



Designation: ~~E1259—18~~ E1259 – 23

## Standard Practice for Evaluation of Antimicrobials in Liquid Fuels Boiling Below 390 °C<sup>1</sup>

This standard is issued under the fixed designation E1259; 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 reappraisal. A superscript epsilon ( $\epsilon$ ) indicates an editorial change since the last revision or reappraisal.

### 1. Scope\*

1.1 This practice is designed to evaluate antimicrobial agents for the prevention of microbially influenced deterioration of liquid fuels (as defined by Specification [D396](#), [D910](#), [D975](#), [D1655](#), [D2069](#), [D2880](#), [D3699](#), [D4814](#), [D6227](#), [D6751](#), and [D7467](#)), system deterioration, or both.

1.2 Knowledge of microbiological techniques is required for these procedures.

1.3 It is the responsibility of the investigator to determine whether Good Laboratory Practice (GLP) is required and to follow them where appropriate (40 CFR, 160), or as revised.

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

1.5 *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.*

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:<sup>2</sup>

- [D396 Specification for Fuel Oils](#)
- [D910 Specification for Leaded Aviation Gasolines](#)
- [D975 Specification for Diesel Fuel](#)
- [D1655 Specification for Aviation Turbine Fuels](#)
- [D2069 Specification for Marine Fuels \(Withdrawn 2003\)<sup>3</sup>](#)
- [D2880 Specification for Gas Turbine Fuel Oils](#)
- [D3699 Specification for Kerosine](#)
- [D4814 Specification for Automotive Spark-Ignition Engine Fuel](#)

<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.

Current edition approved Oct. 1, 2018; June 1, 2023. Published October 2018; June 2023. Originally approved in 1988. Last previous edition approved in 2016 as ~~E1259—16~~ E1259 – 18. DOI: [10.1520/E1259-18](https://doi.org/10.1520/E1259-18); [10.1520/E1259-23](https://doi.org/10.1520/E1259-23).

<sup>2</sup> For referenced ASTM standards, visit the ASTM website, [www.astm.org](http://www.astm.org), or contact ASTM Customer Service at [service@astm.org](mailto:service@astm.org). For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

<sup>3</sup> The last approved version of this historical standard is referenced on [www.astm.org](http://www.astm.org).

\*A Summary of Changes section appears at the end of this standard

- D5465 Practices for Determining Microbial Colony Counts from Waters Analyzed by Plating Methods
- D6227 Specification for Unleaded Aviation Gasoline Containing a Non-hydrocarbon Component
- D6293 Test Method for Oxygenates and Paraffin, Olefin, Naphthene, Aromatic(O-PONA) Hydrocarbon Types in Low-Olefin Spark Ignition Engine Fuels by Gas Chromatography (Withdrawn 2009)<sup>3</sup>
- D6469 Guide for Microbial Contamination in Fuels and Fuel Systems
- D6729 Test Method for Determination of Individual Components in Spark Ignition Engine Fuels by 100 Metre Capillary High Resolution Gas Chromatography
- D6733 Test Method for Determination of Individual Components in Spark Ignition Engine Fuels by 50-Metre Capillary High Resolution Gas Chromatography
- D6751 Specification for Biodiesel Fuel Blendstock (B100) for Middle Distillate Fuels
- D6974 Practice for Enumeration of Viable Bacteria and Fungi in Liquid Fuels—Filtration and Culture Procedures
- D7463 Test Method for Adenosine Triphosphate (ATP) Content of Microorganisms in Fuel, Fuel/Water Mixtures, and Fuel Associated Water
- D7464 Practice for Manual Sampling of Liquid Fuels, Associated Materials and Fuel System Components for Microbiological Testing
- D7467 Specification for Diesel Fuel Oil, Biodiesel Blend (B6 to B20)
- D7687 Test Method for Measurement of Cellular Adenosine Triphosphate in Fuel and Fuel-associated Water With Sample Concentration by Filtration
- D7978 Test Method for Determination of the Viable Aerobic Microbial Content of Fuels and Associated Water—Thixotropic Gel Culture Method
- D8412 Guide for Quantification of Microbial Contamination in Liquid Fuels and Fuel-Associated Water by Quantitative Polymerase Chain Reaction (qPCR)
- E1054 Practices for Evaluation of Inactivators of Antimicrobial Agents
- E1259 Practice for Evaluation of Antimicrobials in Liquid Fuels Boiling Below 390 °C
- E1326 Guide for Evaluating Non-culture Microbiological Tests

