



Designation: E1053 – 20

# Standard Practice to Assess Virucidal Activity of Chemicals Intended for Disinfection of Inanimate, Nonporous Environmental Surfaces<sup>1</sup>

This standard is issued under the fixed designation E1053; 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 ( $\epsilon$ ) indicates an editorial change since the last revision or reapproval.

## 1. Scope

1.1 This practice is used to evaluate the virucidal efficacy of liquid, aerosol, or trigger-spray microbicides intended for use on inanimate, nonporous environmental surfaces. This practice may be employed with most viruses, which can be grown in cultured cells.<sup>2</sup> However, other host systems (for example, embryonic eggs) may be used with proper justification and documentation.

1.2 This practice should be performed only by those trained in microbiological and virological techniques in facilities designed and equipped for work with infectious agents at the appropriate biosafety level.

1.3 It is the responsibility of the investigator to determine whether Good Laboratory Practice regulations (GLPs) are required and to follow them where appropriate (40 CFR, Part 160 for EPA submissions and 21 CFR, Part 58 for FDA submissions). Refer to the appropriate regulatory agency for performance standards of virucidal efficacy.

1.4 *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, health, and environmental practices and determine the applicability of regulatory limitations prior to use.* The user should consult a reference for laboratory safety recommendations.<sup>2</sup>

1.5 *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.*

<sup>1</sup> 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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<sup>2</sup> Public Health Service, Centers for Disease Control and Prevention, and National Institutes of Health, *Biosafety in Microbiological and Biomedical Laboratories*, U.S. Department of Health and Human Services, Washington, DC, December 2009, 422 pp.

## 2. Referenced Documents

2.1 *ASTM Standards*:<sup>3</sup>

D1129 Terminology Relating to Water

E1153 Test Method for Efficacy of Sanitizers Recommended for Inanimate, Hard, Nonporous Non-Food Contact Surfaces

E1482 Practice for Use of Gel Filtration Columns for Cytotoxicity Reduction and Neutralization

E2197 Quantitative Disk Carrier Test Method for Determining Bactericidal, Virucidal, Fungicidal, Mycobactericidal, and Sporocidal Activities of Chemicals

2.2 *Federal Standards*:<sup>4</sup>

Title 21, Code of Federal Regulations (CFR), Food and Drug Administration, Part 58, Laboratory Practice for Nonclinical Laboratory Studies

Title 40, Code of Federal Regulations (CFR), Environmental Protection Agency, Subchapter E, Pesticide Programs; Part 160, Good Laboratory Practice Standards

## 3. Terminology

3.1 *Definitions*—For definitions of general terms used in this test method, refer to Terminology D1129.

3.2 *Definitions of Terms Specific to This Standard*:

3.2.1 *hard water, n*—water with a standard hardness as calcium carbonate.

3.2.2 *neutralization, n*—a process which results in quenching the microbicidal activity of a test substance. This may be achieved through dilution of the test substance to reduce the microbicidal activity, or through the use of chemical agents, called neutralizers, to eliminate microbicidal activity.

3.2.3 *soil load, n*—a solution of one or more organic and/or inorganic substances added to the suspension of the test

<sup>3</sup> 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.

<sup>4</sup> Available from U.S. Government Printing Office Superintendent of Documents, 732 N. Capitol St., NW, Mail Stop: SDE, Washington, DC 20401, http://www.access.gpo.gov.

organism to simulate the presence of body secretions, excretions or other extraneous substances.

3.2.4 *test substance or test formulation, n*—a formulation which incorporates microbicidal ingredients.

#### 4. Summary of Practice

4.1 The virus suspension is dried on an inanimate, nonporous surface. The test substance is added over the dried film at its use-dilution or sprayed from an aerosol can or trigger-sprayer following the manufacturer's directions. Control carriers receive an equivalent volume of a buffer harmless to the test virus and its host cells. After exposure at the appropriate temperature (usually  $20 \pm 2^\circ\text{C}$ ) for the recommended time, the eluates from control and test carriers are assayed for infectivity.

4.2 This practice is designed to be performed by a person trained in culturing and assaying infectious viruses who is responsible for choosing the appropriate host system for the test virus and applying the techniques necessary for propagation and maintenance of the host system and test virus. For a reference text, refer to Lennette et al.<sup>5</sup>

#### 5. Significance and Use

5.1 This practice may be used to determine the effectiveness of liquid, aerosols/foams, and trigger-spray products against designated prototype viruses.

5.2 The number of lots of the test substance and the number of replicates in each test will depend on the requirements of the target regulatory agency.

5.3 Certain regulatory agencies may require additional testing using other carrier tests for product registration purposes.

#### 6. Materials and Reagents

6.1 *Host System and Assay of Infectious Virus*—See **Note 2**.

