What's New in the 2006 Standards for Antimicrobial Susceptibility Testing (AST)?

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At the conclusion of this talk, you will be able to.....

- Outline the major changes found in the new CLSI tables (M100-S16) and standards for disk diffusion (M2-A9) and MIC testing (M7-A7).
- Discuss how to optimally use the new Disk Diffusion and MIC QC Troubleshooting Guides.
- Describe a strategy for implementing the new practice guidelines in your laboratory, as appropriate.

CLSI Standards - 2006

◆ M100-S16 Tables (2006)*



.....to be used with text documents explaining how to perform the tests....

M2-A9 Disk Diffusion (2006)**

M7-A7 MIC (2006)**

* M100 updated yearly **M2, M7 updated every 3 years

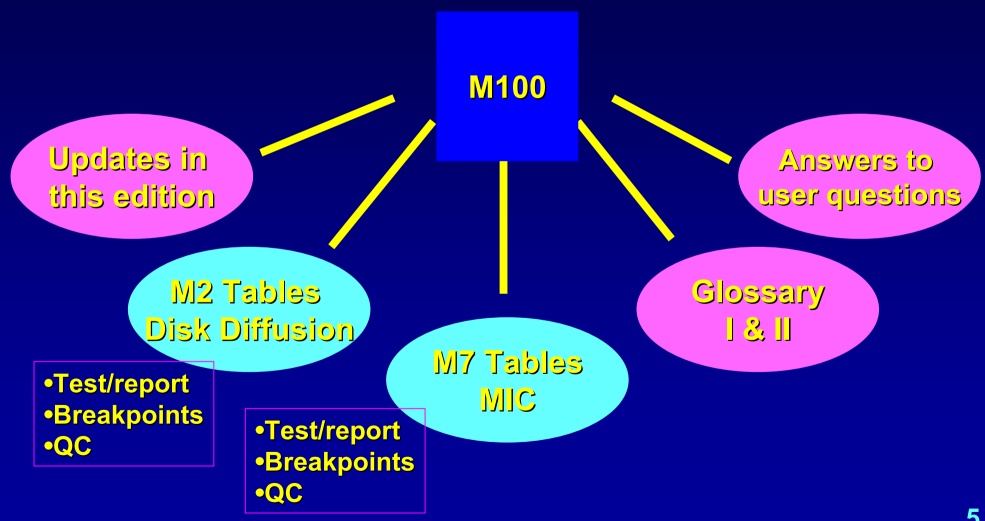


Reference Terminology

....when I refer to....

- ◆ M100 -- this means the new tables M100-S16
- M2 -- this means the new disk diffusion document M2-A9
- ◆ M7 -- this means the new MIC document M7-A7
- M45 -- this means the new fastidious organism testing document M45-P
- CLSI = formerly NCCLS

CLSI M100 contains.....



Updated information in M100-S16

Vol. 26 No. 3 M100-S16

Updated Information in This Edition

This document includes all of the tables from the Clinical and Laboratory Standards Institute Disk Diffusion (M2) susceptibility testing and Aerobic Dilution (M7) susceptibility testing documents. There are several important changes to the tables that have resulted from meetings of the Subcommittee on Antimicrobial Susceptibility Testing during 2005. Included below is a summary of the changes in this document, which supersede the tables published in 2005 and in earlier years.

Summary of Major Changes in This Document

The list includes the "major" changes in this document. Other minor or editorial changes have been made to the general formatting and to some of the table footnotes. Boldface type is used to highlight the changes in each table.

Additions/Changes/Deletions

The following are additions or changes unless otherwise noted as a "deletion."

All Tables Throughout (M2, Tables 2A-2J; M7, Tables 2A-2L)

Clarification of incubation temperature (M2 and M7)

Introduction to Tables:

Expanded rationale for listing drugs in Tables 1 and 1A (M2 n. 17 and M7 n. 91).

Changes 2006 CLSI M100-S16

7

CLSI Standard Reference Method and Breakpoints (M100-S16, page 14)

January 2006 M100-S16

It is important for users of M2-A9, M7-A7, and the M100 Informational 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 clinical isolates, for evaluation of commercial devices that will be used in clinical 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 that 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 the following: different data bases, differences in interpretation of data, differences in doses utilized in different parts of the world and public health policies. Differences also exist because the CLSI proactively evaluates the need for changing breakpoints. The reasons why breakpoints may change and the manner in which the CLSI evaluates data and determines breakpoints are outlined in CLSI document M23—Development of In Vitro Susceptibility Testing Criteria and Quality Control Parameters.

Following a decision by CLSI to shange an existing breakpoint, monletony authorities may also

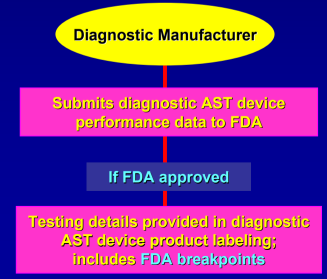
CLSI M2, M7, M100

- Describe standard consensus "reference methods"
- U.S. clinical labs can use:
 - -CLSI test method as written
 - Method that performs comparably to CLSI "reference method" (e.g. FDA-cleared diagnostic AST device)

Diagnostic AST device = commercial instrument or test used to determine antimicrobial susceptibility in vitro

New Antimicrobial Agent Pathway Pharmaceutical Company Submits "New Drug Approval" (NDA) Submits condensed version of NDA request packet to FDA request packet to CLSI -clinical outcome data -clinical outcome data -microbiological data -microbiological data -breakpoints -breakpoints -QC ranges -QC ranges -other -other If FDA approved If CLSI approved Therapeutic details provided in **Testing details and CLSI breakpoints** pharmaceutical product labeling; provided in laboratory testing "reference standards"....CLSI M2, M7 and M100 includes FDA breakpoints

Testing to establish FDA breakpoints done using CLSI "standard reference methods"



Diagnostic AST device performance data is based on manufacturer demonstrating that their device produces results comparable to results produced with CLSI "standard reference methods"

FDA vs. CLSI Breakpoints

- Nearly always agree!
- Sometimes disagree
- Sometimes only FDA breakpoints (e.g. tigecycline)
- Sometimes only CLSI breakpoints (before drug is FDA cleared or if drug used in other countries)
- Sometimes modified by CLSI (e.g., 2006, vancomycin – S. aureus)

2006 Vancomycin MIC (µg/ml) Breakpoints – *S. aureus*

CLSI ≤2 4-8 ≥16 FDA ≤4 8-16 ≥32

Presently, diagnostic manufacturers must use FDA breakpoints

Clinical laboratories can use CLSI or FDA breakpoints
Caveat: if using commercial AST instrument, system uses
FDA breakpoints

CLSI Introduction to Tables 1-1B; 2A-2L

For Use With M7-A7-MIC Testing

M100-S16

Introduction to Tables 1 Through 1B and 2A Through 2L for Use With M7-A7—MIC Testing

On the following pages, you will find:

- 1. Tables 1 and 1A—Suggested groupings of antimicrobial agents that should be considered for routine testing and reporting by clinical microbiology laboratories. These guidelines are based on drugs with clinical indications approved by the Food and Drug Administration (FDA) in the United States. In other countries, placement in Table 1 or 1A of antimicrobial agents should be based on available drugs approved for clinical use by relevant regulatory agencies.
- 2. For each organism group, an additional table (Tables 2A through 2L) that contains:
 - Recommended testing conditions.
 - b. Minimal QC recommendations. (See also the M7-A7 text document, Section 16.)
 - c. General comments for testing the organism group and specific comments for testing particular drug/organism combinations.
 - d. Suggested agents that should be considered for routine testing and reporting by clinical microbiology laboratories as specified in Tables 1 and 1A (test/report groups A, B, C, U; the latter for "urine").
 - e. Additional drugs that have an approved indication for the respective organism group, but would generally not warrant routine testing by a clinical microbiology laboratory in the United States (test/report group O for "other"; test/report group Inv. for "investigational" [not yet FDA approved]).
 - Minimal inhibitory concentration (MIC) interpretive standards.