## 2.2 NACE Standard:

TM0172 Determining Corrosive Properties of Cargoes in Petroleum Product Pipelines<sup>4</sup>

## 2.3 Federal Standards:

40 CFR Part 79 Fuels and Fuel Additives Registration Regulations<sup>5</sup>

40 CFR Part 152 Pesticide Registration and Classification Procedures<sup>5</sup>

## 3. Terminology

### 3.1 Definitions of Terms Specific to This Standard:

3.1.1 *antimicrobial, n*—see *biocide*.

3.1.2 *biocide, n*—a physical or chemical agent that kills living organisms.

#### 3.1.2.1 Discussion—

Biocides are further classified as bactericides (kill bacteria), fungicides (kill fungi), and microbicides (kill both bacterial and fungi). They are also referred to as *antimicrobials*.

3.1.3 *microbially-influenced deterioration, n*—decomposition /degradation of material (fuel) or making unsuitable for use, as a result of metabolic activity or the presence of microbes.

3.1.4 *microbicide, n*—see *biocide*.

3.1.5 *microcosm, n*—a miniature system used to model larger systems.

#### 3.1.5.1 Discussion—

It is generally impractical to evaluate microbicide performance in large fuel storage system capacities (> 24 000 m<sup>3</sup>), consequently small volume (1.0 to 208 L capacity) microcosms are used as model systems.

<sup>4</sup> Item No. 21204, available from NACE International (NACE), 1440 South Creek Dr., Houston, TX 77084-4906, <http://www.nace.org>.

<sup>5</sup> Available from U.S. Government Printing Office Superintendent of Documents, 732 N. Capitol St., NW, Mail Stop: SDE, Washington, DC 20401.

#### 4. Summary of Practice

4.1 This practice is conducted on a fuel representative of the grade to be treated, and determines the antimicrobial efficacy under well-defined conditions that may include specific inocula or an uncharacterized inoculum from a microbially contaminated fuel system.

4.1.1 Water/fuel ratios and containment time are also defined. This practice allows for impact of fuel/water partitioning and time, on the antimicrobial agent, as well as the effect of continual rechallenge.

4.1.2 At each sampling time interval, treated and untreated aliquots are checked for the treated population ~~survival~~ survival, proliferation, or both. Microbiological testing is coupled with gross observations of each system for biofilm formation and interfacial growth.

4.1.3 The size of the test system, total volume of fluid, fuel to bottom-water ~~ratio and test duration~~ ratio, test duration, and test system incubation temperature may vary depending on the specific objectives of the test.

4.1.4 Before beginning any test plan intended to meet performance testing compliance requirements, confirm that the cognizant authority accepts the test protocol.

#### 5. Significance and Use

5.1 Guide **D6469** details the types of problems associated with uncontrolled microbial growth in fuels and fuel systems. Treatment with effective antimicrobial agents is one element of contamination control strategy.

5.2 The procedure should be used to evaluate the relative efficacy of microbicides in liquid fuels boiling below 390 °C. The effect of environmental conditions, such as a variety of fuel additives, metal surfaces, and climatology, are variables that can be included in specific tests using this protocol.

5.3 This practice addresses product performance issues only. Regulatory Agencies restrict and control the use of both pesticides (in the U.S.: 40 CFR 152) and fuel additives (40 CFR 79). Regardless of performance in this method, antimicrobials must only be used in compliance with applicable regulations. Specific industries, for example, the aviation industry, may place further restrictions on chemicals used for fuel treatment.

#### 6. Apparatus [standards.iteh.ai/catalog/standards/sist/0054e30d-9b3e-4b0f-9d05-b35cd15d495/astm-e1259-23](https://standards.iteh.ai/catalog/standards/sist/0054e30d-9b3e-4b0f-9d05-b35cd15d495/astm-e1259-23)

6.1 *Colony Counter*—Any of several types, for example, a Quebec Colony Counter may be used.

