  
    - **Appendix B. Conditions for Dilution Antimicrobial Susceptibility Tests** (formerly Appendix C)
  
    - **Appendix C. Quality Control Strain Maintenance** (formerly Appendix E)
  
    - **Appendix D. Quality Control Protocol Flow Charts** (formerly Appendix A)
  
  - Deleted **Quality Control Strains for Antimicrobial Susceptibility Tests** (formerly Appendix D) (see M100<sup>2</sup> Appendix C)
  
- **Appendix A. Preparation of Supplements, Media, and Reagents:**
  - Reorganized procedures into step-action tables
  
- **Appendix C. Quality Control Strain Maintenance:**
  - Clarified maintenance and subculture of QC strains
  
- **Appendix D. Quality Control Protocol Flow Charts:**
  - Recreated QC flow charts in black-and-white format for easier viewing
  - Revised Appendixes D1 and D2 flow charts

## 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 is found at [www.clsi.org](http://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.<sup>5</sup>

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 standard are found in the meeting summary minutes of the Subcommittee on Antimicrobial Susceptibility Testing at [www.clsi.org](http://www.clsi.org).

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

It is important for users of M02,<sup>3</sup> M07, and the M100<sup>2</sup> supplement 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 evaluation of commercial devices that will be used in medical laboratories, or by drug or device manufacturers for testing of new agents or systems. Results generated by reference methods, such as those contained 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.<sup>5</sup>

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 control committees of the medical staff, and antimicrobial stewardship teams), newly approved or revised breakpoints may be implemented by laboratories. Following verification, CLSI broth dilution and agar dilution test breakpoints may be implemented as soon as they are published in M100.<sup>2</sup> 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.

## 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 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.

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

### Key Words

Agar dilution, antimicrobial susceptibility, broth dilution, broth macrodilution, broth microdilution, minimal inhibitory concentration

# Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically

## Chapter 1: Introduction

This chapter includes:

- Standard's scope and applicable exclusions
- Background information pertinent to the standard's content
- Standard precautions information
- Terms and definitions used in the standard
- Abbreviations and acronyms used in the standard

### 1.1 Scope

This standard describes standard broth (macrodilution and microdilution) and agar dilution methods for determining *in vitro* susceptibility to antimicrobial agents for bacteria that grow aerobically and includes:

- Broth and agar dilution test preparation
- Testing conditions, including inoculum preparation and standardization, incubation time, and incubation temperature
- Reporting minimal inhibitory concentration (MIC) results
- QC procedures
- Dilution test method limitations

To assist the medical laboratory, suggestions are provided for selecting antimicrobial agents for routine testing and reporting.

Standards for testing the *in vitro* antimicrobial susceptibility of bacteria that grow aerobically using the antimicrobial disk testing method are found in CLSI document M02.<sup>3</sup> Standards for testing the *in vitro* antimicrobial susceptibility of bacteria that grow anaerobically are found in CLSI document M11.<sup>6</sup> Guidelines for standardized antimicrobial susceptibility testing (AST) of infrequently isolated or fastidious bacteria that are not included in CLSI documents M02,<sup>3</sup> M07, or M11<sup>6</sup> are available in CLSI document M45.<sup>7</sup> The AST methods provided in this standard can be used in laboratories around the world including but not limited to:

- Medical laboratories
- Public health laboratories
- Research laboratories
- Food laboratories
- Environmental laboratories

# 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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Clinical and Laboratory Standards Institute  
950 West Valley Road, Suite 2500  
Wayne, PA 19087 USA  
P: +1.610.688.0100  
F: +1.610.688.0700  
[www.clsi.org](http://www.clsi.org)  
[standard@clsi.org](mailto:standard@clsi.org)

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## Performance Standards for Antimicrobial Susceptibility Testing

Melvin P. Weinstein, MD  
Jean B. Patel, PhD, D(ABMM)  
April M. Bobenchik, PhD, D(ABMM)  
Shelley Campeau, PhD, D(ABMM)  
Sharon K. Cullen, BS, RAC  
Marcelo F. Galas  
Howard Gold, MD, FIDSA  
Romney M. Humphries, PhD, D(ABMM)  
Thomas J. Kirn, Jr., MD, PhD