CLSI Introduction to Tables "Clinical Indications"

"These guidelines are based on drugs with clinical indications approved by the Food & Drug Administration (FDA) in the United States. In other countries, placement in Tables 1 and 1A of antimicrobial agents should be based on available drugs approved for clinical use by relevant regulatory agencies."



Rx only

To reduce the development of drug-resistant bacteria and maintain the effectiveness of Cubicin and other antibacterial drugs,

Example of Drug Labeling

le antibacterial agent derived from the fermentation of *Streptomyces* -L-tryptophyl-L-asparaginyl-L-aspartyl-L-threonylglycyl-L-ornithyl-L-aspartyl-I-L-glutamyl-3-anthraniloyl-L-alanine ε₁-lactone. The chemical *s*tructure is:

HO
$$_{2}$$
C $_{NH}$ $_{0}$ $_{0}$ $_{NH}$ $_{0}$ $_$

removed by 4 hours of hemodialysis and 48 hours of CAP every 24 hours for patients with $CL_{CR} \ge 30$ mL/min and 4 those on hemodialysis and CAPD. Daptomycin should b hemodialysis days (see **DOSAGE AND ADMINISTRATION**

Table 2. Mean (SD) Daptomycin Population Pharmacokine Intravenous Infusion of 4 mg/kg to Infected Patients and N Renal Function

| Renal Function | AUC₀⊷ |
|---|----------|
| | (μg*h/mL |
| Normal | 417 (158 |
| (CL _{CR} >80mL/min) (N=165) | |
| Mild Renal Impairment (CL _{CR} 50-80 mL/min) (N=64) | 466 (17) |
| Moderate Renal Impairment | 560 (258 |
| (CL _{CR} 30-<50 mL/min) (N=24) | |
| Severe Renal Impairment | 925 (467 |
| (CL _{CR} <30 mL/min) (N=8) | |
| Hemodialysis and CAPD (N=21) | 1244 (37 |

Note: CL_{CR}= Creatinine clearance estimated using the

Hepatic Insufficiency

The pharmacokinetics of daptomycin were evaluated in 10 and compared with healthy volunteers (n=9) matched for not altered in subjects with moderate hepatic impairment. mycin to patients with mild to moderate hepatic impairment hepatic insufficiency have not been evaluated.

Gender

No clinically significant gender-related differences in dapte

Daptomycin product labeling...
http://www.cubicin.com/PDF/1004-2 MARKETING gt.pdf.

Drug product labeling....

- Has many sections, including...
 - (Clinical) indications and usage
 - Based on demonstrated clinical efficacy
 - Examines in vitro susceptibility test and clinical outcome data
 - Microbiological activity (in vitro)
 - Does not mean clinical efficacy

Provides information for

- Clinicians
- Pharmacists
- Diagnostic AST manufacturers
- Patients
- Others

"Clinical Indication and Usage"

 Clinical indication = a disease entity that has a specific set of signs, symptoms and laboratory findings that can be described to clinicians in product labeling.

Example:

Daptomycin has clinical indication for complicated skin and skin structure infections This means...

- Daptomycin can be used to treat S. aureus (including MRSA) wound infections as noted in package insert
- Clinical indication is linked to specific pathogens

..........

INDICATIONS AND USAGE

Subjection (daptomycin for injection) is indicated for the treatment of complicated skin and skin structure infections caused by susceptible strains of the following Gram-positive microorganisms (see also DOSAGE AND ADMINISTRATION): Staphylococcus aureus (including methicillin-resistant strains), Streptococcus pyogenes, Streptococcus agalactiae, Streptococcus dysgalactiae subsp. equisimilis and Enterococcus faecalis (vancomycin-susceptible strains only). Combination therapy may be clinically indicated if the documented or presumed pathogens include Gram-negative or anaerobic organisms. (see CLINICAL STUDIES).

Daptomycin is not indicated for the treatment of pneumonia.

Appropriate specimens for microbiological examination should be obtained in order to isolate and identify the causative pathogens and to determine their susceptibility to daptomycin. Empiric therapy may be initiated while awaiting test results. Antimicrobial therapy should be adjusted as needed based upon test results.

To reduce the development of drug-resistant bacteria and maintain the effectiveness of Cubicin and other antibacterial drugs, Cubicin should be used only to treat or prevent infections that are proven or strongly suspected to be caused by susceptible bacteria. When culture and susceptibility information are available, they should be considered in selecting or modifying antibacterial therapy. In the absence of such data, local epidemiology and susceptibility patterns may contribute to the empiric selection of therapy.

CONTRAINDICATIONS

Cubicin is contraindicated in patients with known hypersensitivity to daptomycin.

WARNINGS

Pseudomembranous colitis has been reported with nearly all antibacterial agents, including daptomycin, and may range in

Daptomycin product labeling... http://www.cubicin.com/PDF/1004-2 MARKETING gt.pdf.

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is co-administered with probenecid.

MICROBIOLOGY

Daptomycin is an antibacterial agent of a new class of antibiotics, the cyclic lipopeptides. Daptomycin is a natural product which has clinical utility in the treatment of infections caused by aerobic Gram-positive bacteria. The *in vitro* spectrum of activity of daptomycin appropriate postage.

Daptomycin has been shown to be active against most isolates of the following microorganisms both in vitro and in clinical infections, as described in the INDICATIONS AND USAGE section.

Aerobic and facultative Gram-positive microorganisms:

Enterococcus faecalis (vancomycin-susceptible strains only)

Staphylococcus aureus (including methicillin-resistant

Streptococcus agalactiae

both in vitro and in clinical infections

Streptococcus dysgalactiae subsp. equisimilis

Streptococcus pyogenes

The following in vitro data are available, but their clinical significance is unknown. Greater than 90% of the following microorganisms demonstrate an in vitro MIC less than or equal to the susceptible breakpoint for daptomycin versus the bacterial genus. The efficacy or daptomycin in treating clinical infections due to these microorganisms has not been established in adequate and well-controlled clinical trials.

Aerobic and facultative Gram-positive microorganisms

in vitro

Corynebacterium jeikeium

Enterococcus faecalis (vancomycin-resistant strains)

Enterococcus faecium (including vancomycin-resistant strains)

Staphylococcus epidermidis (including methicillin-resistant strains)

Staphylococcus haemolyticus

Daptomycin product labeling...
http://www.cubicin.com/PDF/1004-2 MARKETING gt.pdf

Daptomycin FDA MIC Breakpoints (same as CLSI) 2006 - CLSI eliminated disk diffusion breakpoints

Table 3. Susceptibility Interpretive Criteria for Daptomycin

| Pathogen | Minimal inhibitory concentration (μg/mL) ^a | | | Disk diffusion zone Diameter (mm) ^b | | |
|--|--|-----|-----|---|-----|-----|
| | s | I | R | s | _ | R |
| Staphylococcus aureus (methicillin-susceptible and methicillin-resistant) | ⊴1 | (c) | (c) | ≥16 | (c) | (c) |
| Streptococcus pyogenes, Streptococcus agalactiae, and Streptococcus dysgalactiae subsp. equisimilis | ⊴1 | (c) | (c) | ≥16 | (c) | (c) |
| Enterococcus faecalis (vancomycin-susceptible only) | ≤4 | (c) | (c) | ≥11 | (c) | (c) |

- a. The MIC interpretive criteria for S. aureus and E. faecalis are applicable only to tests performed by broth microdilution using Mueller-Hinton broth adjusted to a calcium content of 50 mg/L; the MIC interpretive criteria for Streptococcus spp. other than S. pneumoniae are applicable only to tests performed by broth microdilution using Mueller-Hinton broth adjusted to a calcium content of 50 mg/L, supplemented with 2 to 5% lysed horse blood, inoculated with a direct colony suspension and incubated in ambient air at 35°C for 20 to 24 hours.
- The zone diameter interpretive criteria for Streptococcus spp. other than S. pneumoniae are applicable only to tests

Daptomycin product labeling...

http://www.cubicin.com/PDF/1004-2 MARKETING gt.pd20

Table 1. Suggested Groupings of Antimicrobial Agents With FDA Clinical Indications That Should Be Considered for Routine Testing and Reporting on Nonfastidious Organisms by Clinical Microbiology Laboratories in the U.S.