6.1.1 *Cell Cultures in multi-well plates*, appropriate for test virus.

6.1.2 *Growth and Maintenance Media*, any growth and maintenance media suitable for work with the virus and its host cells.

NOTE 1—Materials and reagents for cell culture may be purchased from biological supply houses.

6.1.3 *Diluent*, Earle's Balanced Salt Solution (EBSS) or other appropriate media.

6.1.4 *Plastic Cell Culture Ware*.

NOTE 2—Plastic cell culture ware may be purchased from most laboratory supply houses.

6.1.5 *Incubator*, capable of maintaining  $36 \pm 1^\circ\text{C}$  or other temperature appropriate for replication of the specific test virus; an incubator with 5 to 7 %  $\text{CO}_2$  will be needed if an open system is being used for cell culture and virus assay.

6.1.6 *Refrigerator*,  $4 \pm 2^\circ\text{C}$ .

6.1.7 *Test Tubes*, screw-capped or snap capped or similar.

6.1.8 *Pipettes*, serological, various volumes.

6.1.9 *Micropipettors and Sterile Tips*

6.1.10 *96-Well Dilution Plates*, (only required for use in a 96-well assay).

6.1.11 *Microtitration Kit*, if applicable.

NOTE 3—Microtitration kits may be purchased from most laboratory supply houses.

6.1.12 *Petri Plates*, glass, 100-mm diameter, 10-15 mm deep.

6.1.13 *Microscope*.

6.1.14 *Biosafety cabinet*.

#### 7. Test Viruses

7.1 **Appendix X1** lists viruses and their respective host cells as examples for use in this practice. Other viruses and cell lines may be used.

7.2 To demonstrate that the test substance has broad virucidal activity, it should be shown to be effective against at least one non-enveloped virus.

#### 8. Virus Stock

8.1 The titer of the test virus suspension must be sufficiently high so that at least  $10^{4.8}$  infective units/carrier can be recovered from the inoculated carriers after the inoculum has dried. The host system employed for virus propagation need not be the same as that used for virus recovery and the infectivity assay.

#### 9. Operating Technique

9.1 *Test Substance Diluent*—For test substances requiring dilution in water to obtain a use-dilution, water with a standardized and specified level of hardness, or otherwise recommended by manufacturer, shall be used as the diluent.<sup>6</sup>

9.2 *Cytotoxicity Control*—The objective of this control is to (1) determine the dilution of the test substance post-neutralization at which it causes no apparent degeneration (cytotoxicity) of the cell line to be used for measuring virus infectivity and (2) assess if the neutralizer in any way reduces or enhances such cytotoxicity. Make an initial 1:2 dilution of the use-dilution of the test substance in the neutralizer and three further ten-fold dilutions of the neutralized test substance in the diluent. Remove the culture medium from the monolayers of the host cell line(s) and put into each test monolayer separately the same volume of inoculum as used in virus titration; control monolayers receive an equivalent amount of diluent (without any neutralizer) only. Another set of monolayers should be exposed to the neutralizer alone. Hold the cultures for the same period of time as used in virus titration and examine them under an inverted microscope for any cytotoxicity. In case of cytotoxicity, a different neutralizer or gel filtration (see Test Method **E1482**) of the neutralized virus-test substance mixture may be needed.

NOTE 4—If gel filtration is used in the virucidal activity test runs, the

<sup>5</sup> Schmidt, N. J., Lennette, D. A., Lennette, E. T., and Lennette, E. H. eds., *Diagnostic Procedures for Viral, Rickettsial and Chlamydial Infections*, Seventh Edition, American Public Health Association, Washington, DC, 1995.

<sup>6</sup> AOAC International, *Official Methods of Analysis of the AOAC*, Arlington, VA, 1990.

neutralized and gel-filtered test substance should be evaluated for the cytotoxicity control.

**9.3 Test Substance Neutralization Control**—To determine the dilution at which neutralization of the test substance has occurred, prepare and inoculate an additional set of neutralization/cytotoxicity controls following the same procedure as the test substance efficacy evaluation runs using a mock inoculum (for example, dilution medium) in lieu of virus. To validate the neutralization, add equal volumes of the neutralized test substance, and a control fluid (for example, PBS) a relatively low number (for example, 1000 to 5000) infective units of the test virus and hold the mixtures for at least the contact time at the contact temperature. Titrate the mixtures for infectious virus. Comparable levels of infective units must be recovered from the control and the neutralized test substance for the neutralization to be successful. In case of incomplete neutralization, either another neutralizer may be needed or the gel filtration method (Test Method **E1482**) may be used. If gel filtration columns are used as the only neutralization method, the test substance will be added to a previously prepared gel filtration column. The solution is pushed through the column utilizing the syringe plunger or by centrifugation. Then a relatively low number (for example, 1000 to 5000) infective units of the test virus will be added to the eluate; and the mixtures held for at least the contact time at room temperature. Titrate the mixtures for infectious virus. Comparable levels of infective units must be recovered from the control and the post-column eluate for the neutralization to be successful.

