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THE INSTITUTE OF PETROLEUM Designation: D 2276 - 00

Designation: 216/97

Standard Test Method for Particulate Contaminant in Aviation Fuel by Line Sampling¹

This standard is issued under the fixed designation D 2276; 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.

This test method has been approved by the sponsoring committees and accepted by the Cooperating Societies in accordance with established procedures.

This standard has been approved for use by agencies of the Department of Defense.

1. Scope

1.1 This test method covers the determination of particulate contaminant in aviation turbine fuel using a field monitor.

1.2 There are two test methods described. The basic test method is used to evaluate the level of contamination gravimetrically. The second test method, presented in Appendix X1, describes a color rating technique that is used for rapid qualitative assessment of changes in contamination level without the time delay required for the gravimetric determinations by stringent laboratory procedures.

1.3 There are two Annexes and two Appendixes in this test method.

1.3.1 Annex A1 provides some precautionary information regarding the use of the required reagents.

1.3.2 Annex A2 describes a standard practice for obtaining a sample of the particulates present in a flowing stream of aviation turbine fuel.

1.3.3 Appendix X1 describes a test method for rating the particulate level in an aviation turbine fuel on the basis of the color of a filter membrane after sampling the fuel in the field.

1.3.4 Appendix X2 provides some safety precautions to avoid static discharge resulting from the accumulation of electrical charges in the fuel and on the equipment while following the procedures.

1.4 The values stated in SI units are to be regarded as the standard. The values given in parentheses are for information only.

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 and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

- 2.1 ASTM Standards:
- D 1193 Specification for Reagent Water²
- D 1535 Practice for Specifying Color by the Munsell System³
- D 1655 Specification for Aviation Turbine Fuels⁴
- D 2244 Test Method for Calculation of Color Differences from Instrumentally Measured Color Coordinates³
- D 4865 Guide for Generation and Dissipation of Static Electricity in Petroleum Fuel Systems⁵
- D 5452 Test Method for Particulate Contamination in Aviation Fuels by Laboratory Filtration⁶

3. Terminology

3.1 Definitions:

3.1.1 *membrane color*, *n*—a visual rating of particulate on a filter membrane against ASTM Color Standards.

3.1.2 *membrane filter*, n—a porous article of closely controlled pore size through which a liquid is passed to separate matter in suspension.

3.1.2.1 *Discussion*— RR: D02-1012 contains information on membrane filters that meet the requirements therein.

3.1.3 *monitor*, *n*—something that reminds or warns.

3.1.3.1 *Discussion*—A plastic holder for a membrane filter held in a field sampling apparatus.

3.1.4 *particulate, adj*—of or relating to minute separate particles.

3.1.4.1 *Discussion*—Solids generally composed of oxides, silicates, and fuel insoluble salts.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *volatile fuels, n*—relatively wide boiling range volatile distillate.

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¹ This test method is under the jurisdiction of ASTM Committee D02 on Petroleum Products and Lubricants and is the direct responsibility of Subcommittee D02.J0 on Aviation Fuels.

Current edition approved June 10, 2000. Published August 2000. Originally published as D 2276–64T. Last previous edition D 2276–99.

² Annual Book of ASTM Standards, Vol 11.01.

³ Annual Book of ASTM Standards, Vol 06.01.

⁴ Annual Book of ASTM Standards, Vol 05.01.

⁵ Annual Book of ASTM Standards, Vol 05.02.

⁶ Annual Book of ASTM Standards, Vol 05.03.

3.2.1.1 *Discussion*—These are identified as Jet B in Specification D 1655 or the military grade known as JP-4.

4. Summary of Test Method

4.1 A known volume of fuel is filtered through a preweighed test membrane filter in a field monitor and the increase in membrane filter mass determined after washing and drying. The change in mass of a control membrane filter located immediately below the test membrane filter is also determined. The particulate contaminant is determined from the increase in mass of the test membrane filter relative to the control membrane filter.