| ST RT | Enterobacteriaceae ^o | Pseudomonas aeruginosa and Other Non- Enterobacteriaceae ¹ | Staphylococcus spp. | Enterococcus spp. | |
|---------------------------------------|--|---|-------------------------------------|-----------------------------|--|
| UP A Y TE :POF | Ampicillin ^a | Ceftazidime | Oxacill in ^t | Penicillin°or ampicillin | |
| GROUP A PRIMARY TEST AND REPORT | Cefazolin ^a Cephalothin ^a | Gentamicin | Penicillin ^t | 1 | |
| PRII | Gentamicin | Mezlocillin or ticarcillin | | | |
| | | Piperacillin | | | |
| | Amikacin | Amikacin | Azithromycin ^d or | Daptomycin | |
| | | | clarithromγcin ^d or | Linezolid Quinupristin- | |
| | | | · | dalfopristin ^q | |
| | Amoxicillin-clavulanic acid or ampicillin-sulbactam Piperacillin-tazobactam Ticarcillin-clavulanic acid | Cefepime | — erythromycin [₫] | Vancomycin | |
| | Cefamandole or | Aztreonam | Clindamycin ^a | | |
| OUP B° ARY TEST SELECTIVELY | cefonicid or cefuroxime | Cefoperazone Ciprofloxacin Levofloxacin | Daptomycin Linezolid Telithromycin® | - | |
| o Be TES ECTI | Cefepime | Imipenem Meropenem | CLSI Table | e 1 (M7) | |
| ROUP IARY SELE | Cefmetazole Cefoperazone | Ticarcillin-clavulanic acid ^k | Drugs to Te | | |

Why are some drugs listed in Table 2 but not in Table 1?

Table 1. Suggested Gro Be Considered for Ro CLSI Table 1 (M7)

Drugs to Test/Report

n FDA Clinical Indications That Should Nonfastidious Organisms by Clinical

Microbiology Laboratories in the U.S.

JUP A RY TEST REPORT

| Enterobacteriaceae ^a | <i>Pseudomonas aeruginosa</i> and Other Non- Enterobacteriaceae ¹ | Staphylococcus spp. | Enterococcus spp." |
|---------------------------------|--|-------------------------|-----------------------------|
| Ampicillin ^a | Ceftazidime | Oxacillin' | Penicillin°or ampicillin |
| Coforolin ^a | Contomicin | Dopioillip ¹ | |

| | Test/Report | rt Antimicrobial Agent | | MIC (µg/mL) Interpretive Standard | | | Comments | |
|--|--------------|-----------------------------|------|--------------------------------------|-----------|------------------------|---|--|
| | Group | | | S | | ¦ R | | |
| | PENICILLINS | | | | | • | | |
| | А | l Mezlocillin or | | ≤ 64 | - | ; ≥128 | For P. aeruginosa only | |
| <u> </u> | | - 4 (5 (-) | | ≤16 | 32-64 | ! ≥128 | For other non-Enterobacteriace ae | |
| | Ibla 2 | B-1 (M7) | | ≤ 64 | - | ; ≥128 | For P. aeruginosa only | |
| | | ענונו) ניים | | ≤16 | 32-64 | ¦ ≥128 | For other non-Enterobacteriace ae | |
| | | 4 9 | | ≤ 64 | - | ! ≥128 | For P. aeruginosa only | |
| | | eiterisicesie | | ≤16 | 32-64 | : ≥128 | For other non-Enterobacteriaceae | |
| Non-Enterobacteriaceae — | | | ≤128 | 256 | ≥ 512 | For P. aeruginosa only | | |
| - | | P 6 | | ≤16 | 32 | ≥64 | For other non-Enterobacteriace ae | |
| = 3 <mark>(=</mark> | 351 (0) (0) | Ints | | ≤ 64 | - | ≥128 | For P. aeruginosa only | |
| Single Single | | | | | | | | |
| | В | Ticarcillin-clavulanic acid | | ≤16/2 | 32/2-64/2 | : ≥ 128/2 | For other non-Enterobacteriaceae | |
| | | | | | | | (4) May be indicated for testing of some <i>Pseudomonas</i> spp. (other than <i>P. aeruginosa</i>) | |
| | 0 | Ticarcillin-clavulanic acid | | ≤ 64/2 | - | ≥ 128/2 | For P. aeruginosa only | |
| | | Pineracillin-tazohadam | | < 84/4 | | > 10874 | For P aeguninosa only | |

Possible Reasons why drug may be in Table 2 but not Table 1...

- Drug does not have an FDA clinical indication for organism
- Drug may not be used in the USA
- Drug is not a first-choice or alternative drug suggested for routine testing for organism

Example:

piperacillin-tazobactam and P. aeruginosa

Susceptible ("S") Reworded

- ♦ New -...implies that isolates are inhibited by the usually achievable concentrations of antimicrobial agent when the recommended dosage is used for the site of infection.
- ♦ Old -...implies that an infection due to the strain may be appropriately treated with the dosage of antimicrobial agent recommended for that type of infection and infecting species, unless otherwise contraindicated.

Intermediate ("I")

The "intermediate" category includes isolates with antimicrobial agent MICs that approach usually attainable blood and tissue levels and for which response rates may be lower than for susceptible isolates. The intermediate category implies clinical efficacy in body sites where the drugs are physiologically concentrated (e.g., quinolones and β-lactams in urine) or when a higher than normal* dosage of a drug can be used (e.g., β-lactams). This category also includes a buffer zone, which should prevent small, uncontrolled, technical factors from causing major discrepancies in interpretations, especially for drugs with narrow pharmacotoxicity margins.

*previously stated "high dosage"

Resistant ("R") Reworded

- New —...implies that isolates are not inhibited by the usually achievable concentrations of the agent with normal dosage schedules and/or that demonstrate MICs that fall in the range where specific microbial resistance mechanisms are likely (e.g., β-lactamases), and clinical efficacy of that agent against the isolate has not been reliably shown in treatment studies.
- Old ...strains are not inhibited by the usually achievable systemic concentrations of the agent with normal dosage schedules and/or fall in the range where specific microbial resistance mechanisms (e.g., β-lactamases) and clinical efficacy has not been reliable in treatment studies.
 CLSI M100-S16; Introduction (M2, M7) 26

"S" only Definition

ener occurrence

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If only "S" criteria are specified:

For some organism/antimicrobial combinations, 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. Subsequently, the isolates should be saved and submitted to a reference laboratory that will confirm results using a CLSI reference dilution method.

- "Absence" vs. "rare occurrence"....

 How do we know if there has EVER
 been a "NS" isolate?
 - Check CLSI M100-S16 "Suggestions for Verification of AST Results and Confirmation of Organism Identification" [Table 4 (M2) or Table 8 (M7)]

NS, not susceptible

Excerpt from: "Suggestions for Verification of AST Results and Confirmation of Organism Identification"

| Organism or Group | Category I Verify at all labs | Category II Verify – institution specific |
|--------------------------|---|---|
| Staphylococcus aureus | daptomycin - NS linezolid – NS quin-dalfo – I or R vancomycin – I or R | oxacillin – R |
| Streptococcus pneumoniae | fluoroquinolone - NS linezolid ^C – NS vancomycin ^C – NS | penicillin - R 3 rd -gen cephalosporin- R |

NS, not susceptible Note: superscript "c" means "never reported" ^cNS Example
NS, not susceptible

S. pneumoniae - Vancomycin^c MIC (µg/ml) Susc Int Res

vancomycin

≤ 1.0

_ .

