

Designation: E2149 – 01

Standard Test Method for Determining the Antimicrobial Activity of Immobilized Antimicrobial Agents Under Dynamic Contact Conditions¹

This standard is issued under the fixed designation E2149; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ε) indicates an editorial change since the last revision or reapproval.

1. Scope

1.1 This test method is designed to evaluate the resistance of non-leaching antimicrobial treated specimens to the growth of microbes under dynamic contact conditions. This dynamic shake flask test was developed for routine quality control and screening tests in order to overcome difficulties in using classical antimicrobial test methods to evaluate substratebound antimicrobials. These difficulties include ensuring contact of inoculum to treated surface (as in AATCC 100), flexibility of retrieval at different contact times, use of inappropriately applied static conditions (as in AATCC 147), sensitivity, and reproducibility. This test also allows for the versatility of testing contamination due to such things as hard water, proteins, blood, serum, various chemicals, and other contaminates or physical/chemical stresses or manipulations of the specimens of interest.

1.2 Surface antimicrobial activity is determined by comparing results from the test sample to simultaneously run controls.

1.3 The presence of a leaching antimicrobial is both preand post-determined by the presence of a zone of inhibition.

1.4 This test method should be performed only by those trained in microbiological techniques.

1.5 This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 ASTM Standards:²

E1054 Test Methods for Evaluation of Inactivators of Antimicrobial Agents

2.2 Other Documents:

- AATCC Test Method 147-1998 Antibacterial Activity Assessment of Textile Materials: Parallel Streak Method. American Association of Textile Chemists and Colorists, RTP, NC
- AATCC Test Method 100-1999 Antibacterial Finishes on Fabrics, Evaluation of American Association of Textile Chemists and Colorists, RTP, NC

3. Summary of Test Method

3.1 Immobilized antimicrobial agents, such as surface bonded materials, are not free to diffuse into their environment under normal conditions of use. Test methods such as AATCC 147 that are directly dependent on the ready leachability of the antimicrobial agent from the treated fabric are inappropriate for evaluating immobilized antimicrobial agents. The following test method ensures good contact between the bacteria and the treated fiber, fabric, or other substrate by constant agitation of the test specimen in a bacterial suspension during the test period. The test is suitable for evaluating stressed or modified specimens when accompanied by adequate controls.

NOTE 1—Stresses may include laundry, wear and abrasion, radiation and steam sterilization, UV exposure, solvent manipulation, temperature susceptibility, or similar physical or chemical manipulation.

4. Significance and Use

4.1 The antimicrobial activity of a substrate-bound antimicrobial is dependent upon direct contact of microbes with the active chemical agent. This test determines the antimicrobial activity of treated specimen by shaking samples of surface bound materials in a concentrated bacterial suspension for a one hour contact time or other contact times as specified by the investigator. The suspension is serially diluted both . before and after contact and cultured. The number of viable organisms in the suspension is determined and the percent reduction is calculated based on initial counts or on retrievals from appropriate untreated controls.

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¹ This test method is under the jurisdiction of ASTM Committee E35 on Pesticides and is the direct responsibility of Subcommittee E35.15 on Antimicrobial Agents.

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² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

Note 2—This method is intended for those surfaces having a percent reduction activity of 50 % to 100 % for the specified contact time.

5. Apparatus

5.1 Sterilizer,

5.2 Incubator,

5.3 Spectrophotometer,

5.4 *Shaker, Wrist Action*—A Wrist Action Shaker is recommended but other means of agitation such as reciprocal action shakers may be satisfactory for routine testing. Shaker must ensure good agitation. Rotary shakers are unacceptable.

5.5 Water Bath,

5.6 Vortex Mixer,

5.7 Glassware,

5.7.1 Contact Flask, 250 ml Erlenmeyer flask, capped, autoclavable.

5.7.2 Dilution Vessels,

5.7.3 Pipettes,

5.8 Agar, bore 8-mm diameter.

6. Reagents

6.1 Buffer Solution-For test specimen which might alter the pH of the system, Sorensen's Phosphate Buffer (pH 6.8) is recommended.³ Other appropriate buffers must be shown not to cause a reduction or increase in bacterial numbers by prior testing at the intended use concentration. For all other samples, the following solution is recommended and is prepared from reagent grade chemicals. For buffer stock solution (0.25M KH_2PO_4): Prepare a fresh solution at least once every 6 months as follows: Weigh 34 ± 0.1 g of potassium dihydrogen phosphate into a 100 ml beaker. Add 500 ml of distilled water. Adjust pH to 7.2 \pm 0.1 with a dilute solution of NaOH. Dilute to 1000 ml; transfer to a flask and store at 4°C. For working buffer solution (0.3mM KH₂PO₄): Prepare a fresh solution at least once every 2 months as follows: Transfer 1 ml \pm 0.01 ml of stock buffer solution, with a sterile pipette to flask containing 800 ml of distilled water. Cap and sterilize.

