



Designation: E1173 – 23

Standard Practice for Evaluation of Preoperative, Precatheterization, or Preinjection Skin Preparations¹

This standard is issued under the fixed designation E1173; 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 The practice is designed to measure the reduction of the microflora of the skin.

1.2 A knowledge of microbiological techniques is required for these procedures.

1.3 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.3.1 *Exception*—In this practice, metric units are used for all applications except for linear measure, in which case inches are used, and metric units follow in parentheses.

1.4 Performance of this procedure requires a knowledge of regulations pertaining to the protection of human subjects (1).²

NOTE 1—Importantly, it must be noted that the FDA currently does not accept data resulting from procedures in this Method for testing products for approval as Vascular Precatheterization Skin Preparations.

1.5 *This standard does not purport to address all of the safety problems, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety, health, and environmental practices and determine the applicability of regulatory limitations prior to use.*

1.6 *This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.*

2. Referenced Documents

2.1 ASTM Standards:³

¹ This practice is under the jurisdiction of ASTM Committee E35 on Pesticides, Antimicrobials, and Alternative Control Agents and is the direct responsibility of Subcommittee E35.15 on Antimicrobial Agents.

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² The boldface numbers in parentheses refer to the list of references at the end of this standard.

³ 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.

E1054 Practices for Evaluation of Inactivators of Antimicrobial Agents

E1874 Practice for Recovery of Microorganisms From Skin using the Cup Scrub Technique

E2756 Terminology Relating to Antimicrobial and Antiviral Agents

3. Terminology

3.1 Terms used in this standard are defined in E2756, Standard Terminology Relating to Antimicrobial and Antiviral Agents. Others defined below are specific to their use in this document.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *active ingredient, n*—a substance added to a formulation specifically for the inhibition or inactivation of microorganisms.

3.2.2 *inguen, n*—groin: the junctional region between the abdomen and thigh; pl. *inguina*.

3.2.3 *inguinal crease, n*—the discrete region of flexure between the abdomen and the thigh.

3.2.4 *sampling fluid, n*—a recovery fluid that contains a neutralizer demonstrated to inactivate or quench the active ingredient(s) in the test formulation and the positive and negative control formulations. See Practices E1054.

3.2.5 *test formulation, n*—a formulation containing an active ingredient(s).

4. Summary of Test Method

4.1 This practice is conducted on human subjects selected randomly from a group of volunteers who, after refraining voluntarily from using topical and oral antimicrobials for at least two weeks (14 days), exhibit acceptably high normal flora counts on the skin sites to be used in testing (see Section 8).

4.2 The antimicrobial activity of preoperative, vascular precatheterization, or preinjection skin preparations and of controls is measured by comparing microbial counts, obtained at various time intervals after application of a formulation to skin sites to counts obtained from those same sites prior to application of a formulation. Skin sites recommended for use in testing are: (1) the inguinal region and the abdomen for preoperative skin preparations; (2) the inguinal region, the

subclavian (clavicular) region, and/or the median cubital region of the arm for vascular precatheterization preparations; and (3) the median cubital region of the arm for preinjection skin preparations.

4.2.1 *Preoperative Skin Preparation*—Microbial samples are collected from the test sites a minimum of three (3) times after treatment application on both moist and dry skin sites. The recommended sample times are 30 s, 10 min, and 6 h post-treatment, but other relevant times may be selected.

4.2.2 *Vascular Precatheterization Preparation*—Microbial samples are collected from the test sites a minimum of three (3) times after treatment application on skin sites. The recommended sample times are “immediate,” 12 h, and 24 h post-treatment, but other relevant times may be selected.

4.2.3 *Preinjection Preparation*—Microbial samples are collected from the skin sites 30 s post-treatment.

4.3 The fluid used for sampling the test sites must effectively quench (neutralize) the antimicrobial action of all formulations tested. The effectiveness of the inactivator must be demonstrated prior to initiation of product-testing, as described in Practices E1054, and using in-vivo techniques consistent with the cup-scrub technique (see Section 10).

4.4 A positive control formulation having performance characteristics known to the laboratory (internal validity of test) and a negative control (test formulation without active ingredient or a physiological saline solution) must be tested in parallel with the test formulation.