- *investigate any NS isolate
- ..Repeat ID and AST
- ..Save isolate
- ..Send to reference lab (test by CLSI MIC method)

 Note: vancomycin-NS *S. pneumoniae* have NEVER been reported

NS Example NS, not susceptible

S. aureus - Linezolid MIC (µg/ml)

Susc Int Res

linezolid

 ≤ 4.0

*investigate any NS isolate

..Repeat ID and AST

..Save isolate

..Send to reference lab (CLSI MIC method)

Note: linezolid-NS *S. aureus* have been reported on rare occasions

CLSI M100-S16; Table 2C (M7)

CLSI Introduction to Tables Added ESBLs to "Warning" List

M100-S16

V. 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."

"Warning": The following antimicrobial agent/organism combinations may appear active *in vitro* but are not effective clinically and should not be reported as susceptible.

| Location | Organism | Antimicrobial Agents That Must Not |
|---------------------|---------------------------------------|---|
| | - | be Reported as Susceptible |
| Table 2A | ESBL-producing K. pneumoniae, | penicillins, cephalosporins, and |
| | K. oxytoca, E. coli, and P. mirabilis | aztreonam |
| Table 2A | Salmonella spp., Shigella spp | 1st- and 2nd-generation cephalcsporins, |
| | | cephamycins, and aminoglycosides |
| Table 2C | oxacillin-resistant Staphylococcus | penicillins, β-lactam/β-lactamase inhibitor |
| | spp. | combinations, cephems, and carbapenems |
| Table 2D | Enterococcus spp. | aminoglycosides (except high |
| | | concentrations), cephalospories, |
| | | clindamyc:n, and trimethoprim- |
| | | sulfamethcxazole |
| Table 2K (Table 2A) | Yersinia pestis | β-lactam antimicrobial agents |
| Table 7 | Listeria spp. | cephalosporins |

Clarification of Incubation Temperature Range

 35°C +/- 2°C (except staphylococci as shown below and *Neisseria gonorrhoeae*)

Table 2C. MIC Interpretive Standards (μq/mL) for Staphylococcus spp.

Testing Conditions

Medium: Broth dilution: Cation-adjusted Mueller-Hinton broth (CAMHB)

CAMHB plus 2% NaCl for oxacillin, methicillin, and nafcillin; CAMHB supplemented to 50 µg/mL calcium for daptomycin Agar dilution: Mueller-Hinton agar (MHA); Agar dilution **has not**

been validated for daptomycin.

MHA plus 2% NaCl for oxacillin, methicillin, and nafcillin.

Inoculum: Direct colony suspension, equivalent to a 0.5 McFarland standard

Incubation: 35 ± 2 °C; ambient air; 16 to 20 hours; 24 hours for exacillin,

methicillin, nafcillin, and vancomycin. Testing at temperatures

above 35 °C may not detect MRS.

Minimal

ranges.)

Staphylo Escherio combina

Staphylo ATCC® E test)

CLSI M45-P Guideline

"Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently-Isolated or Fastidious Bacteria"





CLSI M45-P Guideline

- "Guideline" vs. "Standard"
- ◆ M45 is....
 - Based upon data in published literature
 - Based on MIC distributions and resistance mechanisms of organisms

Limited clinical data available to support decisions

- M100 is a "standard" and is based on substantial clinical data in addition to in vitro data (see CLSI M23)
- "P" = proposed; will likely become M45-"A" (A= approved)

On inside cover of each CLSI document is description of Standard, Guideline, Proposed, Approved and more....

Clinical and Laboratory Standards Institute

Providing NCCLS standards and guidelines, ISO/TC 212 standards, and ISO/TC 76 standards

The Clinical and Laboratory Standards Institute (CLSI, formerly NCCLS) is an international, interdisciplinary, nonprofit, standards-developing, and educational organization that promotes the development and use of voluntary consensus standards and guidelines within the healthcare community. It is recognized worldwide for the application of its unique consensus process in the development of standards and guidelines for patient testing and related healthcare issues. Our process is based on the principle that consensus is an effective and cost-effective way to improve patient testing and healthcare services.

In addition to developing and promoting the use of voluntary consensus standards and guidelines, we provide an open and unbiased forum to address critical issues affecting the quality of patient testing and health care.

PUBLICATIONS

A document is published as a standard, guideline, or committee report.

Standard A document developed through the consensus process that clearly identifies specific, essential

Most documents are subject to two levels of consensus— "proposed" and "approved." Depending on the need for field evaluation or data collection, documents may also be made available for review at an intermediate consensus level.

Proposed A consensus document undergoes the first stage of review by the healthcare community as a proposed standard or guideline. The document should receive a wide and thorough technical review, including an overall review of its scope, approach, and utility, and a line-by-line review of its technical and editorial content.

Approved An approved standard or guideline has achieved consensus within the healthcare community. It should be reviewed to assess the utility of the final document, to ensure attainment of consensus (i.e., that comments on earlier versions have been satisfactorily addressed), and to identify the need for additional consensus documents.

Our standards and guidelines represent a consensus opinion on good practices and reflect the substantial agreement by materially affected, competent, and interested parties obtained by following CLSI's established consensus procedures. Provisions in CLSI standards and guidelines

CLSI M45-P Guideline

| Abiotrophia / Granulicatella | Lactobacillus |
|------------------------------|----------------------------|
| *Aeromonas / Plesiomonas | Leuconostoc |
| Bacillus spp. (not anthrax) | Listeria monocytogenes |
| Campylobacter jejuni / coli | Moraxella catarrhalis |
| Corynebacterium | *Pasteurella |
| Erysipelothrix | Pediococcus |
| HACEK Group | *Vibrio spp. (not cholera) |

^{*}disk diffusion method described in addition to MIC method

CLSI M45-P Guideline

Testing should only be undertaken in consultation with infectious diseases or other expert clinicians that can assist in determining if susceptibility testing is needed in the management of a specific patient."

CLSI M45-P; Indications

<u>Table 5, Corynebacterium</u> spp. (Including *C. diphtheriae*)—Information and Interpretive Criteria for Broth Microdilution Susceptibility Testing

Testing Conditions

Medium Cation adjusted Mueller-Hinton broth with lysed horse

blood (2.5-5%)

Inoculum Direct colony suspension, equivalent to a 0.5 Mc Farland

standard

Incubation 35 °C; ambient air for 24-48 hours

Minimal QC Recommendations

Streptococcus pneumoniae ATCC= 49619 Agents to Consider for Primary Testing

Penicillin Vancomycin Erythromycin Gentamicin

General Comments

- (1) Growth characteristics on routine media:
 Nonfastidious; grows well on BAP; ambient air; 20-24 hours
- For some organism/antimicrobial combinations, 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. Subsequently, the isolates should be saved and submitted to a reference laboratory for confirmation.

NOTE: Information in boldface type is considered tentative for one year.

| Antimierobial | Antimicrobial Agent | MIC (mg/mL) Interpretive Criteria | | riteria | Comments | |
|---------------|---------------------|--------------------------------------|--------|---------|--------------|--|
| Class | | S | l I | J | ¦ R | |
| PENICILLINS | | | | | | (3) Interpretive oritena may not apply to meningitis |
| | Penicillin | 51 | 1 | 2 | ; 24 | |
| CEPHEMS | | | | | • | (3) Interpretive ortena may not apply to meningtis |
| | Cefepime | _51 | | 2 | <u>† 24</u> | |
| | Cefotaxime | 51 | l | 2 | ; 24 | |
| | Cettraxone | 51 | I | 2 | 1 24 | |
| CARBAPENEM | IS | | | | | (3) Interpretive ortena may not apply to meningtis |
| | lmipenem | 54- | I | 8 | ı 216 | CI CI MAR D Table R |
| | Meropenem | _≤4 | 1 | 8 | <u>† 216</u> | CLSI M45-P Table 5 |
| GLYC OPEPTIC | | | | | | |
| | Vancomycin | 54- | 1 | - | ! - | Corynebacterium spp. 3 |
| | Teicoplanin | 58 | ı | 16 | 232 | |

CLSI M45-P Table 5 (con't) Corynebacterium spp.

Supplemental Information

Resistance:

Some species of Corynebacterium may exhibit resistance to multiple drug classes.

Reasons for Testing/Not Testing:

Testing of isolates from normally sterile sources (blood cultures, deep tissue, implanted prosthetic devices) may be warranted, especially in immunodeficient patients.

