



Designation: G131 – 96 (Reapproved 2023)^{ε1}

Standard Practice for Cleaning of Materials and Components by Ultrasonic Techniques¹

This standard is issued under the fixed designation G131; 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.

^{ε1} NOTE—Editorially updated formatting in 7.1 in July 2023.

1. Scope

1.1 This practice covers a procedure for the cleaning of materials and components used in systems requiring a high level of cleanliness, such as oxygen, by ultrasonic techniques.

1.2 This practice may be used for cleaning small parts, components, softgoods, etc.

1.3 The values stated in SI units are standard.

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.* Specific precautionary statements are given in **Note 1**.

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.*

2. Referenced Documents

2.1 *ASTM Standards:*²

D1193 Specification for Reagent Water

E1235 Test Method for Gravimetric Determination of Nonvolatile Residue (NVR) in Environmentally Controlled Areas for Spacecraft

F311 Practice for Processing Aerospace Liquid Samples for Particulate Contamination Analysis Using Membrane Filters

F324 Test Method for Nonvolatile Residue of Volatile

Cleaning Solvents Using the Solvent Purity Meter (Withdrawn 1987)³

F331 Test Method for Nonvolatile Residue of Solvent Extract from Aerospace Components (Using Flash Evaporator)

G121 Practice for Preparation of Contaminated Test Coupons for the Evaluation of Cleaning Agents

G122 Test Method for Evaluating the Effectiveness of Cleaning Agents and Processes

3. Terminology

3.1 *Definitions of Terms Specific to This Standard:*

3.1.1 *contaminant (contamination)*, *n*—unwanted molecular and particulate matter that could affect or degrade the performance of the components upon which they reside.

3.1.2 *contaminate*, *v*—a process of applying a contaminant.

3.1.3 *control coupon (witness coupon)*, *n*—a coupon made from the same material and prepared in exactly the same way as the test coupons, which is used to verify the validity of the method or part thereof.

3.1.3.1 *Discussion*—In this practice, the control coupon will be contaminated in the same manner as the test coupons and will be subjected to the identical cleaning procedure.

3.1.4 *degas*, *v*—the process of removing gases from a liquid.

3.1.5 *nonvolatile residue (NVR)*, *n*—residual molecular and particulate matter remaining following the filtration and controlled evaporation of a solvent containing contaminants.

3.1.6 *particle (particulate contaminant)*, *n*—a piece of matter in a solid state with observable length, width, and thickness.

3.1.6.1 *Discussion*—The size of a particle is usually defined by its greatest dimension and is specified in micrometres.

4. Summary of Practice

4.1 A part, material or component is placed in a container containing the cleaning agent. This container is then placed in an ultrasonic cleaner and treated for a given period of time at

¹ This practice is under the jurisdiction of ASTM Committee G04 on Compatibility and Sensitivity of Materials in Oxygen Enriched Atmospheres and is the direct responsibility of Subcommittee G04.02 on Recommended Practices.

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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.

³ The last approved version of this historical standard is referenced on www.astm.org.

the recommended temperature for the cleaning agent. This results in a solution if the contaminant is soluble in the test fluid or a emulsion if the contaminant is not soluble in the test fluid. The cleaning solution combined with the rinse solutions may then be analyzed for particulate, NVR, or total carbon (TC).

4.1.1 In the case of aqueous-based agents, the parts are rinsed after the removal from the cleaning bath and ultrasonically cleaned in reagent water to provide a solution for TC analysis using G TC.

4.1.2 In the case of solvent-based agents, the parts are rinsed with fresh solvent, which is collected and combined with the solvent used in the cleaning process, and the NVR determined using Test Method E1235, Test Method F324, or Test Method F331, as appropriate.

4.1.3 Particulate analyses may be performed by filtering the final cleaning solution. The particles captured by the filter are then counted using Practice F311.

5. Significance and Use

5.1 This practice is suitable for the removal of contaminants found on materials, parts, and components used in systems requiring a high level of cleanliness, such as oxygen. Parts shall have been precleaned to remove visible contaminants prior to using this procedure. Softgoods such as seals and valve seats may be cleaned without precleaning.

5.2 This procedure may also be used as the cleanliness verification technique for coupons used during cleaning effectiveness tests as in Test Method G122.

5.3 The cleaning efficiency has been shown to vary with the frequency and power density of the ultrasonic unit. Low frequencies in the 20 kHz to 25 kHz range have been found to damage soft metals such as aluminum and silver. Therefore, the specifications of the unit and the frequencies available must be considered in order to optimize the cleaning conditions without damaging the parts.

6. Apparatus

6.1 *Ultrasonic Cleaner*, with an operating frequency range between 25 kHz and 90 kHz, a typical power range between 10 W/L and 25 W/L, and a temperature-controlled bath capable of maintaining a temperature between ambient and 70 °C with an accuracy of 2 °C.

6.2 *Parts Pans*, stainless steel container with volumes between 1 L and 4 L.

6.3 *Bracket*, stainless steel device capable of supporting the parts pans in the ultrasonic bath.

NOTE 1—The bracket should be designed to hang in the ultrasonic bath without contact with the bottom.

7. Reagents

7.1 Solvents such as the following may be used: tetrachloroethylene (perchloroethylene), trichloroethylene, methylene chloride, and perfluorinated carbon fluids. **Warning**—Solvents such as tetrachloroethylene (perchloroethylene), trichloroethylene, and methylene chloride have relative low threshold limit values and the user should refer to appropriate

safe handling procedures, particularly in open tanks. Many solvents are not considered to be compatible with oxygen and must be completely removed from cleaned components prior to the use of these components in oxygen systems. The preferred method of removal shall be determined by the user.

7.2 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society where such specifications are available.⁴ Other grades may be used, provided it is first ascertained that the reagent is of sufficiently high purity to permit its use without lessening the accuracy of the determination. Detergents used shall be identified by manufacturer and name (registered trademark, if any).

7.3 *Purity of Water*—The water used shall meet the requirements of Specification D1193, Type II.

8. Procedure

8.1 Sample Preparation:

8.1.1 If cleanliness verification is to be performed on control or test coupons, prepare the coupons in accordance with Practice G121.

8.1.2 If cleanliness verification is to be performed on small parts, measure the total surface area (*S*) in square centimetres or the mass in grams, or both, as applicable, to the nearest tenth of a milligram (*MI*). Record the surface area (*S*) and mass (*MI*).

8.2 Preliminary Procedure:

8.2.1 If a cleaning agent is being used that requires dilution or special preparation, carefully follow the manufacturer's instructions. Use Type II water to prepare the aqueous cleaning agent solutions or as the actual cleaning agent.

NOTE 2—It has been found that many common hydrocarbon based lubricants are effectively removed to acceptable levels using Type II water at 50 °C to 55 °C. Contaminants more difficult to remove, such as fluorinated or silicone based lubricants, have typically been found to require the use of surface active agents. Use Test Method G122 to evaluate the cleaning effectiveness of the proposed cleaning agent.

8.2.2 Fill the ultrasonic bath to the level specified by the manufacturer with water. Place the support bracket in the ultrasonic bath, heat the ultrasonic bath to the desired temperature, and degas the water for 10 min.

8.2.3 Clean the stainless steel sample parts pan to be used. Conduct the sampling procedure using the selected cleaning agent without parts to verify the cleanliness of the parts pan. Use the same sampling and analysis procedures that will be used on the actual parts. Determine the contamination level of the parts pan, the blank value (*B*), which shall be less than the allowable contamination level for the items being cleaned or extracted. If the contamination level of the parts pan is greater than that specified for the parts, reclean the parts pan until the

⁴ 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.