6.2 *Drums; Steel*—208 L (55 gal) 16 ga. steel, open-head drum with removable 16 ga. lid fitted with 2.05 cm and ~~1.90 cm~~ 1.90 cm threaded ports for venting and sampling.

6.3 *Incubator*—Any incubator capable of maintaining temperature of ~~30~~ 30 °C to 35 °C may be used.

6.4 *Glass Jars*—French square or similar configuration.

NOTE 1—Jar capacity should be determined based on the test plan designed fuel to water ratio and the expected sample volume size needed for weekly testing (9.5 and 9.9).

6.5 *Pails; Steel*—18.9 L (5 gal) steel, open-head pail with removable 16 ga. lid fitted with 2.05 cm and 1.90 cm threaded ports for venting and sampling.

6.6 *Sterilizer*—Any suitable steam sterilizer capable of producing the conditions of sterility is acceptable. A pressurized filter sterilization apparatus of appropriate capacity to filter sterilize the test fuels and bottom-water used in the negative control microcosms. A 0.2 µm pore-size methyl cellulose or cellulose acetate membrane should be used as the filtration medium.

6.7 *Vortex*—Mixer.

## 7. Reagents and Materials<sup>6</sup>

7.1 *Petri Dishes*—~~100 by 15 mm~~ 100 mm by 15 mm required for performing standard plate count.

7.2 *Bacteriological Pipets*—10.0 mL and ~~1.1, or 2.2 mL~~ 1.1 mL, or 2.2 mL capacity.

7.3 *Water Dilution Bottles*—Any sterilizable glass container having a ~~150 to 200 mL~~ 150 mL to 200 mL capacity and tight closure may be used.

7.4 *Fuel*.

NOTE 2—Representative fuel samples from each product grade are available from all petroleum refiners.

7.5 *Synthetic Bottom Water*.

NOTE 3—In order to promote microbial growth of the inoculum when using the fuel as the sole source of organic nutrients, synthetic bottom water may contain various inorganic nutrients. An example, of a commonly used synthetic bottom water is Bushnell-Haas Mineral Salts medium (BHMSS),<sup>7</sup> with the concentration adjusted to simulate a particular type of bottoms-water (marine, brackish, fresh, etc.).

7.6 *Soy Peptone Casein Digest Agar*.

7.7 *Sabouraud Dextrose Agar*.

7.8 *Agar, Bacteriological Grade*.

7.9 *Potassium Tellurite Solution*—sterile 1 %.

7.10 *Gentamicin Sulfate*—50 µg/mL.

7.11 *Plate Count Agar*.

7.12 *Potato Dextrose Agar*.

NOTE 4—Items 7.5 – 7.12 are available from a variety of media manufacturers and chemical supply companies.

## 8. Inoculum

8.1 *Inoculum Selection*:

8.1.1 Depending on the objectives of a test plan, one or more characterized cultures (for example: bacterium, yeast and mold) can be selected or microbially contaminated bottoms-water collected from a fuel system can be used.

8.1.2 Contaminated fuel system microbial communities can be quite diverse and contain >50 different taxa. Consequently, when Practice E1259 is to be used in order to assess a product's general antimicrobial performance properties in fuel systems, multi-taxa inocula provide a more realistic challenge population than either single or commonly used, three taxa inocula.

8.1.3 The use of standardized cultures to prepare microcosm inocula facilitates corroborative testing.

8.1.4 Inoculum taxa should be selected from cultures known to grow using fuel as their sole carbon source.

8.1.5 Depending on microcosm design, it can be appropriate to include aerobic and anaerobic taxa. If inhibition of

<sup>6</sup> ACS Reagent Chemicals, Specifications and Procedures for Reagents and Standard-Grade Reference Materials, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see *Analar Standards for Laboratory Chemicals*, BDH Ltd., Poole, Dorset, U.K., and the *United States Pharmacopeia and National Formulary*, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.

<sup>7</sup> Bushnell, L.D. and Haas, H.F. 1941. The utilization of certain hydrocarbons by microorganisms. *J. Bacteriol.* 41: 653- 673.

microbiologically influence corrosion is to be assessed, the challenge population should include iron related bacteria, acid producing bacteria and sulfate reducing bacteria as part of the inoculum mixture.

