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Designation: D 2228 – 02

Standard Test Method for Rubber Property—Relative Abrasion Resistance (by the Pico Abrader) Method¹

This standard is issued under the fixed designation D 2228; 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.

¹ This test method is under the jurisdiction of ASTM Committee D-11 on Rubber and is the direct responsibility of Subcommittee D11.15 on Degradation Tests .
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1. Scope

1.1 This test method covers the determination of the abrasion resistance of soft vulcanized rubber compounds (thermoset rubbers, thermoplastic elastomers, and elastomeric and similar materials to a standardized reference standard system. A reference or standard standardized set of reference compounds is used to calculate relative abrasion resistance. These reference compounds are also used to determine the relative performance, within a permissible range, of the cutting knives used in performing the test.

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 All materials, instruments, or equipment used for the determination of mass, force, or dimension shall have traceability to the National Institute for Standards and Technology,² or other internationally recognized organization parallel in nature.

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

2. Referenced Documents

2.1 *ASTM Standards:*

D 613498 Practice for Rubber—Standard Temperatures Conditioning Plastics for Testing³

D 3182 Practice for Rubber—Materials, Equipment, and Procedures for Mixing Standard Compounds and Preparing Standard Vulcanized Sheets⁴

D 4483 Practice for Determining Precision for Test Method Standards in the Rubber and Carbon Black Industries⁴

3. Summary of Test Method

3.1 In this test method, a pair of tungsten carbide cutting knives of a specified geometry and sharpness is configuration are used to abrade the surface of ~~rubber~~ the specimen. The knives are lowered onto a circular test specimen, or button, which is rotated under controlled conditions of speed, time, and force on the cutting knives. A dusting powder is used ~~at the~~ as an interface between the cutting knives and the specimen to engulf the abraded rubber particles; and to maintain the cutting knives relatively free from oils, resins, etc., which ~~might may~~ be present in the ~~specimen~~. ~~These oils, resins, etc., specimen and may~~ interfere with the abrasion assessment. A series of five calibration compounds ~~is~~ are used to determine that the sharpness of the knives and hence, the calibration of the instrument, are within the specified limits, and additionally, as reference standards to which the abrasion resistance, determined by volumetric loss, of any experimental a subject material may be compared.

4. Significance and Use

4.1 The test method may be used to estimate the relative abrasion resistance of ~~different rubber compounds~~ subject materials as described in 1.1. No correlation between this accelerated test and service performance is given or implied, due, in part, to the widely varying nature of service conditions.

4.2 The formulas, for which the mixing and curing specifications are given in Annex A1, once prepared, are referred to as calibration compounds. These calibration compounds may be used to determine the performance status of the cutting knives as described herein.

4.3 The performance of the cutting knives may also be determined by periodically determining their dimensions as described in 6.1.7.

4.4 The calibration compounds are used as reference standards to which the abrasion resistance, determined by volume loss of a subject material, may be compared.

4.5 Once the resistance to abrasion is established, using this methodology, for a specific material, the results achieved may be used as a basis for future comparative analysis of identical materials, either as agreed upon between laboratories, or between customer and supplier.

5. Interference

5.1 This test method is conducted under controlled conditions, except for the sharpness of the ~~abrasion~~ cutting knives. ~~Different rubbers behave differently~~ The behavior of the materials, as described in 1.1, yield varying results with respect to the sharpness of the cutting knives.³ ~~For the most uniform test results, the knives must~~ This variation can be maintained within minimized by maintaining the specifications of 9.5. knives in accordance with recommendations outlined in Section 9.

³ For further information, see Research Report RR: D-11-1022. A copy is available from

³ *Annual Book of ASTM Headquarters, 100 Barr harbor Drive, West Conshohocken, PA 19428-2959; Standards, Vol 08.01.*

⁴ Alon-C undensed aluminum oxide, manufactured by the Cabot Corp., Boston, MA, has been found satisfactory.