Derivation of Interpretive Criteria:

Interpretive criteria for penicillin and enythromycin are based primarily on MIC distributions following testing of a large number of isolates. Cephalosporin interpretive criteria are adapted from those for *Streptococcus* spp.; linezolid interpretive criteria are adapted from those for *Staphylococcus* spp.; remaining interpretive criteria are adapted from those for *Staphylococcus* spp. as published in the current edition of CLSI document M100.

Testing Notes:

Resistant results can be reported at 24 hours. Isolates demonstrating susceptible results for beta-lactams should be reincubated and results reported at 48 hours.

Changes 2006 CLSI M100-S16 Gram Negatives



Enterobacteriaceae

CLARIFICATIONS...

- ESBL screening breakpoints for *Proteus* mirabilis
- Warning comment for Salmonella/Shigella
- AST of Salmonella spp. from feces

Proteus mirabilis ESBL Screen Test

| Drug | Disk Screen (mm) | MIC Screen (μg/ml) |
|-------------|---------------------|-----------------------|
| Cefpodoxime | ≤22* | >1* |
| Ceftazidime | ≤22 | >1 |
| Aztreonam | NA | NA |
| Cefotaxime | ≤27 | >1 |
| Ceftriaxone | NA | NA |

*unique for *P. mirabilis* (as compared to ≤17 mm and >4 μg/ml for *E. coli* and *Klebsiella* spp.); NA, not applicable

"Warning" Comment Salmonella/Shigella

◆ "Warning: For Salmonella and Shigella spp., aminoglycosides, 1st - and 2nd -generation cephalosporins and cephamycins may appear active in vitro but are not effective clinically and should not be reported as "S".

| | | ticarcillin-clavulanic acid |
|----------------------|--|--|
| cephems (parenteral) | cephalosporin Isa | cefazolin |
| | | cephalothin. |
| | | cephapirin |
| | | cephradine |
| | cephalosporin II 68 | cefamandole |
| | | cefonicid |
| | | cefuroxime (sodium) |
| | cephalosporin III ca | cefoperazone |
| | | cefotaxime |
| ssarv | | ceftazidime |
| | | ceftizoxime |
| † 1) | | ceftriaxone |
| | | cefepime |
| ems | cephamycin ⁴ | cefmetazole |
| | | cefotetan |
| | | cefo xitin |
| | | moxalactam |
| cephems (oral) | cephalosporin ^a | cefac lor |
| | | cefadro xil |
| | | cefdinir |
| | | cefditoren |
| | | cefetamet |
| | | cefixime |
| | | cefpodoxime |
| | | cefprozil |
| | | ceftibuten |
| | | cefuroxime (axetil) |
| | | cephalexin |
| | | cephradine 4.5 |
| | carbacephem | loracarb ef |
| | cephems (parenteral) ssary 1 1) ms cephems (oral) | cephalosporin III cephalosporin III cephalosporin III cephalosporin IV c. cephalosporin IV c. cephamycin cephamycin cephamycin cephamycin cephamycin cephalosporin cephalo |

Salmonella Reporting

♦ When fecal isolates of Salmonella and Shigella spp. are tested only ampicillin, a quinolone, and trimeth-sulfa should be tested and reported routinely. In addition, chloramphenicol and a 3rd-generation cephalosporin should be tested and reported for extraintestinal isolates of Salmonella spp.

Salmonella spp. (feces)

ampicillin S

ciprofloxacin S

trimeth-sulfa S

Notes:

- Add chloramphenicol (if your MDs want this) and 3rdgeneration cephalosporin for extraintestinal isolates
- May not need to do AST on Salmonella spp. isolated from patients with mild diarrhea as disease is self-limiting

Non-Enterobacteriaceae

- Added separate breakpoint tables:
 - Pseudomonas aeruginosa (disk diffusion)
 - Pseudomonas aeruginosa and other non-Enterobacteriaceae (MIC)
 - Acinetobacter spp.
 - Burkholderia cepacia
 - Stenotrophomonas maltophilia
- ◆ Delete P. aeruginosa "Rx" comment
- Colistin / polymyxin B
 - Deleted disk diffusion QC ranges
 - Added colistin MIC breakpoints for Acinetobacter spp.

CLSI Table 1 (M7) Drugs to Test/Report

January 2006 Vol. 26 No. 3

Table 1. Suggested Groupings of Antimicrobial Agents With FDA Clinical Indications That Should Be Considered for Routine Testing and Reporting on Nonfastidious Organisms by Clinical Misrobial and Indications in the U.S.

Microbiology Laboratories in the U.S.

GROUP A PRIMARY TEST AND REPORT

| Enterobacteriaceae ^a | Pseudomonas aeruginosa and Other Non- Enterobacteriaceae ¹ | Staphylococcus spp. | Enterococcus spp." |
|--|---|---------------------|-----------------------------|
| Ampicillin ^a | Ceftazidime | Oxacillin' | Penicillin°or ampicillin |
| Cefazolin ^a Cephalothin ^a | Gentamicin | Penicillin' | |
| Gentamicin | Mezlocillin or ticarcillin Pineracillin | | |

For Use With M7-A7-MIC Testing

M100-S16

Table 1. (Continued)

| | A TEST ORT | Acinetobacter.spp. | Burkholderia cepacia ¹ | Stenotrophomonas maltophilia ¹ |
|-----|--------------------|-----------------------|-----------------------------------|---|
| | م نے ق | Ceftazidime | Trimethoprim- sulfamethoxazole | Trim ethoprim-sulfamethoxazole |
| | REI REI | lmipenem Meropenem | | |
| | GF PRIIM AND | | | |
| ++- | | | | |
| ** | | Amikacin | Ceftazidime | Ceftazidime |

Table 2B-4 (M7) MIC Breakpoints for Stenotrophomonas maltophilia

Table 2B.4. MIC Interpretive Standards (µg/mL) for Breakpoints for Stenotrophomonas maltophilia

Testing Conditions

Medium: Broth dilution: Cation-adjusted Mueller-Hinton broth (CAMHB)

Agar dilution: Mueller-Hinton agar (MHA)

Inoculum: Growth method or direct colony suspension, equivalent to a 0.5

McFarland standard

Incubation: 35±2°C; ambient air; 20 to 24 hours

Minimal QC Recommendations (See Table 3 for Acceptable QC Ranges.)

Pseudomonas aeruginosa ATCC® 27853

Escherichia coli ATCC® 25922

Escherichia coli ATCC® 35218 (for ß-lactam/ß-lactamase inhibitor

combinations)

General Comments

(1) Some of these agents may require testing in those institutions that harbor endemic or epidemic strains resistant to other drugs or for reporting to infection control as an epidemiologic aid.

| # | NOTE: Ir | nformation in boldface type is considered tentative for one year. | | | | | | | | |
|---|---------------------|---|--------------------------------------|----------------|-----------------|----------|--|--|--|--|
| П | Test/Report | Antimicrobial Agent | MIC (μg/mL) Interpretive Standard | | | Comments | | | | |
| | Group | | int | erpretive 5to | andard | | | | | |
| | | | S I R | | | | | | | |
| | β-LACTAM/β-L | ACTAMASE INHIBITOR COMBINATIONS | | | | | | | | |
| | В | Ticarcillin-clavulanic acid | ≤ 16/2 | 32/2-64/2 | ≥ 128/2 | | | | | |
| | CEPHEMS (PA | RENTERAL) (Including cephalosporins I, II, II | I, and IV. F | Please refer t | to Glossary I.) | | | | | |
| | В | Cettazidime | ≤8 | 16 | ≥ 32 | | | | | |
| | TETRACYCLIN | IES | | | | | | | | |
| | В | Minocycline | ≤ 4 | 8 | ≥16 | | | | | |
| | FLUOROQUINOLONES | | | | | | | | | |
| | A | Levofloxacin | ≤2 | 4 | ≥8 | | | | | |
| | FOLATE PATH | FOLATE PATHWAY INHIBITORS | | | | | | | | |
| | А | Trimethoprim-sulfamethoxazole | ≤ 2/38 | - | ≥ 4/76 | | | | | |

S. maltophilia (blood)

MIC (μg/ml)

| lovoflovosin | 9 | 7 |
|--------------|------------|---|
| levofloxacin | Z 3 | |

minocycline 1 S

ticarcillin-clav 16 S

trimeth-sulfa 0.5/9.5 S

These are the only drugs for which there are MIC breakpoints in M100-S16 Table 2B-4 (M7)

What if MD asks for results for other drugs on S. maltophilia?