8.1.6 Uncharacterized, bottoms-water, contaminant populations are most appropriate when Practice E1259 is to be used to evaluate microbicide performance efficacy in a single system or family of systems (for example, bulk storage tanks for a specific fuel grade at a specific facility).

## 8.2 Inoculum Preparation and Maintenance:

8.2.1 *Inoculum Revitalization*—Commonly used cultures are *Pseudomonas aeruginosa*, ATCC No. 33988, *Hormoconis resinae*, ATCC No. 20495, and *Candida viswanathii* (formerly *Candida tropicalis* and *Yarrowia tropicalis*), ATCC No. 48138. However, in accordance with 8.1, additional cultures can be used.

8.2.1.1 Obtain cultures from ATCC. Before initiating fuel antimicrobial tests, revitalize each of the three cultures in accordance with the instructions contained with each culture.

8.2.2 *Maintenance and Preparation of Pre-Inocula*—All cultures are transferred from slants of a specified agar, (for example, *a*) *Pseudomonas aeruginosa* (Plate Count Agar), (*b*) *Hormoconis resinae* Potato Dextrose Agar, and (*c*) *Yarrowia tropicali* (Potato Dextrose Agar)) to synthetic bottom water medium in a suitable size screw-cap glass bottle (6.4).

8.2.2.1 Overlay inoculated bottom water with fuel to give a final fuel to water ratio of 10.

8.2.2.2 Keep this two-phase system at room temperature ( $\pm 20(20\text{ }^{\circ}\text{C}$  to  $30\text{ }^{\circ}\text{C}$ ) for seven days.

8.2.2.3 Weekly, transfer the interface, along with half the bottom water to a similar system until the inoculum used.

8.2.2.4 During this inoculum preparation period the bacterial levels should be maintained at approximately  $10^7$  CFU/mL or non-culture test bioburden equivalent, the yeast levels at approximately  $10^6$  CFU/mL, and mold levels at approximately  $10^4$  spores/mL.

8.2.2.5 Freshly collected, microbially contaminated bottoms-water can be maintained per 8.2.2.1 – 8.2.2.4.

## 8.2.3 Preparation of Challenge (Test) Inoculum:

8.2.3.1 To prepare the test inoculum, dilute bacterial pre-inocula 1:100 to achieve a population equivalent to approximately  $10^5$  CFU/mL. Dilute yeast and molds 1:10 to achieve a population equivalent to approximately  $10^3$  CFU/mL.

8.2.3.2 At time zero, just prior to adding inoculum to each setup, and at each subsequent time point, determine the microbial population density (9.9).

8.2.3.3 If test systems larger than 1.0 L will be used, the challenge inoculum should first be acclimated to growth in systems that contain the same volume and fuel to bottom-water ratio as the test systems.

## 9. Procedure

9.1 *Test Array Determination*—The test plan determines the number and capacities of microcosms needed for the test plan. Preferably, duplicate microcosms will be set up for each control and test treatment.

9.1.1 Controls may include any combination of:

9.1.1.1 Filter sterilized fuel over filter sterilized water.

9.1.1.2 Challenged, microbicide-free fuel over water.

NOTE 5—Some commercially available fuels contain additives with antimicrobial properties. It may be necessary to filter such fuels through activated carbon filters before using them for microbicide performance testing.

9.1.1.3 *Reference Control*—Microbicide treated fuel over bottom-water.

9.1.2 *Microbicide Treatment Dose*—Testing may be performed using a single dose or a range of doses. Typically the minimum and maximum doses permitted under the microbicide’s FIFRA registration are used. One or intermediate concentrations may also be used. For cost-effectiveness comparisons, dose selection may be based on the treatment costs of the microbicide against which the test product is being evaluated.

9.1.3 To determine the number of microcosms needed for the test array, add the total number of control and test treatments and multiply by the number of replicate microcosms required.

9.2 *Determine Microcosm Volume*—Microcosm volume will depend on test objectives.