⁴ *Annual Book of ASTM Standards, Vol 09.01.*

6. Apparatus

6.1 *Pico Tester*—An overall view of the apparatus is illustrated in Fig. 1. The pico tester itself is illustrated in Fig. 2.

6.1.1 *Turntable*, on which the test specimen is mounted and rotated, having the capability of maintaining 1.00 ± 0.03 Hz (rps) throughout the duration of a test cycle (see Section 10).

6.1.2 *Instrument Frame*, with armature assembly that holds and lifts the cutting knives. Mounted on top of the assembly is a “dead-weight load box” in which masses (weights) may be placed to regulate the force on the cutting knives (see Section 10). The assembly moves freely in a bearing housing that permits vertical motion but counteracts the reaction torque on the cutting knives, thus preventing rotation. Vertical travel, once knives have been lowered onto the test specimen, is restricted by an arm lock.

6.1.3 *Drive Motor*, with forward, reverse, and stop controls to govern the operation of the turntable.

6.1.4 *Dusting Powder Reservoir and Feeder Tubes*, capable of supplying a uniform flow of dusting powder at the rate of 5 mg/s to the interface of the cutting knives and test specimen during operation (see Section 10).

6.1.5 *Vacuum Dust Collector*, with vacuum sweeper hose of rubber tubing capable of rapid removal of the dusting powder and debris from the specimen and cutting knife interface.

6.1.6 *Digital Counters*,— a pair mounted diametrically opposed to one another on the turntable support, capable of displaying a total count of no less than 80 and determining the rotations of the turntable to within 1.00 ± 0.03 Hz (rps) throughout the duration of a test cycle (see Section 10).

6.1.7 *Cutting Knives*, tungsten carbide knives manufactured to the specifications in Fig. 2 of Grade 831 Carboly or an equivalent material, at the time of manufacture or resharpening.

6.1.7.1 The cutting knives shall have a “cutting edge” formed by the angle of the two bevels. The apex of the angle shall have a blunted edge, or “flat,” with a width of $10 \pm 5 \mu\text{m}$.

6.1.7.2 At the time of manufacture or resharpening, the “cutting edge” may be less than $10 \pm 5 \mu\text{m}$. The blunted edge, or “flat,” shall then be produced by the end user, manufacturer or resharpening service supplier, by dulling with a diamond dust, or other suitable method to $10 \pm 5 \mu\text{m}$ prior to first use.

NOTE 1—The supplier shall notify the end user of new or resharpened cutting knives of the final specifications of the cutting edges and other dimensional specifications.