4.2 This test method employs a field monitor to filter a sample of fuel that is taken in the field by the sampling procedure detailed in Test Method D 5452.

4.3 For situations where it is not possible to take a field monitor sample, procedures are given in Test Method D 5452 for the determination of particulate contaminant in a fuel sample by laboratory filtration.

4.4 Appendix X1 describes a method for color-rating used filter membranes.

5. Significance and Use

5.1 This test method provides a gravimetric measurement of the particulate matter present in a sample of aviation turbine fuel by line sampling. The objective is to minimize these contaminants to avoid filter plugging and other operational problems. Although tolerable levels of particulate contaminants have not yet been established for all points in fuel distribution systems, the total contaminant measurement is normally of most interest. The Appendix X1 color rating method is useful for fuel system monitoring purposes. No quantitative relationship exists between gravimetric and color rating test results.

RR:D02-1012 is available from ASTM Headquarters. To obtain the research report.

data, and list of suppliers request RR:D02-1012, 1994.

NOTE 1—Matched weight membrane filters,⁷ 37-mm diameter, nominal pore size $0.8 \mu m$, may be used as test and control membrane filters if so desired. Use of matched-weight membrane filters precludes the necessity for carrying out subsequently the procedures detailed in Section 8.

6.8 *Dispenser for Flushing Fluid*, 0.45- μ m membrane filters to be provided in the delivery line (see Fig. 1).

6.9 *Field Monitors*, ⁷ complete with protective plugs and 34-mm support pads.

6.10 *Air Ionizer*, for the balance case (see Note 2 and Note 3).

NOTE 2—When using a solid-pan balance, the air ionizer may be omitted provided that, when weighing a membrane filter, it is placed on the pan so that no part protrudes over the edge of the pan.

Note 3-Air ionizers should be replaced within 1 year of manufacture.

6.11 *Multimeter/VOM*, used for determining whether electrical continuity is 10 Ω or less between 2 points.

6.12 *Flushing Apparatus*, of the type shown in Fig. 2. It consists of a receiving flask large enough to contain the flushing fluid and shall be equipped with a side arm to connect to the vacuum system. Reagent resistant tubing shall be arranged to allow passage of a grounding wire. An assembly of reagent grade resistant tubing and bung fitted with a glass tube shall be assembled as shown in Fig. 2 to attach to a field monitor.

6.13 *Ground/Bond Wire*, Nos. 10 through 19, (0.912 to 2.59 mm) bare stranded flexible stainless steel or copper installed in the flask and grounded as shown in Fig. 2.

7. Reagents

7.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents shall conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society,

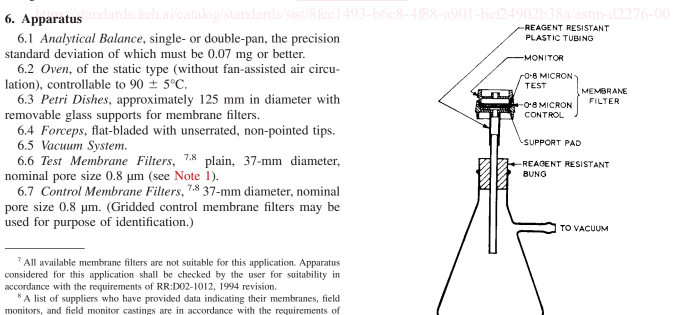
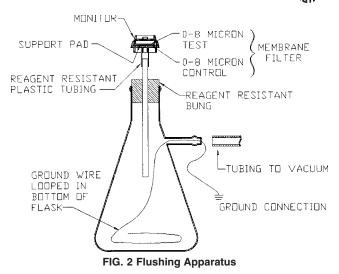


FIG. 1 Dispenser for Flushing Liquid

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where such specifications are available.⁹ 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.

7.2 *Purity of Water*— Unless otherwise indicated references to water shall be understood to mean reagent water as defined by Type III of Specification D 1193.

