



Standard Test Method for Silica—Oil Absorption Number (OAN)¹

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1. Scope

1.1 This test method covers the determination of the oil absorption number (OAN) of silica.

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

1.3 *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:*²

[D1799 Practice for Carbon Black—Sampling Packaged Shipments](#)

[D1900 Practice for Carbon Black—Sampling Bulk Shipments](#)

[D2414 Test Method for Carbon Black—Oil Absorption Number \(OAN\)](#)

[D6738 Test Method for Precipitated Silica—Volatile Content](#)

[E177 Practice for Use of the Terms Precision and Bias in ASTM Test Methods](#)

[E691 Practice for Conducting an Interlaboratory Study to Determine the Precision of a Test Method](#)

3. Summary of Test Method

3.1 In this test method, oil is added by means of a constant-rate burette to a sample of silica in the mixer chamber of an absorptometer. As the sample absorbs the oil, the mixture changes from a free-flowing state to one of a semiplastic agglomeration, with an accompanying increase in viscosity.

¹ This test method is under the jurisdiction of ASTM Committee D11 on Rubber and is the direct responsibility of Subcommittee D11.20 on Compounding Materials and Procedures.

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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.

This increased viscosity is transmitted to the torque-sensing system of the absorptometer. The test is stopped when a torque level has been reached. Preferably the torque versus volume of oil is recorded by a penwriter or by a data acquisition system allowing a reliable determination of the endpoint. The volume of oil per unit mass of silica is the oil absorption number (OAN).

4. Significance and Use

4.1 The oil absorption number of a specific silica is related to the processing and vulcanizate properties of rubber compounds containing the silica.

5. Apparatus³

5.1 *Balance*, analytical, with a sensitivity of 0.001 g.

5.2 *Sieve*, 500 μm (U.S. standard No. 35), having a diameter of 200 mm (8 in.) and a height of 25 mm (1 in.).

5.3 *Bottom Receiver Pan*.

5.4 *Oven*, gravity-convection type, capable of temperature regulation within $\pm 1^\circ\text{C}$ at 105°C and temperature uniformity within $\pm 5^\circ\text{C}$.

5.5 *Spatula*, rubber, 100-mm.

5.6 *Absorptometer*,⁴ equipped with a constant-rate burette that delivers $4 \pm 0.024 \text{ cm}^3/\text{min}$.

5.7 *Desiccator*, with silica gel as desiccant.

6. Reagents and Standards

6.1 *Oil*:

6.1.1 *n-Dibutyl Phthalate*,⁵ having a density of 1.042 to $1.047 \text{ mg}/\text{m}^3$ (g/cm^3) at 25°C .

6.1.2 *di-octyl-adipate (DOA)*, having a density of $0.9255 \text{ g}/\text{cm}^3$ at 20°C , a refractive index of 1.447 at 20°C , and a kinematic viscosity of 10 to $34 \text{ mm}^2/\text{s}$ (cSt) at 40°C .

6.1.3 *Epoxidized fatty acid ester (EFA)*, meeting the specifications listed in Test Method [D2414](#), Annex A4.

³ All apparatus is to be operated and maintained in accordance with the manufacturer's directions for optimum performance.

⁴ Available from C. W. Brabender Instruments, Inc., 50 E. Wesley St., Hackensack, NJ 07606, website: www.cwbrabender.com, and HITEC Luxembourg, 5 rue de l'Eglise, L-1458 Luxembourg, website: www.hitec.lu.

⁵ Technical grade has turned out to be suitable for the test, provided that the density is in the specified range.

6.2 *Silica*, commercial grade with a nitrogen surface area of $175 \pm 10 \text{ m}^2/\text{g}$.

7. Sampling

7.1 Samples shall be taken in accordance with Practices [D1799](#) and [D1900](#).

8. Calibration

8.1 *Absorptometer*⁶—The absorptometer is composed of components that influence calibration: the dynamometer torque spring or the load cell, the torque-limit or the indicator set point, the oil damper (absorptometers Type C, E, and H are equipped with electronic damping), and the mixer-measuring head.⁷ It is necessary that each of the components be in good condition or proper adjustment to achieve acceptable calibration.

NOTE 1—Stainless steel mixing chambers⁸ have been found satisfactory for this test when they are manufactured to a roughness average (Ra) of $2.5 \pm 0.4 \mu\text{m}$ ($100 \pm 15 \mu\text{in.}$) based upon eight measurements. No single measurement should be greater than $3.6 \mu\text{m}$ ($140 \mu\text{in.}$) or less than $1.5 \mu\text{m}$ ($60 \mu\text{in.}$). Stainless steel bowls purchased with an absorptometer have been pre-polished for 16 h to minimize bowl surface changes affecting calibration during their initial use. It is recommended that new replacement stainless steel bowls should also be pre-polished to minimize the bowl surface effects on calibration (see Annex A1).

8.1.1 The torque indicator is the primary component used to correct calibration. The load cell tension is adjusted by varying the alarm shut-off set point. Proper adjustment on the torque indicator should provide repeatable values for a silica sample dedicated to internal reference.