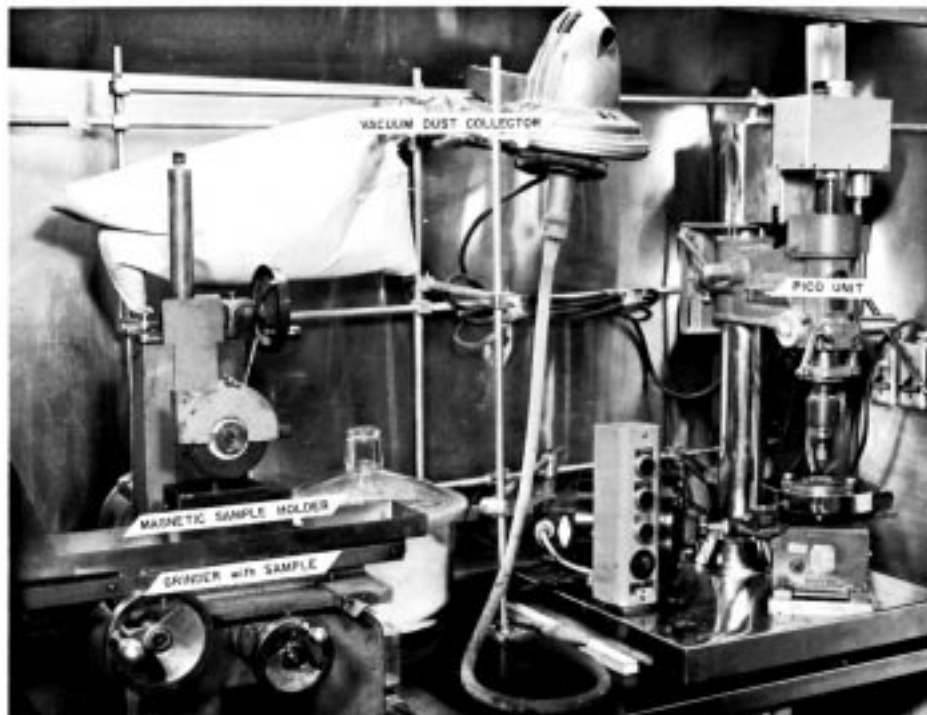


FIG. 1 Typical Pico Tester with Auxiliary Apparatus

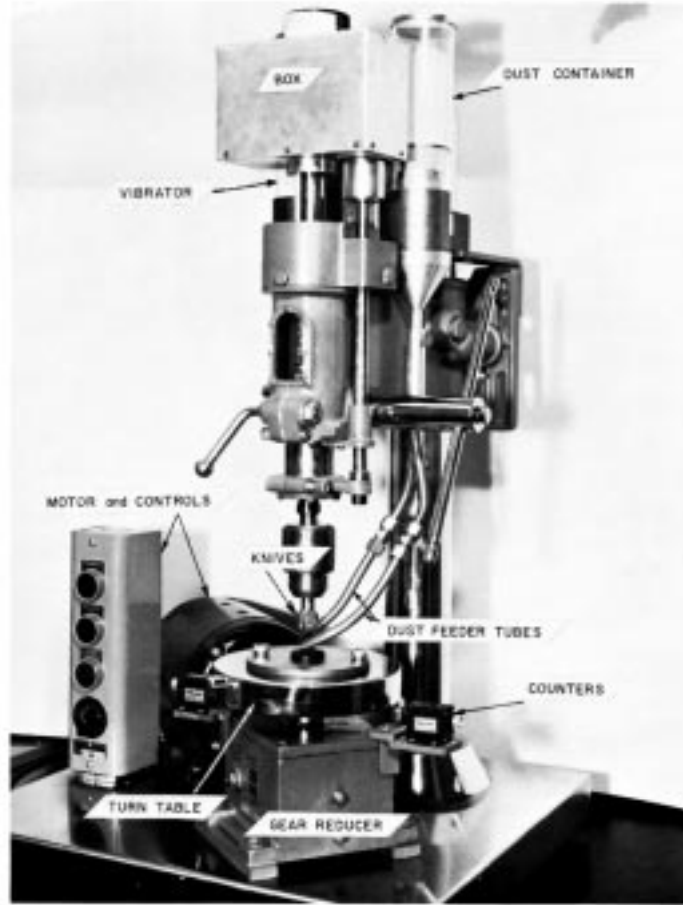


FIG. 2.1 Typical Pico Tester (continued)