5. Significance and Use

5.1 These procedures should be used to test topical antimicrobial-containing preparations that are intended to be fast-acting in reducing significantly the number of microorganisms on intact skin immediately and, for preoperative and vascular precatheterization preparations, maintenance of some degree of reduction for an extended time.

6. Apparatus

6.1 *Colony Counter*—Any of several types may be used; for example, Quebec colony counters and similar devices, or automated, computerized plater/counter systems.

6.2 *Incubator*—Any incubator that can maintain a temperature of $30^{\circ} \pm 2^{\circ}\text{C}$ may be used.

6.3 *Sterilizer*—Any steam sterilizer that can produce the conditions of sterilization is acceptable.

6.4 *Timer (stopwatch)*—One that displays hours, minutes, and seconds.

6.5 *Examining Table*—Any elevated surface, such as a 3-by-6-ft (0.9-by-1.8-meter) table with mattress or similar padding to allow the subject to recline comfortably.

7. Reagents and Materials

7.1 *Bacteriological Pipettes*—10.0 and 2.2-mL or 1.1-mL capacity, available from most laboratory supply houses.

7.2 *Petri Dishes*—100 mm by 15 mm for performing standard plate counts, available from most laboratory supply houses.

7.3 *Scrubbing Cups*—Autoclavable cylinders, height approximately 1 in (2.5 cm) and inside diameter of a size convenient to placement on the skin of the anatomical area to be sampled. Useful diameters range from approximately 0.5 to 1.5 in (1.3 to 3.8 cm), depending on sites to be sampled.

7.4 *Rubber Policeman, TFE-fluorocarbon Scrubbers, or other appropriate device*—Can be fashioned in the laboratory or purchased from most laboratory supply houses. Whichever type is selected, it should be used throughout the course of testing.

7.5 *Testing Formulation, including directions for use.*

7.6 *Sterile Gauge Pads*—Used to cover treated skin sites.

7.7 *Sterile Dressings*⁴—Used to cover treated skin sites.

7.8 *Sampling Fluid*—Dissolve 0.4 g KH_2PO_4 , 10.1 g Na_2HPO_4 , and 1.0 g isooctylphenoxypolyethoxyethanol in 1 L of distilled water. Inactivator(s) specific for the antimicrobial active(s) in the test and control formulations must be included (See Practices E1054). Adjust to pH 7.8. Dispense in appropriate volumes and sterilize.

7.9 *Dilution Fluid*—Butterfield’s (2) phosphate-buffered water adjusted to pH 7.2, or other suitable diluent, which must contain antimicrobial inactivators specific for the test, positive and negative control formulations (see Practices E1054).

7.10 *Plating Medium*—Soybean-casein digest agar (3), with or without antimicrobial inactivators.

7.11 *Sterile Template Material*—Used to demarcate the skin sites; made from paper, plastic, or cloth, for example.

7.12 *Surgical Skin Marker*—Used to delineate mark the skin sites to be used in testing.

NOTE 2—Because some markers contain crystal violet or other fluids that are inhibitory to many skin microflora, a marker should be proven non-antimicrobial prior to use in testing.

8. Skin Sites to be Used in Testing

8.1 Preoperative Skin Preparations:

8.1.1 The skin sites selected for evaluation of the effectiveness of preoperative skin preparations should include both moist and dry skin areas. Bacterial baseline populations should be at least $3.0 \log_{10}/\text{cm}^2$ greater on moist skin sites than the detection limit of the sampling procedure, and at least $2.0 \log_{10}/\text{cm}^2$ greater than the detection limit on dry skin sites. The preferred moist-skin areas are the inguina, in which skin-to-skin contact results in a moist environment conducive to higher populations of microflora. The preferred dry-skin area is the lower abdomen adjacent to the umbilicus. These areas are illustrated in Fig. 1.

8.1.2 Using a 1.5-by-5-in (3.8-by-12.7-cm) sterile template (for example, paper, plastic, cloth), treatment sites in the

⁴ The sole source of supply of the apparatus (TELEA non-adherent dressing, No. 3279) known to the committee at this time is Kendall Co.; Hospital Products; Boston, MA 02101. This product is not sterile, but can be steam-sterilized prior to use. If you are aware of alternative suppliers of appropriate dressings, please provide this information to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee,¹ which you may attend.