7.3 *Isopropyl Alcohol*, reagent grade. (Warning– Flammable. See A1.1.)

7.4 Liquid Detergent, water-soluble.

7.5 Flushing Fluids:

7.5.1 Petroleum Spirit (also known as petroleum ether or IP Petroleum Spirit 40/60) (Warning—Extremely flammable. Harmful if inhaled. Vapors are easily ignited by electrostatic discharges, causing flash fire. See A1.2.), having boiling range from 35 to 60°C.

8. Preparation of Test and Control Membrane Filters and Field Monitors Prior to Sampling

8.1 Two 37-mm membrane filters of nominal pore size 0.8 µm are required: a test and a control membrane filter. Matched-weight membrane filters may be used if so desired (see Note 1). If matched-weight membrane filters are used, it is unnecessary to carry out the procedures detailed in this section because they have been carried out previously by the membrane filter supplier. The two membrane filters used for each individual test should be identified by marking the petri dishes used as containers. Glassware used in preparation of membrane filters shall be cleaned as described in Section 10.

8.1.1 Using forceps, place the test and control membrane filters side by side in a clean petri dish. To facilitate handling the membrane filters should rest on clean glass support rods in the petri dish.

8.1.2 Place the petri dish with its lid slightly ajar, in an oven at 90 \pm 5°C and leave it for 30 min.

8.1.3 Remove the petri dish from the oven and place it near the balance. The petri dish cover should be ajar but still protecting the membrane filters from contamination from the atmosphere. Allow 30 min for the membrane filters to come to equilibrium with the ambient air temperature and humidity.

8.1.4 Remove the control membrane filter from the petri dish with forceps, handling by the edge only, and place it centrally on the weighing pan. Weigh it and return it to the petri dish.

8.1.5 Repeat 8.1.4 for the test membrane filter. Record the membrane filter masses.

8.1.6 Take a clean field monitor, mark for identification, rinse with filtered flushing fluid, and insert a clean support pad.

8.1.7 Using clean forceps, place the weighed control membrane filter centrally on the support pad in the field monitor and place the weighed test membrane filter on top of the control membrane filter. Assemble the two parts of the field monitor, ensuring that the membrane filters are firmly clamped inside and the protective plugs are in position.

8.1.8 Record the monitor identification.

9. Sampling

9.1 When possible, 3.785 L (1 gal) to 5 L of fuel (Warning—Jet A, combustible. Vapor harmful. See A1.4.) (Warning—Jet B, extremely flammable. Harmful if inhaled. Vapors may cause flash fire. See A1.4.) should be passed through the monitor during field sampling. The sample volume actually employed shall be reported. See Annex A2 for specific details of sampling practices.

10. Preparation of Apparatus

10.1 Follow the procedure utilizing the apparatus shown in Fig. 2. Alternative apparatus may be used provided that it achieves the same end.

10.1.1 Wash the petri dishes and supports with warm water containing detergent. Then rinse with warm water and finally with distilled water.

10.1.2 Rinse thoroughly with filtered isopropyl alcohol.

10.1.3 Rinse thoroughly with filtered flushing fluid.

10.1.4 Drain for a few seconds, and then air or oven dry.

10.2 Ensure that all glass and plastic tubing attached to the solvent filtering dispenser is clean by flushing thoroughly with filtered flushing fluid.

11. Procedure

11.1 Assemble the apparatus shown in Fig. 2 with the field monitor in place on the stopper of the vacuum flask.

NOTE 4—Take care to ensure that monitors are tightly closed and preferably clamped. Spring paper clips have been found suitable for this purpose.

11.2 Place the tip of the delivery spout of the solvent filtering dispenser in direct contact with the monitor inlet hole. Introduce filtered flushing fluid.

11.3 Apply vacuum to the flask and allow approximately 250 mL of filtered flushing fluid to pass from the solvent filtering dispenser through the monitor and into the vacuum flask.