James S. Lewis II, PharmD, FIDSA  
Brandi Limbago, PhD  
Amy J. Mathers, MD, D(ABMM)  
Tony Mazzulli, MD, FACP, FRCP(C)  
Sandra S. Richter, MD, D(ABMM), FCAP, FIDSA  
Michael Satlin, MD, MS  
Audrey N. Schuetz, MD, MPH, D(ABMM)  
Jana M. Swenson, MMSc  
Pranita D. Tamma, MD, MHS

### Abstract

The data in the tables are valid only if the methodologies in CLSI documents M02,<sup>1</sup> M07,<sup>2</sup> and M11<sup>3</sup> are followed. These standards contain information about disk diffusion (M02<sup>1</sup>) and dilution (M07<sup>2</sup> and M11<sup>3</sup>) 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,<sup>1</sup> M07,<sup>2</sup> and M11.<sup>3</sup> 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*. 29th ed. CLSI supplement M100 (ISBN 978-1-68440-032-4 [Print]; ISBN 978-1-68440-033-1 [Electronic]). Clinical and Laboratory Standards Institute, 950 West Valley Road, Suite 2500, Wayne, Pennsylvania 19087 USA, 2019.

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### **Suggested Citation**

CLSI. *Performance Standards for Antimicrobial Susceptibility Testing*. 29th ed. CLSI supplement M100. Wayne, PA: Clinical and Laboratory Standards Institute; 2019.

### **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 2009, January 2010, June 2010, January 2011, January 2012, January 2013, January 2014, January 2015, January 2016, January 2017, January 2018

ISBN 978-1-68440-032-4 (Print)  
ISBN 978-1-68440-033-1 (Electronic)  
ISSN 1558-6502 (Print)  
ISSN 2162-2914 (Electronic)

Volume 39, Number 1

## Committee Membership

### Subcommittee on Antimicrobial Susceptibility Testing

<b>Melvin P. Weinstein, MD</b> <b>Chairholder</b> <b>Rutgers Robert Wood Johnson Medical School</b> USA	Romney M. Humphries, PhD, D(ABMM) Accelerate Diagnostics, Inc. USA	Tony Mazzulli, MD, FACP, FRCP(C) Mount Sinai Hospital Canada
<b>Jean B. Patel, PhD, D(ABMM)</b> <b>Vice-Chairholder</b> <b>Centers for Disease Control and Prevention</b> USA	Thomas J. Kirm, Jr., MD, PhD Rutgers Robert Wood Johnson Medical School USA	Sandra S. Richter, MD, D(ABMM), FCAP, FIDSA Cleveland Clinic USA
Sharon K. Cullen, BS, RAC Beckman Coulter, Inc. Microbiology Business USA	James S. Lewis II, PharmD, FIDSA Oregon Health and Science University USA	Michael Satlin, MD, MS New York Presbyterian Hospital USA
Marcelo F. Galas Pan American Health Organization USA	Brandi Limbago, PhD Centers for Disease Control and Prevention USA	Audrey N. Schuetz, MD, MPH, D(ABMM) Mayo Clinic USA
Howard Gold, MD, FIDSA Beth Israel Deaconess Medical Center USA	Amy J. Mathers, MD, D(ABMM) University of Virginia Medical Center USA	Pranita D. Tamma, MD, MHS Johns Hopkins University School of Medicine USA

### Acknowledgment

CLSI and the Subcommittee on Antimicrobial Susceptibility Testing gratefully acknowledge the following volunteers for their important contributions to the development of this document:

April M. Bobenchik, PhD, D(ABMM) Lifespan Academic Medical Center USA	Shelley Campeau, PhD, D(ABMM) Accelerate Diagnostics, Inc. USA	Jana M. Swenson, MMSc USA
---	--	------------------------------