6.2 *Media*:

6.2.1 *Nutrient Broth* (Difco Laboratories, Detroit, MI or equivalent) or media appropriate of organism selected.

6.2.2 *Tryptone Glucose Extract Agar* (Difco Laboratories, Detroit, MI or equivalent) or media appropriate for organism selected.

6.3 *Wetting Agent Surfactant*—Agents must be shown not to cause a reduction or increase in bacterial numbers by prior testing at the intended use concentration.

Note 3—Dow Corning, Midland, MI Q2-5211 at 0.01 % final dilution or equivalent can be used.

7. Test Organism

7.1 *Klebsiella pneumoniae*, American Type Culture Collection No. 4352. Other organisms may be used at the discretion of the investigator.

7.1.1 Cultures of the test organism should be maintained according to good microbiological practice and checked for

purity, on a routine basis. Consistent and accurate testing requires maintenance of a pure, uncontaminated test culture. Avoid contamination by use of good sterile technique in plating and transferring. Avoid mutation or reversion by strict adherence to monthly stock transfers. Check culture purity by making streak plates periodically and observing for a single species characteristic type of colonies.

NOTE 4—A glossiness in the broth culture of Klebsiella pneumoniae is a sign of reversion and the culture should not be used.

8. Parameters

8.1 Aerobic organisms and/or contact times used must be specified.

8.2 Surface preparation or conditioning must be specified. Prior manipulation of the specimen in order to demonstrate maximum activity in desired time frame must be recorded and compared to identically handled controls.

8.3 The weight, size, and material of construction of specimen must be specified.

8.4 Specimens should be prepared such that they can maximize agitation and are reflective of a recordable ratio of surface area to test titer.

8.5 Wetting agent surfactants must be used with highly hydrophobic specimen.

9. Preparation of Bacterial Inoculum

9.1 Grow a fresh 18 hour shake culture of *Klebsiella pneumoniae*Klebsiella pneumoniae in sterile nutrient broth for each series of samples. If other organisms are specified, they should be prepared in the same manner, unless other media and different calibration techniques are specified.

9.2 Dilute the culture with the sterile buffer solution until the solution has an absorbance of 0.28 ± 0.01 at 475 nm, as measured spectrophotometrically. This has a concentration of $1.5-3.0 \times 10^8$ CFU/ml. Dilute appropriately into sterile buffer solution to obtain a final concentration of $1.5-3.0 \times 10^5$ CFU/ml. This solution will be the working bacterial dilution.

Note 5—For other organisms, adjust final concentration to 1.5-3.0 \times 10^5 CFU/ml by appropriate methods.

10. Test Specimen

10.1 Preparation of Test Specimen:

10.1.1 *Fabric and Paper*—Samples are selected on weight basis at the discretion of the investigator and weighed to \pm 0.1 g.

NOTE 6—Weight, usually between 0.5 and 2.0 g, must ensure strong agitation during contact period. Specimen should be cut into small enough portions to ensure maximum agitation and must be identical in size between treated and untreated controls. Clumping of specimen negatively affects reproducibility.

10.1.2 *Powder and Granular Material*—Weigh to \pm 0.1 g. The material must settle after shaking so that no specimen interferes with the retrieval and counting techniques.

10.1.3 Other Solids (Surface Treatment)—Reduce the solid in size to fit into the flask or use a sterile wide-mouth bottle. Use a specimen that gives 9 sq. in. (58 cm²) of treated surface area. Specimen may also be selected on weight basis at the discretion of the investigator, weigh to \pm 0.1 g. Care must be

 $^{^{3}}$ Clinical Chemistry: Principle and Technics, Second Edition 1974, Table A-3e, p 1592.