⁹ Reagent Chemicals, American Chemical Society Specifications, 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.

11.4 Remove the solvent filtering dispenser and slowly release the vacuum.

11.5 Remove the monitor from the stopper of the vacuum flask and carefully dismantle it in an upright position.

11.6 Carefully remove the test and control membrane filters, and place side by side on clean glass supports in a clean, covered petri dish.

NOTE 5-The test and control membrane filters can be removed from the monitor by pushing upwards against the support pad through the outlet orifice with a thin dowel.

11.7 Dry and reweigh the membrane filters as described in 8.1.2-8.1.5, taking care not to disturb the contaminant on the surface of the test membrane filter.

12. Calculation and Report

12.1 Subtract the initial mass of the test membrane filter, W_1 , from the final mass, W_2 .

12.2 Subtract the initial mass of the control membrane filter, W_3 , from the final mass, W_4 .

12.3 Divide the correct mass of contaminant ($W_2 - W$ 1) – $(W_4 – W_3)$ by the volume of sample filtered and report the result as total contaminant, expressed in milligrams per litre.

NOTE 6-If matched-weight membrane filters have been used for the test (see Note 1), then $W_1 = W_3$ and the corrected mass of contaminant in 12.3 becomes $W_2 - W_4$.

12.4 Report the result to the nearest 0.01 mg/L, and also the sample volume used in the test.

13. Precision and Bias ¹⁰

13.1 The precision of this test method is not known to have been obtained in accordance with currently accepted guidelines in Committee D-2 RR:D02-1007.

13.2 These precision data have been obtained by statistical examination of test results using 5-L samples and were first published in 1966.

¹⁰ Supporting data are available from ASTM Headquarters. Request RR:D02-1197

13.3 Repeatability— The difference between successive results obtained by the same operator with the same apparatus under constant operating conditions on identical test material would, in the long run, in the normal and correct operation of the test method, exceed the following values in only one case in twenty:

Range	Repeatability
0.0 to 2.0 mg/L	0.175x + 0.070

where x is the average value of two results.

13.4 Reproducibility- The difference between two single and independent results obtained by different operators working in different laboratories on identical test material would, in the long run, exceed the following values in only one case in twenty:

Range	Reproducibility
0.0 to 2.0 mg/L	0.444 <i>x</i> + 0.178

where x is the average value of two results.

13.5 Typical values are given in Table 1.

NOTE 7-Reproducibility values were determined through cooperative testing by different operators using separate apparatus working at the same location using identical test material. This procedure was adopted as it is highly improbable, if not impossible, to ensure the obtaining of "identical test material" when testing at different locations.

13.6 Bias—The procedure given for the determination of particulate contaminant in aviation turbine fuels has no bias since this property can only be defined in terms of this test method.

14. Keywords

14.1 aviation fuel; color rating; field monitor; gravimetric contaminant; membrane color; membrane filter; particulate

TABLE 1 Statistical Information for Particulate Contaminant											
Average Result, mg/L	0.0	0.1	0.2	0.3	0.5	0.7	1.0	1.5	2.0		
Repeatability	0.07	0.09	0.11	0.12	0.16	0.19	0.25	0.33	0.42		
Reproducibility	0.18	0.22	0.27	0.31	0.40	0.49	0.62	0.84	1.07		

ANNEXES

(Mandatory Information)

A1. PRECAUTIONARY STATEMENTS

A1.1 Isopropyl Alcohol

A1.1.1 Keep away from heat, sparks, and open flame.

- A1.1.2 Keep container closed.
- A1.1.3 Use with adequate ventilation.
- A1.1.4 Avoid prolonged breathing of vapor or spray mist.
- A1.1.5 Avoid contact with eyes and skin.

A1.1.6 Do not take internally.

A1.2 Petroleum Ether

A1.2.1 Keep away from heat, sparks, and open flame.