8.1.2 The maximum torque span is set at 10 000 mNm (10 000 units) torque value. The torque-limit alarm is initially set at 5000 mNm (5000 units), but for testing silicas it will be necessary to adjust this setting to a lower value in order to obtain reproducible results. Use an internal silica sample with a nitrogen surface area of $175 \pm 10 \text{ m}^2/\text{g}$ to set the torque limit alarm which should correspond to approximately 70 % of the maximum torque developed during the test. After calibration, this setting should not be changed.

NOTE 2—It is generally recommended to use the absorptometer in conjunction with a penwriter or preferably with a data acquisition system (see [9.10](#) for further details).

8.1.3 All digital signals are preset at 3 s damping for the torque sensing system.

8.1.4 Properly maintain the surface finish of the mixing chamber. If a new mixer chamber is installed, frequently monitor the instrument for any drift in calibration.

8.2 *Constant-Rate Burette*—The delivery rate of the burette is to be $4 \text{ cm}^3/\text{min}$. See [Annex A1](#) for detailed instructions on the procedure for calibration check of the constant-rate burette.

⁶ Mechanical absorptometers (type A or type B) can be used for the test; however, they are no longer commercially available. Refer to the instructions of the supplier for calibration procedure.

⁷ The rotor motor speed is 1.31 rad/s (125 r/min).

⁸ Replacement stainless steel bowls which have been found to be satisfactory are available from Titan Specialties, Inc., P.O. Box 2316, Pampa, TX 79066-2316, and C. W. Brabender Instruments, Inc., 50 E. Wesley St., S. Hackensack, NJ 07606, website: www.cwbrabender.com, HITEC Luxembourg, 5 rue de l'Eglise, L-1458 Luxembourg, website: www.hitec.lu, and Titan Specialties, Inc. P.O. Box 2316, Pampa, TX 79066-2316.

9. Procedure

9.1 Pass a suitable amount of the sample through Sieve No. 35 (500 μm), using a brush in order to deagglomerate larger particles. Use 2 g of the sieved material to test the moisture content (see [9.3](#)) as volatile matter according to Test Method [D6738](#).

9.2 Determine the amount of moisture in the silica under test by weighing 2 g of the sieved silica (see [9.1](#)) into a dish to the nearest 0.001 g. Place the dish into an oven set at 105°C , leave it inside for 2 h, cool in a desiccator and weigh to the nearest 0.001 g. See Section [10](#) (Calculation) for details of moisture calculation.

9.3 Weigh 12.5 g of the sample to the nearest 0.01 g.

NOTE 3—For silicas with an extraordinary high pour density it may be necessary to increase the sample mass used for the test. This modification has to be mentioned in the test report.

9.4 It is recommended that a testing temperature of $23 \pm 5^\circ\text{C}$ be maintained, as measured by a thermocouple in the mixing bowl. If a temperature controllable mixing bowl is not available, keep the bowl temperature below 30°C and comply with [Note 4](#) while running the samples.

NOTE 4—If the absorptometer has remained idle for more than 15 min and a temperature controllable bowl is not being used, a 10-min warm-up sample must be run before beginning a test. It is important that the mixer chamber temperature be kept uniform. Preferably, allow 5 min between the end of one test and the start of another.

9.5 Transfer the sample to the absorptometer mixer chamber and replace the cover.

9.6 Place a waste receptacle under the delivery tube. Make sure that the tube is free of air bubbles by delivering approximately 1 cm^3 of oil into the waste receptacle.

9.7 Verify the drive speed is set to 1.31 rad/s (125 r/min).

9.8 Position the burette delivery tube over the hole in the mixer chamber cover or use the accessory funnel. Set the burette digital counter to zero.

9.9 Activate the “start” buttons simultaneously or use the start procedure given in the software. The apparatus will operate until sufficient torque has developed to activate the torque-limit switch, which will halt the absorptometer and burette.

9.10 Record the volume of oil used as indicated by the burette digital counter.

NOTE 5—If a penwriter is used to record the torque curve, deactivate the automatic cut-off by setting the torque limit to 10 000. Stop the test when the torque maximum has been recorded unequivocally. Mark on the curve the oil volume corresponding to the maximum torque and measure the height (in mm or in.) of the maximum. At the left side of the maximum, identify the point corresponding to a height of 70 % of the maximum of the curve. Measure the distance on the x -axis from the start point to this point and convert the value to volume of oil as follows:

$$\text{Volume oil} = \text{delivery rate of burette} \cdot \text{distance} / \text{speed of penwriter}$$

NOTE 6—If a data acquisition system⁹ is used, the absorptometer will stop after having recorded the torque maximum, and the test result (in

⁹ OAN Data Acquisition Systems are available from C.W. Brabender Instruments, Inc., 50 E. Wesley St., S. Hackensack, NJ 07606, website: www.cwbrabender.com, and HITEC Luxembourg, 5 rue de l'Eglise, L-1458 Luxembourg, website: www.hitec.lu.