9.2.1 Preliminary microbicidal product screening can be performed in 1 L or ~~2~~2 L microcosms.

9.2.2 Microbicide partitioning between fuel and water phases, in test microcosms and under field conditions, is likely to be affected by fuel to water ratios.

9.2.2.1 Use of a fuel to water ratio of 1000 to 1 is recommended, although fuel to water ratios between 50:1 and 500:1 may also be used, depending on factors such as sample ~~availability~~availability and test system volume.

NOTE 6—All fuel-grades covered by this practice have sufficiently high vapor pressures to permit off-gassing of noxious, potentially toxic volatile organic carbon (VOC) molecules. Small microcosms should be set up inside a fume hood. Microcosms too large to be stored inside a fume hood should be equipped with a vapor trapping system. A simple system can be designed from polyvinylchloride (PVC) piping and buckets filled with activated carbon (see Fig. 1).

9.3 *Determine Bottom-Water Composition*—Depending on the anticipated end-use application, bottom-water composition may

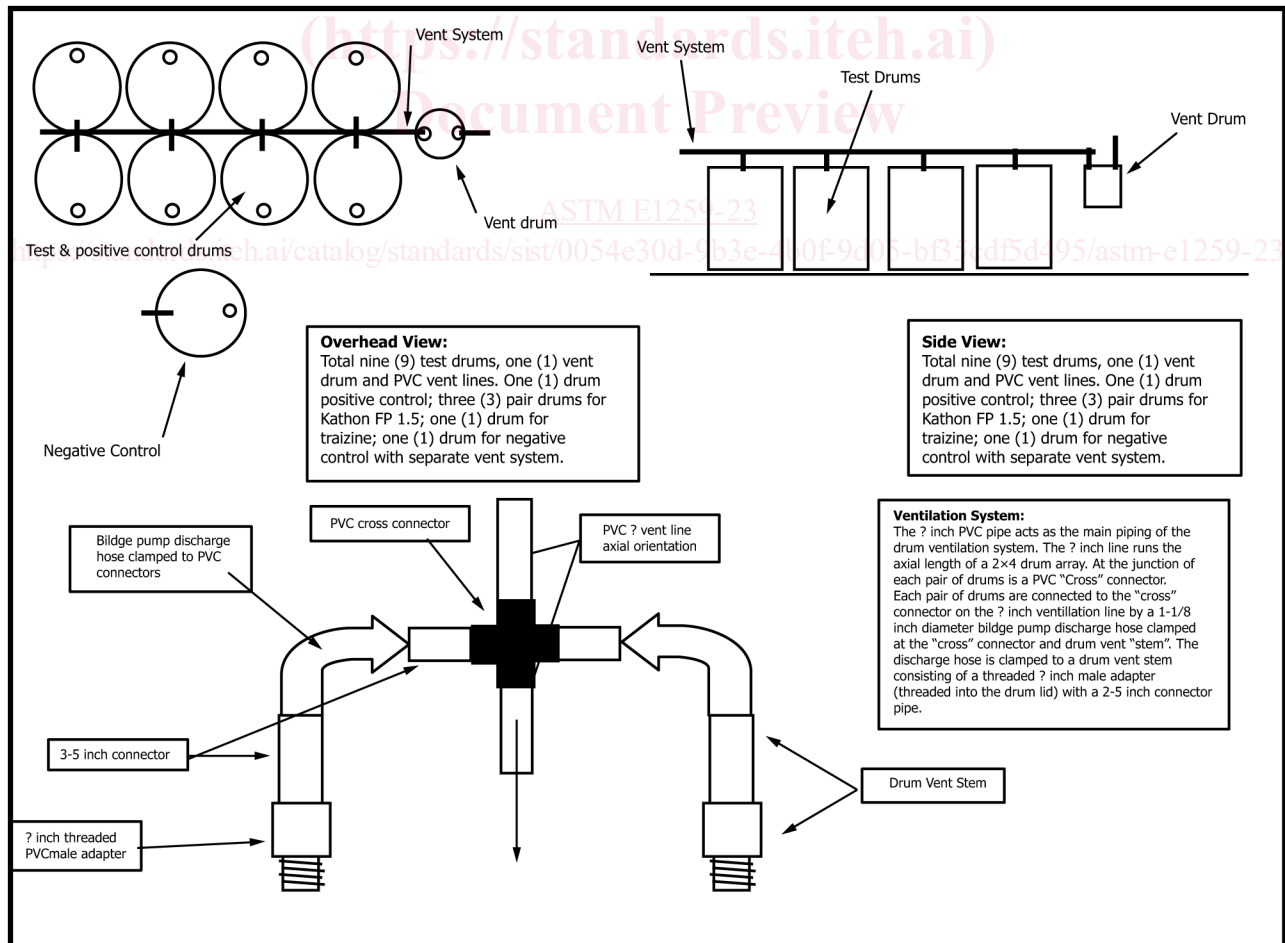


FIG. 1 Schematic Drawing for an Eight-Drum Microcosm Array Ventilation System

range from distilled water (simulating condensate-water accumulation) to sea-water. Recognizing that bottom-water chemistry varies substantially amongst fuel tanks, site-specific testing should be performed using filter-sterilized water from fuel tanks.

9.4 *Determine Challenge Frequency*—The test plan may include a single challenge or repeated challenges. Typically, when repeated challenges are used, they are scheduled for immediately after each sample collection time.

9.5 *Determine Sampling Schedule:*

9.5.1 *Kill-Rate Testing*—For speed of kill or kill-rate testing, collect samples at appropriate time points, such as after 30 min; 4, 8, 16, 24, 48, and 72 h.

9.5.2 *Persistence of Effect Testing*—Sample at periodically at daily, weekly, or monthly intervals until microcosm with highest microbicide dose fails (see 10.2.3).