- Option to consider.....
- Get request in writing, preferably from Infectious Diseases clinician
- Test by MIC only
- Report results without interpretation
- Qualify results

S. maltophilia (blood)

Report Option

MIC (μg/ml)

*aztreonam 8
ceftazidime 32 R
levofloxacin 2 S
minocycline 1 S
ticarcillin-clav 16 S
trimeth-sulfa ≤0.5/9.5 S

*No MIC interpretive criteria available; reported per Dr. Jones request; Infectious Diseases consult suggested

Rx: Comment Deleted Pseudomonas aeruginosa

"Rx: P. aeruginosa infections in granulocytopenic patients and serious infections in other patients should be treated with maximum doses of the selected antipseudomonal penicillin (carboxypenicillin or ureidopenicillin) or ceftazidime in combination with an aminoglycoside."

Rationale for deletion – there are currently other options for treatment

Colistin / Polymyxin B

- Currently, NO disk diffusion recommendations
- Acinetobacter spp.

| Antimicrobial | MIC (µg/ml) | | | | |
|-------------------------|-------------|---|----|--|--|
| Agent | S | I | R | | |
| Polymyxin B or colistin | ≤2 | - | ≥4 | | |

- Breakpoints for other bugs forthcoming
- No FDA-cleared test

Haemophilus spp. Neisseria meningitidis

- Procedures in M2, M7 and M100 are for *H. influenzae* and *H. parainfluenzae*
 - CLSI M45-P for other *Haemophilus* species
- Neisseria meningitidis
 - Added disk diffusion procedure

Standard Disk Diffusion Method for Neisseria meningitidis



<u>Table 2.1.</u> Zone Diameter Interpretive Standards and Equivalent Minimal Inhibitory Concentration (MIC) Breakpoints for Neisseria meningitidis

Testing Conditions

Medium: Mueller-Hinton agar with 5% sheep blood

Inoculum: Direct colony suspension, equivalent to a

0.5 McFarland standard

Incubation: 35 ± 2 °C; 5% CO₂; 20 to 24 hours

Minimal QC Recommendations (See Tables 3 and 3A for acceptable QC ranges.)

Streptococcus pneumoniae ATCC® 49619 in 5% CO₂. E. coli ATCC® 25922 (incubated either in ambient air or 5% CO₂) should be used for ciprofloxacin, nalidixic acid, and minocycline.

General Comments

- (1) Recommended precautions: Biosafety Level 2 (BSL2) practices are recommended for this organism. Whenever possible, procedures likely to generate aerosols should be performed within a biological safety cabinet.
- (2) For some organism/antimicrobial agent combinations, 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. Subsequently, the isolates should be saved and submitted to a reference laboratory that will confirm results using a CLSI reference dilution method.

NOTE: Information in boldface type is considered tentative for one year.

| # | | | | | | |
|---|-------------|---------------|---------|------------------------------------|---------------------------------------|----------|
| | Test/Report | Antimicrobial | Disk | Zone Diameter, Nearest Whole mm | Equivalent MIC Breakpoints (µg/mL) | Comments |
| | Group | Agent | Content | RIIS | RS | |

Neisseria meningitidis Disk Diffusion Testing

Medium: MHA with 5% sheep blood

Inoculum: direct colony suspension equivalent

to 0.5 McFarland standard

Incubation: 35°C +/- 2°C; 5% CO₂; 20-24h

QC: S. pneumoniae ATCC 49619

E. coli ATCC 25922 (select drugs)

Biosafety Level 2 (BSL 2) safety practices; use biosafety cabinet

Neisseria meningitidis Disk Diffusion Testing

- Therapeutic agents*
 - *Penicillin
 - *Ampicillin
 - Cefotaxime
 - Ceftriaxone
 - Meropenem
 - Chloramphenicol

All drugs listed are in Test/Report Group C "Supplemental, Report Selectively"

*No disk diffusion breakpoints; must do MIC test

Neisseria meningitidis Disk Diffusion Testing

- "Breakpoints may be appropriate only for prophylaxis of meningococcal case contacts"
 - Azithromycin
 - Ciprofloxacin
 - Minocycline
 - Nalidixic acid (for surveillance only; may detect diminished fluoroquinolone susceptibility)
 - Rifampin
 - Trimethoprim-sulfamethoxazole (predicts susceptibility to sulfonamides also)

All drugs listed are in Test/Report Group C "Supplemental, Report Selectively"

Changes 2006 CLSI M100-S16 Gram Positives



Daptomycin

For....

- Staphylococcus
- Enterococcus
- Streptococcus
- Deleted daptomycin disk diffusion breakpoints
 - Rationale for deletion inability of disk diffusion method to consistently detect those few isolates that are non-susceptible to daptomycin
- Specify agar dilution testing has not been validated for daptomycin
- FDA-cleared for several commercial systems

Staphylococcus spp.

Clarifications...

- For disk diffusion testing, cefoxitin disk is...
 - Preferred over oxacillin disk for detection of mecAmediated resistance
 - A "surrogate" for oxacillin (report oxacillin NOT cefoxitin)
 - Should always be used for S. lugdunensis (do not use oxacillin disk)
- Fluoroquinolones breakpoints for gatifloxacin, levofloxacin, moxifloxacin, and ofloxacin still tentative (for another year)

Cefoxitin Disk for *mecA*-mediated Resistance in Staphylococci

Cefoxitin zone (mm)

Res

Susc

S. aureus

≤19*

≥20**

S. lugdunensis

≤19*

≥20**

CoNS

≤24*

>25**

- * Report as oxacillin resistant
- ** Report as oxacillin susceptible CoNS, coagulase-negative staphylococci

MIC Breakpoints (μg/ml) Staphylococcus and Fluoroquinolones

| | Old (I | M100-S | 514)* | M100-S15, M100-S16 ** | | |
|--------------|--------|--------|-------|-----------------------|---|----|
| | S | I | R | S | I | R |
| Gatifloxacin | ≤2 | 4 | ≥8 | ≤0.5 | 1 | ≥2 |
| Levofloxacin | ≤2 | 4 | ≥8 | ≤1 | 2 | ≥4 |
| Moxifloxacin | | none | | ≤0.5 | 1 | ≥2 |

^{*}Same as current FDA breakpoints

^{**}Tentative for another year Check M100-S16 for corresponding disk diffusion breakpoints

Staphylococcus spp.

Additions/changes...

- Modified vancomycin MIC breakpoints for S. aureus
- Table highlighting use of BHI-vancomycin agar for S. aureus

CLSI Vancomycin MIC (µg/ml) Breakpoints – S. aureus

2006 ≤2 4-8 ≥16

2005 ≤4 8-16 ≥32

Notes:

2005 same as current FDA breakpoints
No change in disk diffusion breakpoints
No change for coagulase-negative staphylococci

Why did CLSI modify vancomycin breakpoints for S. aureus?

- To detect emerging vancomycin resistance
 - Data shows patients fail vancomycin therapy when infected with *S. aureus* with vancomycin MICs of ≥4 µg/mI
 - Clinical labs have been advised by CDC to investigate S. aureus with vancomycin MICs of ≥4 μg/ml as potential VISA or VRSA
 - Some VISA test 4 μg/ml by some methods

Modifying CLSI Breakpoints

CLSI has mechanism for reevaluation of breakpoints when needed as defined in CLSI M23-A2 "Development of In Vitro Susceptibility Testing Criteria and QC Parameters"

Will it matter what vancomycin breakpoints we use for S. aureus?