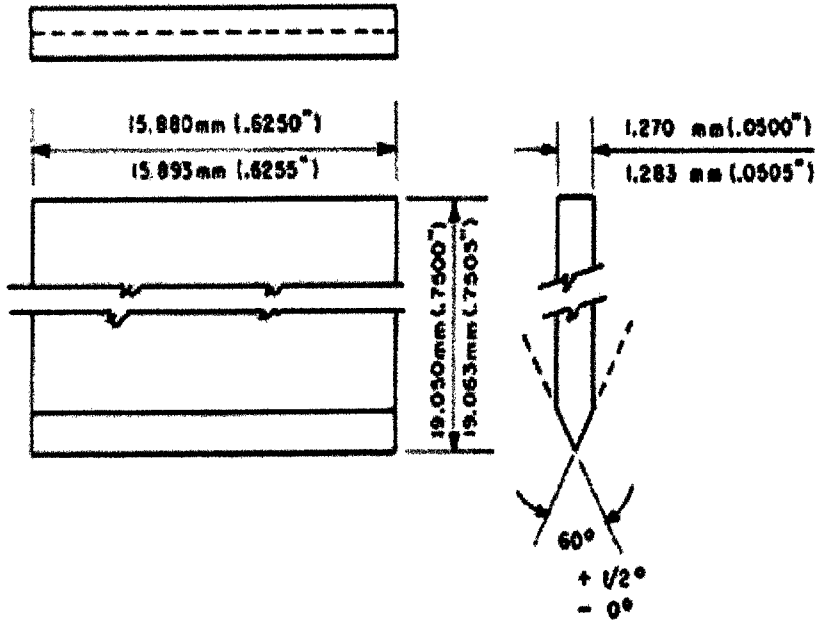


FIG. 2 Pico Cutting Knives

6.1.7.3 The width of the blunted edge, or “flat,” shall be verified either by standard microscopy techniques or scanning electron microscopy.^{5,6}

6.1.7.4 The beveled surfaces shall have a finish equivalent to a No. 4 μm finish at the time of manufacture or resharpener.

6.1.7.5 The cutting knives, at the time of manufacture or resharpener, shall be matched in pairs so that the overall dimensions of each of the three major axis have a difference between them individually no greater than 0.013 mm (0.0005 in.) and that they are parallel to within ± 0.0065 mm (0.00025 in.).

6.2 *Grinder*, for preparing the surfaces of test specimens. The grinder shall be equipped with a:

6.2.1 A magnetic plate for holding the specimen in place, a

6.2.2 A micrometer adjustment for capable of controlling the vertical movement of the abrasive wheel, and a wheel in 0.025 mm (0.001 in.) increments,

6.2.3 A handwheel for traversing the specimen,

6.2.4 An electric motor with a spindle having a rotational frequency shall be of 95 ± 3 Hz (rps). The size of (rps) and equipped with an arbor to secure the abrasive wheel shall be about (see Section 10), and

6.2.5 An abrasive wheel with a diameter of no less than 100 mm (4 in.) in diameter and 12.5 mm (0.5 in.) in width, with new, and a center mounting hole of 12.7 mm (0.5 in.) in diameter. The grit grade of the wheel shall be equivalent to Carborundum C30LB.

6.3 *Vacuum Dust Collector*, with vacuum sweeper hose of gum rubber tubing having a tapered tip to provide rapid removal of the dusting powder and engulfed particles from the specimen.

6.4 *Balance*, accurate to ± 0.0001 g.

7. Auxiliary Materials

7.1 *Dusting Powder*:

7.1.1 The dusting powder used shall be a blend of equal parts by weight of aluminum oxide and diatomaceous earth. The diatomaceous earth should first be passed through a No. 200 (75-μm) screen and the retained material discarded.⁶

7.1.2 An equal part^{7,8}

7.1.2 A mixture of the two powders materials, in equal parts, shall be thoroughly blended, densed; and screened. For When preparing small quantities, the following procedure is satisfactory: i

7.1.2.1 Into a sturdy ceramic wide-mouth, 4-dm³ jar, place 100 g of each of the two pigments. Blend materials.

7.1.2.2 Blend thoroughly by agitation.

7.1.2.3 Add 1000 g of 4.76-mm (3/16-in.) steel balls; and place the closed jar on a roller for approximately 8 h at about 0.33 Hz (rps).

7.1.2.4 Remove the steel balls.

7.1.2.5 Scrape the densed dust out of the jar and pass through a No. 30 (600 μm) screen with the aid of a brush.^{6,9}

7.2 *Calibration Compounds*:

7.2.1 The formulas and the mixing and curing specifications for the five calibration compounds are given in Annex A1. A brief description of the five compounds in terms of rubber and black types is as follows:

Compound	Rubber	Black
A	Styrene-Butadiene-Rubber	Industry Reference-Black
A	Styrene-Butadiene	Industry Reference
B	Styrene-Butadiene-Rubber	Industry Reference-Black
B	Styrene-Butadiene	Industry Reference
C	Natural Rubber	Industry Reference-Black
C	Natural Rubber	Industry Reference
D	Styrene-Butadiene/ Polybutadiene —Blend	Industry Reference-Black
D	Styrene-Butadiene Polybutadiene Blend	Industry Reference
E	Styrene-Butadiene/ Polybutadiene —Blend	Intermediate-Surface-Abra- —sion-Furnace-Black

⁵ “Seneca Standard” Tripoli, Air Float Rose diatomaceous earth, manufactured by the American Tripoli Co., Seneca, MO, or Double Ground Rose, manufactured by the Harshaw Chemical Co., Cleveland, OH, has been found satisfactory.

⁶ Scanning Electron Microscopy (SEM) verification of cutting knife dimensions is available from BF Goodrich Research and Development, Brecksville, OH.

⁷ A preblend mixture

⁸ If you are aware of these pigments is commercially available through alternative suppliers, please provide this information to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the Ferry Machine Co., Kent, OH 44240, responsible technical committee¹, which you may attend.

⁹ Supporting data are available from ASTM Headquarters. Request RR- D11-1022.

¹⁰ “Seneca Standard” Tripoli, Air Float Rose, diatomaceous earth, manufactured by the American Tripoli Co., Seneca, MO, or Double Ground Rose, manufactured by the Harshaw Chemical Co., Cleveland, OH, has been found satisfactory.

¹¹ Alon-C undensd aluminum oxide, manufactured by the Cabot Corp., Boston, MA, has been found satisfactory.

¹² A preblended mixture of these pigments is commercially available through Ferry Industries, Inc., Stow, OH.

E

Styrene- Butadiene
Polybutadiene Blend

Intermediate Surface
Abrasion Furnace Black

7.2.2 The calibration

NOTE 2—The compounds given listed in 7.2.1, 9.1 and Annex A1 are new compounds (1982). The old calibration compounds of this method can no longer be prepared have been modified from the 1982 compounds, due to the unavailability of several materials: some ingredients. The new compounds will not give equivalent results and new “Nominal Indices” and “Permissible Ranges” (see Table 1) are being developed. During the interim period of development, the previous (1982) calibration, or reference compounds, which can be prepared, shall be prepared and used as described in this test method.

7.2.2 The calibration compounds given in Annex A1 are compounds developed in 2000. These compounds have been modified from the 1982 compounds (see Note 2). The 2000 calibration compounds were not intended to give equivalent index results to the old reference materials: calibration compounds described prior to 2000. The “Nominal Indices” of the new 2000 calibration compounds, however, have been developed based on the old reference compounds and on the qualification of in 5.1, and the tolerance ranges have been “Permissible Ranges” are being established based on the specification tolerances specifications for knife sharpness the cutting knives as explained delineated in 9.5. 6.1.7.

7.2.3 The five calibration compounds^{6,10} are used to check verify the operational status of the tester instrument. If within specification tolerances, experimental the “Permissible Ranges” described in Table 1, the subject materials are then tested and compared to the test results of the calibration compounds. In essence, once it has been determined that the tester is in- calibration, the calibration compounds act as reference compounds; and, where applicable, either of the two terms will be used.

8. Test Specimen

8.1 The standard test specimen shall be molded to give the dimensions shown in Fig. 3, and shall be regarded as the standard test specimen.

8.2 An alternative specimen may be used in which a disk of the test material, not less than 1.59 mm (0.062 in.) in thickness and of the same diameter as the standard specimen, is cemented to a previously used specimen, or one that has been buffed down to accommodate the thickness of the disk. The disk may be cut from a product or from a laboratory cured sheet.

8.3 For

8.3 When testing conveyor belt covers or similar items, a specimen may be prepared as described in 8.2 or by cutting cylindrical specimens through the belt and cementing these to buffed down standard specimens with either the top cover or bottom cover facing up, as desired: being the surface to be tested. Covers shall be at least 0.79 mm (0.032 in.) thick.