### Working Group on AST Breakpoints

<b>George M. Eliopoulos, MD</b> <b>Co-Chairholder</b> <b>Beth Israel Deaconess Medical Center</b> USA	Amy J. Mathers, MD, D(ABMM) University of Virginia Medical Center USA	Simone M. Shurland FDA Center for Devices and Radiological Health USA
<b>James S. Lewis II, PharmD, FIDSA</b> <b>Co-Chairholder</b> <b>Oregon Health and Science University</b> USA	David P. Nicolau, PharmD, FCCP, FIDSA Hartford Hospital USA	Lauri D. Thrupp, MD University of California Irvine Medical Center USA
<b>Karen Bush, PhD</b> <b>Committee Secretary</b> <b>Indiana University</b> USA	Robin Patel, MD Mayo Clinic USA	Hui Wang, MD Peking University People's Hospital China
Marcelo F. Galas Pan American Health Organization USA	Michael Satlin, MD, MS New York Presbyterian Hospital USA	Barbara L. Zimmer, PhD Beckman Coulter, Inc. USA

### **Working Group on Methods Application and Interpretation**

**Thomas J. Kirn, Jr., MD, PhD**  
**Co-Chairholder**  
**Rutgers Robert Wood Johnson Medical School**  
**USA**

**Brandi Limbago, PhD**  
**Co-Chairholder**  
**Centers for Disease Control and Prevention**  
**USA**

**Patricia J. Simner, PhD, D(ABMM)**  
**Committee Secretary**  
**Johns Hopkins Hospital - Pathology**  
**USA**

Darcie E. Carpenter, PhD, CIC  
IHMAUSA  
  
Stephen G. Jenkins, PhD, D(ABMM),  
F(AAM)  
Weill Cornell Medicine  
USA

Kristie Johnson, PhD, D(ABMM)  
University of Maryland, Baltimore  
USA

Joseph Kuti, PharmD  
Hartford Hospital  
USA

Samir Patel, PhD, FCCM, D(ABMM)  
Public Health Ontario  
Canada

Virginia M. Pierce, MD  
Massachusetts General Hospital  
USA

Sandra S. Richter, MD, D(ABMM),  
FCAP, FIDSA  
Cleveland Clinic  
USA

Susan Sharp, PhD, D(ABMM),  
F(AAM)  
Copan Diagnostics, Inc.  
USA

### **Working Group on Methods Development and Standardization**

**Dwight J. Hardy, PhD**  
**Co-Chairholder**  
**University of Rochester Medical Center**  
**USA**

**Barbara L. Zimmer, PhD**  
**Co-Chairholder**  
**Beckman Coulter, Inc.**  
**USA**

**Katherine Sei, BS**  
**Committee Secretary**  
**Beckman Coulter, Inc.**  
**USA**

William B. Brasso, BS  
BD Diagnostic Systems  
USA

Susan Butler-Wu, PhD, D(ABMM),  
SM(ASCP)  
LACUSC Medical Center  
USA

Jennifer Dien Bard, PhD, D(ABMM),  
F(CCM)  
Children's Hospital Los Angeles;  
University of Southern California  
USA

Tanis Dingle, PhD, D(ABMM), FCCM  
Provincial Laboratory for Public Health  
Canada

Romney M. Humphries, PhD,  
D(ABMM)  
Accelerate Diagnostics, Inc.  
USA

Laura M. Koeth, MT(ASCP)  
Laboratory Specialists, Inc.  
USA

Ribhi M. Shawar, PhD, D(ABMM)  
FDA Center for Devices and  
Radiological Health  
USA

**Working Group on Outreach**

**Janet A. Hindler, MCLS, MT(ASCP)**  
**Co-Chairholder**  
 USA

April M. Bobenchik, PhD, D(ABMM)  
 Lifespan Academic Medical Center  
 USA

Nicole Scangarella-Oman, MS, BS,  
 GlaxoSmithKline  
 USA

**Audrey N. Schuetz, MD, MPH,**  
**D(ABMM)**  
**Co-Chairholder**  
 Mayo Clinic  
 USA

Angella Charnot-Katsikas, MD  
 The University of Chicago  
 USA

Paula M. Snippes Vagnone,  
 MT(ASCP)  
 Minnesota Department of Health  
 USA

**Stella Antonara, PhD, D(ABMM)**  
**Committee Secretary**  
 OhioHealth  
 USA

Graeme Forrest, MBBS  
 Oregon Health Sciences University  
 USA

Lars F. Westblade, PhD, D(ABMM)  
 New York Presbyterian Hospital -  
 Weill Cornell Campus  
 USA