A1.2.2 Keep container closed.

A1.2.3 Use with adequate ventilation.

A1.2.4 Avoid build-up of vapors and eliminate all sources of ignition, especially nonexplosion-proof electrical apparatus and heaters.

A1.2.5 Avoid prolonged breathing of vapor or spray mist. A1.2.6 Avoid prolonged or repeated skin contact.

A1.3 Aviation Turbine Fuel (Jet A or A-1, see **Specification D 1655**)

A1.3.1 Keep away from heat, sparks, and open flames.

A1.3.2 Keep container closed.

and heaters.

A1.3.3 Use with adequate ventilation.

A1.3.4 Avoid breathing vapor or spray mist.

A1.3.5 Avoid prolonged or repeated contact with skin.

A1.4 Aviation Turbine Fuel (Jet B, see Specification D 1655)

A1.4.1 Keep container closed.

A1.4.2 Use with adequate ventilation.

A2. SAMPLING AVIATION TURBINE FUEL FOR PARTICULATE CONTAMINATION

A2.1 Scope

A2.1.1 This test method covers taking samples of aviation turbine fuels from fuel handling systems under pressure, through field monitors, for the determination of particulate contaminant.

A2.2 Summary of Test Method

A2.2.1 A 3.785 to 5-L sample is taken from a flowing line or pipe and passed under line pressure through a field monitor containing a preweighed 0.8-µm test membrane filter and a preweighed 0.8-µm control membrane filter. After filtration the field monitor is returned to a laboratory for analysis.

A2.3 Apparatus

A2.3.1 Sampling Point, a suitable tapping in the pipe, line, or fitting to accept the sampling valve connection (see A2.3.2.1). If using an existing tapping, it may be necessary to use reducing bushings to ensure the proper tapping size for the sampling connection. Care shall be taken in such cases to avoid trapping or generating contaminant. A sampling probe projecting into the fuel stream aids in guarding against this situation. If a shut-off valve is desired, a stainless steel ball or plug-type valve should be used.

A2.3.2 *Sampling Apparatus*, ⁵ as illustrated in Fig. A2.1 and consisting of the following components:

A2.3.2.1 Sampling Valve Connection, designed to meet the following requirements: (1) It shall be mounted in the sampling point and must incorporate a self-sealing quick action coupling designed to mate with a suitable connection leading to the selector valve of the sampling assembly. (2) It must be completely resistant to fuel and be leak proof up to the maximum working pressures to be encountered. (3) It must have a minimum of internal recesses which could cause the holdup of contaminant. (4) It must be provided with a dust cap.

A2.3.2.2 Flexible Pressure Hose, if used, designed to meet the following requirements: (1) It must be permanently connected to half of the self-sealing sampling connection and be capable of being attached to the selector valve on the other end. (2) It must be completely resistant to fuel. (3) It should be less than 18 in. (457 mm) long and be provided with a dust cap for the sampling connection.

A2.3.2.3 Selector Valve, designed to meet the following requirements: (1) It must have one inlet port and two alternative outlet ports. (2) It may also have an OFF position but this is not mandatory. (3) It must be so designed that it is free from internal pockets in which contaminant may be stored and

subsequently released. (4) It may incorporate a point to which a syringe can be fixed.

A1.4.3 Avoid build-up of vapors and eliminate all sources

of ignition, especially nonexplosion-proof electrical apparatus

A1.4.5 Avoid prolonged or repeated contact with skin.

A1.4.4 Avoid breathing vapor or spray mist.

A2.3.2.4 *Field Monitor Holder*, so constructed that a perfect seal is made between its upper part and the top of the field monitor, and also between its lower part and the bottom of the field monitor. No fuel bypassing can be permitted.

A2.3.2.5 *Field Monitors*, complete with protective plugs and each containing two 37-mm preweighed 0.8-µm membrane filters backed by a 34-mm support pad, prepared as described in Section 8.