NOTE 7—The choice of sampling intervals will depend on the test objectives. Although the protocol, as described, simulates long-term storage, it can be modified to simulate more dynamic fuel systems such as vehicle, vessel, or fueling facility (commercial, depot, fleet or retail) tanks. Dynamic systems can be simulated by exchanging  $\geq 70\%$  of the fuel volume at intervals similar to facility turnover rates (that is, daily, weekly, monthly).

NOTE 8—To simulate long-term storage, replace fuel and bottom-water volumes removed after sampling, but do not re-challenge. To simulate high turnover systems, replace fuel and bottom-water volumes and re-challenge after each sampling.

9.6 *Set Up Microcosms:*

9.6.1 If test will include corrosion testing (NACE TMO172), prepare corrosion coupons and place them in microcosms.

9.6.2 Dispense bottom-water then fuel into each microcosm.

9.6.3 Draw pre-test samples and enumerate fuel and bottom-water viable counts (see Practice D6469 and section 9.9).

9.7 *Add Challenge Inoculum*—Inoculate test and positive control microcosms with challenge population. Draw time zero ( $T_0$ ) fuel and bottom-water samples (see Practice D6469 and section 9.9).

NOTE 9—Viable count data may be replaced by or augmented with non-conventional data (see Guide E1326 and Test Method D7463).

9.8 *Sampling*—Predetermined intervals, the following protocol is observed.

9.8.1 *Small (<5.0 L) Microcosms:*

9.8.1.1 Use a 10.0 mL sterile glass pipet to recover 1.0 mL of bottom-water. Transfer the sample to a sterile sample vial (screw capped test tube or bottle).

9.8.1.2 Use a sterile syringe to draw a fuel-phase sample per Practice D6974.

9.8.2 *Large ( $\geq 5.0$  L) Microcosms:*

9.8.2.1 Draw a fuel-phase sample per Practice D7464.

9.8.2.2 Use the same procedure to draw a bottom-water sample.

9.9 *Microbiological Testing:*

9.9.1 *Bottom-Water*—Enumerate bottom-water bacteria and fungi using either Practice D5465, Test Methods D7687 or D7978, Guide D8412, or an alternative, Guide E1326 validated ~~nonconventional~~ nonculture test method.

9.9.1.1 Use soy casein digest agar for enumerating *Pseudomonas aeruginosa*; Sabouraud Dextrose Agar with gentamycin 0.5  $\mu\text{g/mL}$  for enumerating *Yarrowia tropicalis*, and 0.01 % potassium tellurite in 1.5 % bacteriological agar *Hormoconis resiniae*.