**9.4 Plate Recovery Control**—Vortex the virus suspension thoroughly and place 0.2-mL on the inside bottom surface of each glass petri dish. Allow the virus inoculum to dry under ambient conditions in a laminar flow hood or other suitable chamber with the petri dish cover removed. The drying time of this control should be consistent with that for the test runs. A recovery of at least  $10^{4.8-6.3}$  infective units/control dish should be achieved for the test to be considered valid. Pools of certain types of viruses may require concentration by ultra-centrifugation to obtain titers high enough to give a minimum of  $10^{4.8}$  infective units/dish after the inoculum has been dried. However, any such concentrated virus must be vortexed well to reduce the presence of viral aggregates. If the plate recovery control has a virus endpoint titer of  $>10^{6.3}$  infective units/control dish the virus culture may be standardized by dilution to target a virus endpoint titer of  $10^{4.8}$  to  $10^{6.3}$  infective units/control dish or 3-5  $\text{Log}_{10}$  beyond the level of cytotoxicity. The endpoint titer range may be modified by individual protocols.

**NOTE 5**—The volume of virus inoculum per carrier may be increased depending on the titer of the virus. This volume must be consistent between the plate recovery control and test substance runs. It should be noted, however, that an increased volume will prolong the drying of the inoculum and may lead to increased losses in virus infectivity.

**NOTE 6**—Results of Cytotoxicity should be taken into account when determining if the virus culture should be standardized by dilution to target a virus endpoint titer of  $10^{4.8}$  to  $10^{6.3}$  infective units/control dish.

**9.4.1** After drying, overlay each dried control film with 2.0 mL of PBS or another buffered solution harmless to the virus and its host cells. Let stand for the same contact time as used for the test carriers and then add an equal amount of neutralizer

(2.0 mL). Scrape the inside bottom surface with a sterile cell scraper to resuspend the viral film. If gel filtration columns will be used in addition to the liquid neutralizer, following scraping the suspension will be added to a previously prepared gel filtration column. The solution is pushed through the gel filtration column utilizing the syringe plunger or by centrifugation. This suspension may be considered the 10–1 dilution of the virus. Prepare serial 10-fold dilutions using diluent and inoculate an amount appropriate to the test format to no less than four replicate cell monolayers/dilution. Alternatively, in lieu of adding liquid neutralizer following the contact time, just prior to the end of the exposure time the dish is scraped and immediately following the contact time the solution is added to a gel filtration column which was prepared previously. The solution is pushed through the gel filtration column utilizing the syringe plunger or by centrifugation. This suspension may be considered the  $10^{-1}$  dilution of the virus. See Practice **E1482**.

**9.5 Test for Virucidal Activity**—For each lot of the test substance, treat a dried film carrier with 2.0 mL of the use-dilution of a liquid product or the amount of product released during recommended use of the aerosol or trigger spray. Hold for the required contact time. Upon completion of the contact time, immediately add an equal volume of neutralizer (2.0 mL) to the carrier and mix well. Scrape the film to resuspend the virus/test substance/neutralizer mixture. If gel filtration columns will be used in addition to the liquid neutralizer, following scraping the suspension will be added to a previously prepared gel filtration column. The solution is pushed through the gel filtration column utilizing the syringe plunger or by centrifugation. If gel filtration columns are used as the only neutralization method, the test substance will be added to a previously prepared gel filtration column. The solution is pushed through the column utilizing the syringe plunger or by centrifugation. Prepare serial 10-fold dilutions using diluent and inoculate an amount appropriate to the test format to no less than four replicate cell monolayers/dilution starting from the first ten-fold dilution of the post-neutralized sample.

## **10. Soil Load (refer to Test Method **E2197**)**

**10.1** Fetal Bovine Serum, Horse Serum, or other animal serum as desired, may be used depending on the target regulatory agency. The soil load mixture given in **10.2 – 10.7** contains a level of protein roughly equal to that in 5 % serum. Preliminary screening of albumin and mucin is recommended to ensure compatibility with test organism(s).

**10.2** The soil load to be incorporated in the suspension of the test organism may consist of a mixture of the following stock solutions in phosphate buffer (pH 7.2):

**10.3** Add 0.5 g of tryptone or yeast extract to 10 mL of phosphate buffer.

**10.4** Add 0.5 g of BSA to 10 mL of phosphate buffer.

**10.5** Add 0.04 g of bovine mucin to 10 mL of phosphate buffer.

**10.6** Prepare the solutions separately and sterilize by passage through a 0.22- $\mu\text{m}$  pore diameter membrane filter, aliquot,