- "Either FDA or CLSI susceptibility interpretive breakpoints are acceptable to clinical laboratory accrediting bodies" (CLSI M100-S16, page 14)
- ...and we should all pursue S. aureus with vancomycin MICs ≥4 µg/ml since these could be VISA (or VRSA if ≥16 µg/ml) http://www.cdc.gov/ncidod/dhqp/ar_visavrsa_algo.html
- So, if we ALL diligently pursue vancomycin MICs of ≥4 μg/ml in S. aureus, it really won't matter

S. aureus - Vancomycin MICs* UCLA 1/00-10/05 (n=13981)

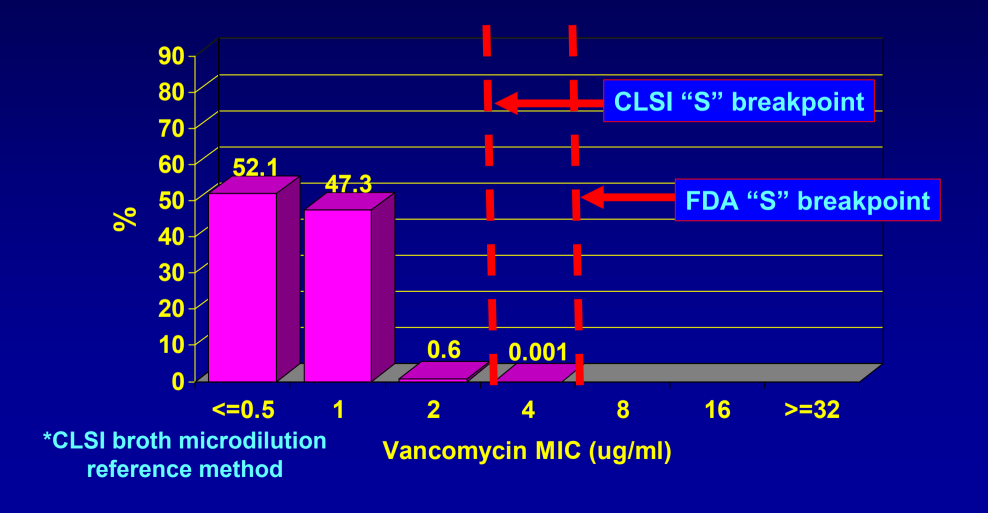


Table 2C. (Continued)

Screening Tests for Oxacillin Resistance and Reduced Susceptibility to Vancomycin in Staphylococcus aureus

| Screen Test | Oxacillin Resistance | Reduced Susceptibility to Vancomycin |
|--------------------------|---|---|
| Medium | MHA with NaCl (4% w/v; 0.68 mol/L) | BHI agar |
| Antimicrobial | 6 µg/mL oxacillin | 6 μg/mL vancomycin |
| concentration | | |
| Inoculum | Direct colony suspension to obtain 0.5 | Direct colony suspension to obtain 0.5 McFarland |
| | McFarland turbidity | turbidity. |
| | Lloing a 1 vl. loop that was dispard in the | Preferably, using a micropipette, spot a 10 µL drop |
| BHI-Vancomycin (6 µg/ml) | | onto agar surface. Alternatively, using a swab |
| REII-Asincom/c | ipped | dipped in the suspension and the excess liquid |
| | | expressed, spot an area 10 to 15 mm in diameter or |
| | nt. | streak a portion of the plate. |
| Screen | | 35 °C; ambient air |
| 0012211 | | 24 hours |
| results | 21 Colony = resistant | >1 colony = presumptive reduced susceptibility |
| | Examine carefully with transmitted light for >1 | Examine carefully with transmitted light for >1 |
| | colony or light film of growth. | colony or light film of growth. Perform vancomycin |
| | | MIC using a validated MIC method to confirm |
| | | reduced susceptibility |
| QC Recommendation | s Staphylococcus aureus ATCC® 29213 - | Enterococcus faecalis ATCC® 29212 - Susceptible |
| | Susceptible | Enterococcus faecalis ATCC® 51299 - Resistant |
| | Staphylococcus aureus ATCC® 43300 - | |
| | Resistant | |

- Add BHI-V screen if using automated system (unless fixed, check with manufacturer) or use CLSI MIC reference method or Etest to detect reduced susceptibility to vancomycin in S. aureus
- For workup of VISA and VRSA, see http://www.cdc.gov/ncidod/dhqp/ar_visavrsa_algo.html

VRSA (as of 12/05)

| | MIC (ug/ml) ¹ | Source | Date | Location |
|---|-----------------------------|------------------|-------|--------------|
| 1 | 1024 | foot ulcer | 4/02 | Michigan |
| 2 | 32 | foot ulcer | 9/02 | Pennsylvania |
| 3 | 64 | nephrostomy tube | 3/04 | New York |
| 4 | 256 | foot ulcer | 2/05 | Michigan |
| 5 | 512 | wound | 10/05 | Michigan |

¹ Reference broth microdilution MIC

Enterococcus spp.

Additions/changes...

- Deleted vancomycin-synergy Rx comment
- Added definitions for high-level aminoglycoside resistance (HLAR) testing for disk diffusion (similar to those for MIC testing)

Enterococcus Rx Comment Deleted

- Rx: If vancomycin is used for serious enterococcal infections, such as endocarditis, combined therapy with an aminoglycoside is usually indicated.
- Rationale for deletion use of vancomycin should not be encouraged; ampicillin or penicillin are the preferred agents to use in combination therapy for enterococci

Lab is ZD Enterococcus app. M2-Disk Diffusion

Enterococcus spp. Disk diffusion screen for HLAR

For Use With M2-A9-Disk Diffusion

M100-S16

Table 20. (Continued)

Disk Diffusion Screening Tests for High-Level Aminoglycoside Resistance (HLAR).

| ₩. | г | 4 | Ŀ | | |
|----|---|---|----|---|--|
| | | а | Į, | ۰ | |
| | • | э | Е | ۰ | |

| ᆜ | | | | | | |
|---|----------------------|------------------------|-----------------|--|--|---|
| | Test/Report Group | Antimicrobial Agent | Disk Content | Zone Diameter, Nearest Whole mm R ^h I ⁴ S ⁴ | Equivalent MIC Breakpoints (µg/mL) R ! S | Comments |
| | U | Gentamion (HLAR) | 120 µg | 6 7-9 210 | > 500 2 500 | (16) If the zone is 7 to 9 mm, the test is inconclusive, and an agar dilution or broth microdilution screen test should be performed to confirm resistance. See comments (2) and (7). |
| | C | Streptomyon (HLAR) | 300 µg | 6 7-9 2 10 | | (16) MIC correlates for streptomycin broth microdilution are resistant >1000 µg/mL and for agar dilution >2000 µg/mL. See comments (2), (7), and (16). |

Footnotes

- For QC of HLAR screen tests, use Enterococcus faecalis ATCC 29212 (see Table 3, Footnote f [Disk Testing] for acceptable QC ranges).
- b. Resistant, will not be synergistic with cell-wall-active agent (e.g., ampicillin, penicillin, vancomycin).
- Inconclusive, perform an agar dilution or broth microdilution test to confirm.
- g. Susceptible, will be synergistic with cell-wall-active agent (e.g., ampicillin, penicillin, vancomycin) that is also susceptible.

See M7, Table 2D (MIC Testing) which summarizes additional screening tests for vancomycin, high-level aminoglycoside resistance, and supplemental tests for identification that may be helpful for vancomycin-resistant enterococci.

Enterococcus spp. Disk diffusion Screen for HLAR

- Resistant -- will not be synergistic with cell-wall-active agent (e.g., ampicillin, penicillin, vancomycin)
- Inconclusive -- perform an agar dilution or broth microdilution test to confirm
- Susceptible -- will be synergistic with cell-wall-active agent (e.g., ampicillin, penicillin, vancomycin) that is also susceptible

Streptococcus pneumoniae Modified reporting recommendations for meropenem...

*"Penicillin and cefotaxime or ceftriaxone or meropenem should be tested by a reliable MIC method and reported routinely with CSF isolates of S. pneumoniae."