A2.3.2.6 *Graduated Receiver*, capable of receiving at least a 5-L fuel sample. The receiver shall be suitably electrically bonded (see Note A2.1).

NOTE A2.1—A metal receiver is preferable to one made of plastic. If a plastic receiver is employed, all metal components shall be grounded and a grounded wire or other conductor shall be inserted in the receiver to pick up electrostatic charges in the fuel.

A2.3.3 *Back Pressure Connection*, for sampling from pipes or lines in which the pressure is too low to obtain a proper fuel sample in a reasonable time. A suitable connection is illustrated in Fig. A2.2. The sampling connection should not be located in this equipment.

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A2.4 General Precautions

A2.4.1 Always handle the sampling equipment with care and ensure that it is maintained in a scrupulously clean condition.

A2.4.2 To avoid extraneous contaminant, field monitor protective plugs must be removed only for sampling and replaced immediately. The monitor must be opened only in a laboratory.

A2.4.3 Under no circumstances should thread-sealing compounds be used. TFE-fluorocarbon pipe thread sealant must be used, but if the apparatus still leaks, abandon the test.

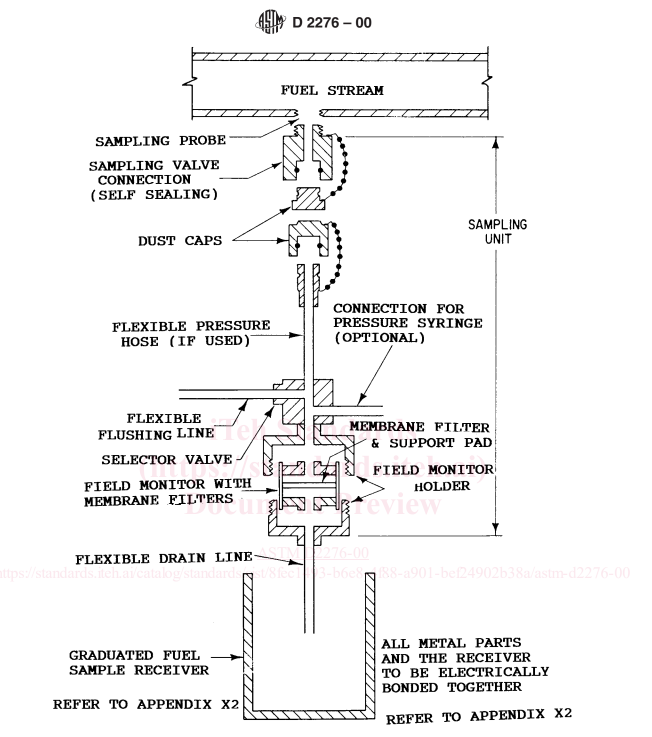
A2.4.4 All metal parts of the sampling apparatus must be electrically bonded together and grounded.

A2.5 Procedure

A2.5.1 Fix the self-sealing connection to the sampling point ensuring that the dust cap at the exposed ends is in place. This sampling connection is normally left in place at all times.

A2.5.2 Unscrew the field monitor holder.

A2.5.3 Remove the bottom protective plug from the monitor and place it in a clean, safe place. Where monitor holder construction requires that the monitor be first placed into the upper half of the casing, remove the upper dust plug.



NOTE 1—All metal parts and the receiver are to be electrically bonded together. FIG. A2.1 Field Sampling Apparatus

A2.5.4 Locate the monitor in its holder.

NOTE A2.2—Examine the monitor carefully to ensure that it is located correctly in its holder (that is, not reversed). The bottom (outlet) of the monitor is the side with the spiderweb and leads directly to the sample receiver. The top (inlet) side of the monitor is the upper portion, which has space for fuel above the filter membrane.

A2.5.5 Remove the other protective plug from the monitor and place it in a clean, safe place.

A2.5.6 Reassemble the monitor holder handtight only. Extreme force is both unnecessary and undesirable.