April Abbott, PhD  
 Deaconess Hospital Laboratory  
 USA

Romney M. Humphries, PhD,  
 D(ABMM)  
 Accelerate Diagnostics, Inc.  
 USA

**Working Group on Quality Control**

**Sharon K. Cullen, BS, RAC**  
**Co-Chairholder**  
 Beckman Coulter, Inc. Microbiology  
 Business  
 USA

Dana C. Dressel, MT(ASCP)  
 International Health Management  
 Associates, Inc.  
 USA

Susan D. Munro, MT(ASCP), CLS  
 USA

**Maria M. Traczewski, BS, MT(ASCP)**  
**Co-Chairholder**  
 The Clinical Microbiology Institute  
 USA

Janet A. Hindler, MCLS, MT(ASCP)  
 USA

David Paisey, BSc  
 Thermo Fisher Scientific  
 United Kingdom

**Michael D. Huband, BS**  
**Committee Secretary**  
 JMI Laboratories  
 USA

Denise Holliday, MT(ASCP)  
 BD Diagnostic Systems  
 USA

Elizabeth Palavecino, MD  
 Wake Forest Baptist Medical Center  
 USA

Erika Matuschek, PhD  
 ESCMID  
 Sweden

Chris Pillar, PhD  
 Micromyx, LLC  
 USA

Patricia S. Conville, MS, MT(ASCP)  
 FDA Center for Devices and  
 Radiological Health  
 USA

Mary K. York, PhD, D(ABMM)  
 MKY Microbiology Consulting  
 USA

## Working Group on Text and Tables

**April M. Bobenchik, PhD, D(ABMM)**  
**Co-Chairholder**  
**Lifespan Academic Medical Center**  
**USA**

Janet A. Hindler, MCLS, MT(ASCP)  
 USA

Dale A. Schwab, PhD,  
 D(ABMM)CM  
 Quest Diagnostics Infectious Disease  
 USA

**Shelley Campeau, PhD, D(ABMM)**  
**Co-Chairholder**  
**Accelerate Diagnostics, Inc.**  
**USA**

Melissa Jones, MT(ASCP), CLS  
 UNC Healthcare  
 USA

Jana M. Swenson, MMSc  
 USA

**Carey-Ann Burnham, PhD, D(ABMM)**  
**Committee Secretary**  
**Washington University School of**  
**Medicine**  
**USA**

Peggy Kohner, BS, MT(ASCP)  
 Mayo Clinic  
 USA

Richard B. Thomson, Jr., PhD,  
 D(ABMM), FAAM  
 Evanston Hospital, NorthShore  
 University HealthSystem  
 USA

Victoria Emma Anikst, BA, CMS,  
 M(ASCP)<sup>CM</sup>  
 UCLA Health  
 USA

Dyan Luper, BS, MT(ASCP)SM, MB  
 BD Diagnostic Systems  
 USA

Maria M. Traczewski, BS,  
 MT(ASCP)  
 The Clinical Microbiology Institute  
 USA

Andrea L. Ferrell, MLS<sup>CM</sup>(ASCP)  
 Becton Dickinson  
 USA

Susan D. Munro, MT(ASCP), CLS  
 USA

Nancy E. Watz, MS, MT(ASCP),  
 CLS  
 Stanford Health Care  
 USA

L. Barth Reller, MD  
 Duke University School of Medicine  
 USA

Flavia Rossi, MD, PhD  
 University of São Paulo  
 Brazil

Mary K. York, PhD, D(ABMM)  
 MKY Microbiology Consulting  
 USA

## Acknowledgment

CLSI and the Subcommittee on Antimicrobial Susceptibility Testing gratefully acknowledge the following volunteers for their important contributions to their respective working groups and the development of this document:

Darcie E. Carpenter, PhD, CIC  
 IHMA  
 USA

Matthew A. Wikler, MD, FIDSA, MBA  
 IDTD Consulting  
 USA

Mariana Castanheira, PhD  
 JMI Laboratories  
 USA

Barbara L. Zimmer, PhD  
 Beckman Coulter, Inc.  
 USA

## Staff

Clinical and Laboratory Standards Institute  
 USA

Megan L. Tertel, MA, ELS  
*Editorial Manager*

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