New Antimicrobial Agents in M100-S16*

| Agent | Drug class | Route of administration | FDA approved |
|--------------|------------|-------------------------|--------------|
| ceftobiprole | cephem | IV | No |
| faropenem | penem | PO | No |

*In Glossary and QC tables only

Ceftobiprole

- Manufacturer:
 - -Johnson & Johnson
- Possible clinical use:
 - -Nosocomial pneumonia
 - Complicated skin and skin structure infections

Ceftobiprole (con't)

Microbiological activity:

- Staphylococci (including MRSA)
- Streptococci (including penicillin-R S. pneumoniae)
- Enterococcus faecalis
- Most Enterobacteriaceae
- Haemophilus influenzae (including BLNAR)
- Many Pseudomonas aeruginosa and Acinetobacter baumannii

Limited activity:

- Many non-Enterobacteriaceae
- Enterococcus faecium
- ESBL-producers
- Metallo-beta-lactamase producers
- Beta-lactamase-producing anaerobes

Faropenem

- Manufacturer:
 - Replidyne
- Possible clinical use:
 - -Bacterial sinusitis
 - Acute exacerbations of chronic bronchitis
 - -Community-acquired pneumonia
 - Uncomplicated skin and skin structure infections

Faropenem (con't)

Microbiological activity:

- Respiratory pathogens (S. pneumoniae, H. influenzae, M. catarrhalis)
- S. pyogenes
- MSSA
- Some Enterobacteriaceae

Limited activity:

- Non-Enterobacteriaceae
- Enterobacter spp.
- Enterococcus faecium
- MRSA

Changes 2006 CLSI M100-S16 Quality Assessment / Quality Control



Quality Control

ADDITIONS / Changes...

- Added QC ranges:
 - Ceftobiprole
 - Faropenem
 - Drugs for testing Campylobacter jejuni ATCC
 33560 using broth dilution method

QC Tables

| Table | M2 (disk diffusion) | M7 (MIC testing) |
|------------|-------------------------------|---|
| 3 | Nonfastidious | Nonfastidious |
| 3 A | Fastidious | Fastidious (broth dilution) |
| 3B | DD QC Testing Frequency | Fastidious (agar dilution) New! |
| 3C | DD Troubleshooting Guide New! | Fastidious (broth dilution + supplement)* |
| 3D | NA | Fastidious (Brucella broth dilution)* |

DD, disk diffusion; NA, not applicable; *when testing potential agents of bioterrorism

QC Tables (con't)

| Table | M2 | M7 |
|-------|----|--------------------------------|
| 3E | NA | MIC QC Testing Frequency |
| 3F | NA | MIC Troubleshooting Guide New! |

^{*}NA, not applicable

Table 3C. Disk Diffusion QC Troubleshooting Guide

Table 3C. Disk Diffusion QC Troubleshooting Guide

This table provides guidance for troubleshooting and corrective action for out-of-range quality control, primarily using antimicrobial susceptibility tests with Mueller Hinton Agar. Refer to M2-A9, Disk Diffusion, Section 15, Quality Control Procedures and Appendix A, Quality Control Protocol Flow Charts for additional information. Out-of-range quality control tests should first be repeated. If the issue is unresolved, this troubleshooting guide provides additional suggestions for troubleshooting out-of-range quality control results. In addition, if unresolved, manufacturers should be notified of potential product problems.

General Comments

(1) QC organism maintenance: avoid repeated subcultures. Retrieve new QC strain from stock. If using lyophilized strains, follow the maintenance recommendations of the manufacturer. Store E. coli ATCC[®] 35218 and E. pneumoniae ATCC[®] 700603 stock cultures at -60°C or below and prepare working stock cultures weekly.

| Antimicrobial Agent | QC Strain | Observation | Probable Cause | Comments/Action |
|------------------------|--------------------------|----------------|---------------------------|-------------------------------|
| Aminoglycosides | Any | Zone too small | pH of media too low | Acceptable pH range = 7.2-7.4 |
| | | | | Avoid CO₂ incubation which |
| | | | | lowers pH. |
| Aminoglycosides | Any | Zone too large | pH of media too high | Acceptable pH range = 7.2-7.4 |
| Aminoglycosides | P. ceruginosa | Zone too small | Ca++ and/or Mg ++ | Use alternative lot of media. |
| | ATC C [©] 27853 | | content too high | |
| Aminoglycosides | Р. сетидіпови | Zone too large | Ca++ and/or Mg ++ | Use alternative lot of media. |
| | ATC C [®] 27853 | | content too low | QQ |
| Amoxicillin-clavulanic | E. coli ATCC® | Zone too small | Clavulanic acid is labile | Use alternative lot of disks. |
| acid | 35218 | | Disk has lost natency | Check storage conditions and |



Disk Diffusion QC Troubleshooting Guide Table 3C (M2) *Examples:*

| Antimicrobial Agent | QC Strain | Observation | Probable Cause | Comments / Action |
|------------------------|-----------|---|-------------------------|--|
| Beta-lactam group | Any | Zone initially acceptable but decreases and possibly out-of-range over time | Disk has lost potency | Use alternative lot of disks. Check storage conditions and package integrity. Imipenem, cefaclor, and clavulanic acid are especially labile. |
| Quinolones | Any | Zone too large | pH of media too high | Acceptable pH range =7.2=7.4 |

CLSI Standards - 2006

♦ M100-S16 Tables (2006)*

.....to be used with text documents explaining how to perform the tests....

M2-A9 Disk Diffusion (2006)**

M7-A7 MIC (2006)**



**M2, M7 updated every 3 years



M2-A9, M7-A7 Primary Changes

- Primarily expanded discussions and detailed recommendations for test procedures in M100-S16 including those for:
 - Oxacillin-resistant staphylococci
 - Streptococcus pneumoniae
 - Streptococcus spp.
 - Neisseria meningitidis

M2-A9, M7-A7 Primary Changes (con't)

- Additions to antimicrobial agent descriptions
- Additional tips for media/reagent preparation
- Supplemental QC suggestions

Last pages of M2 and M7

Summary of Comments and Subcommittee Responses

Clinical and Laboratory Standards Institute consensus procedures include an appeals process that is described in detail in Section 8 of the Administrative Procedures. For further information, contact CLSI or visit our website at www.clsi.org.

Summary of Comments and Subcommittee Responses

M2-A8: Performance Standards for Antimicrobial Disk Susceptibility Tests; Approved Standard—Eighth Edition

<u>General</u>

- 1. Could you please clarify a point for me about cefuroxime testing? Your standard gives different zone size criteria for the parenteral and oral forms of the drug and specifies that different discs should be used. Is the implication that you could get an isolate testing sensitive with the one disk and resistant with the other? If this is true, could you use the parenteral (cefuroxime sodium) disk to predict sensitivity for both forms, accepting that you may miscall some strains resistant that would respond to oral, or is it the other way around? The standard gives no details about these particular recommendations.
- Although there are two formulations for cefuroxime, one for parenteral and one for oral
 administration, there is only one disk for laboratory testing. Different interpretive criteria were
 developed based on the different pharmacodynamic/pharmacokinetic data and clinical
 indications for the two formulations. Tables 2A through 2J should be used to guide
 interpretation for individual organisms.

Q&A Example (paraphrased)

- Q Why is D zone test not recommended for Streptococcus pneumoniae?
- ◆ A Isolates of Streptococcus pneumoniae can have erm-mediated resistance to erythromycin. However, the vast majority of these are also resistant to clindamycin (constitutive phenotype). Rare isolates of pneumococci may have inducible resistance; however the clinical significance of this has not been established. Therefore, routine testing for inducible clindamycin resistance is not recommended for this species.

Issues Under Discussion by CLSI

- Additional recommendations for testing colistin / polymyxin B
- Disk diffusion test for Campylobacter spp.
- Detection of ESBLs
- Review recommendations for drugs to Test / Report (Table 1)
- Improved communication of CLSI AST Subcommittee decisions

Material on your CD-ROM...

- 1. PowerPoint presentation
- 2. PDF of "New Antimicrobial Agent Pathway" slide (slide #10)
- **3.** M100-S16 checklist
- 4. References
- **5.** CLSI information flier
- 6. CLSI catalogue

To Ask a Question....

- Please send to neoffice@nltn.org
- Questions will be compiled and answers will be published on http://www.phppo.cdc.gov/nltn/nphtc s/ast012506.aspx

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