8.4 In all tests, a minimum of two separately cured test specimens per material shall be tested. However, if replicate determinations are required, the specimens they may be rebuffed and retested many times. Replicate determinations of providing the same test material are limited when thin disks are cemented to a standard specimen or when thin conveyor belt covers are tested. minimum specified thickness is maintained and noted in Section 12.

9. Calibration

9.1 The tester is checked calibration of the Pico Abrader shall be verified at least once every for every 30 specimens tested, or when using reformulated, remixed or recured calibration compounds.

NOTE 3—Industry Reference Black No. 5 was used in Compounds A through D when these nominal indices were determined; Compound E contains an ISAF Black. The most recent IRB allotment (No. 6; 1986) does not match its predecessor of 1982 in physical properties performance of vulcanized compounds. The nominal indices of Compounds A through D maintain their relative values with respect to one another with the new black allotment but Compound E is forced to a higher nominal index (148). See Section 8.

¹⁰ Cured specimens will be available from Corporate, Consulting Service and Instruments, Inc. (CCSi) and Akron Rubber Development Laboratory, Inc. (ARDL), both of Akron, OH.

TABLE 1 Calibration Compound (Reference Compound)
Specifications
~~**Note 1—Industry Reference Black No. 5 was used in Compounds A through D when these nominal indices were determined; Compound E contains an ISAF Black. The most recent IRB allotment (No. 6; 1986) does not match its predecessor in physical properties performance of vulcanized compounds. The nominal indices of Compounds A through D maintain their relative values with respect to one another with the new black allotment but Compound E is forced to a higher nominal index (148).**~~

	Compound Designation				
	A	B	C	D	E
Nominal index	76	86	106	113	128
Permissible range	69 to 83	81 to 91	95 to 117	105 to 121	116 to 140

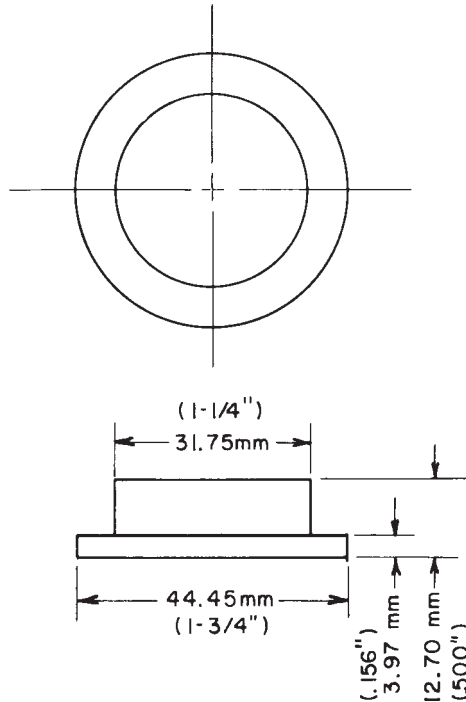


FIG. 3 Standard Test Specimen

9.1.1 The calibration shall be accomplished by testing each of the five calibration compounds enumerated in Table 1.⁹

9.1.1.1 The calibration shall be performed as described in Section 10, and under standard conditions. The results based on the calculation of 9.2 conditions described in Section 11.

9.1.1.2 The results shall be calculated as described in 9.2, 9.3, and 9.4, and shall be in within the acceptance limits of Permissible Range indicated in Table 1-1.

9.2 Calculate the volume loss for each calibration compound by subtracting the final mass from the initial mass and dividing the result by the density of the rubber:

$$L_x = \frac{M_i - M_f}{D} \quad (1)$$

where:

L_x = volume loss in cm^3 of calibration compound X,

M_i = initial mass in g,

M_f = final mass in g, and

D = density, in mg/m^3 .

9.3 Multiply the volume loss for each calibration compound by its nominal index, add these products together, and divide by 500 as follows:

$$\begin{aligned} \text{Standard volume loss} = & \\ & \frac{(L_A \times 76) + (L_B \times 86) + (L_C \times 106) \\ & + (L_D \times 113) + (L_E \times 128)}{5 \times 100} \end{aligned} \quad (2)$$

where: $L_A, L_B, \dots, L_C, L_D,$ and L_E = volume loss, in cm^3 , for Calibration Compounds A, B, ..., C, D and E.

9.4 Divide the standard volume loss resulting from the calculation of Eq 2 by the individual volume losses of each calibration compound to give the index value for each.

NOTE 1—Example: The each.

9.4.1 Example—the following volume losses, in cm^3 , were obtained:

$$\begin{aligned} L_A &= 0.0395 \\ L_B &= 0.0337 \\ L_C &= 0.0272 \\ L_D &= 0.0254 \\ L_E &= 0.0235 \end{aligned} \quad (3)$$

TABLE 2 Type 1—Precision

NOTE 1—

S_r = within laboratory standard deviation,
 r = repeatability (in measurement units),
 (r) = repeatability (in percent),
 S_r = between laboratory standard deviation,
 R = reproducibility (in measurement units), and
 (R) = reproducibility (in percent)

Com- pound	Test Level Average	Part A—Volume Loss, cm ³					
		Within Laboratories			Between Laboratories		
		S_r	r	(r)	S_r	R	(R)
E	0.017	0.0005	0.0014	8.32	0.0037	0.0105	61.6
C	0.017	0.0006	0.0017	10.0	0.0038	0.010	63.3
D	0.019	0.0006	0.0017	8.95	0.0036	0.0102	53.6
B	0.025	0.0012	0.0034	13.6	0.0069	0.0195	78.1
A	0.036	0.0011	0.0031	8.61	0.0043	0.0122	33.8
Part B—Abrasion Index							
A	71.4	6.37	18.0	25.3	7.61	21.5	30.1
B	96.9	5.28	14.9	15.4	4.97	14.1	14.6
C	114.2	4.26	12.1	10.6
D	119.3	6.70	19.0	15.9	5.03	14.2	11.9
E	140.7	6.70	19.0	13.5	14.1	39.9	28.4

$$\begin{aligned}
 \text{Standard volume loss} &= \\
 &= \frac{(0.0395 \times 76) + (0.0337 \times 86) + (0.0272 \times 106) + (0.0254 \times 113) + (0.0235 \times 128)}{5 \times 100} \\
 &= 0.0293 \tag{4}
 \end{aligned}$$

$$\text{Index for calibration compound A} = (0.0293/0.0395) \times 100 = 74$$

$$\text{Index for calibration compound B} = (0.0293/0.0337) \times 100 = 87$$

$$\text{Index for calibration compound C} = (0.0293/0.0272) \times 100 = 108$$

$$\text{Index for calibration compound D} = (0.0293/0.0254) \times 100 = 115$$

$$\text{Index for calibration compound E} = (0.0293/0.0235) \times 100 = 125$$

9.5 To function properly,

9.5 During routine use, the cutting knives must have a “flat” will become dull, affecting test determinations. The cutting knives may be routinely examined (see 6.1.7.3) so that they have, at the apex of 10 to 20 μm on the abrasion edge. To produce this angle formed by the two bevels, a blunted edge, or “flat,” knives of with a width no greater than 20 μm.

9.5.1 When, during use, the geometry illustrated in Fig. 4 can cutting knives are routinely examined, they shall be dulled removed from service if the blunted edge, or “flat,” with a diamond dust to give a “flat” width greater than 20 μm.

9.5.2 When the effectiveness of the cutting knives is determined by reliance upon their comparative performance to 12 μm. Normally, with new or resharpened knives, the five calibration compounds will fall within enumerated in Table 1, they shall be removed from service when a test determination falls outside of the acceptance limits. If one or more “permissible range” for any of the calibration compounds fail to do so, it indicates that the particular compound(s) is out of specification given in one respect or another and it should be remixed, recured, and retested. After continued use, Table 1.

9.6 In the knives will wear so event that the “flat” becomes too wide, that is, wider than 20 μm. When this occurs, Calibration Compounds C and E will become equivalent to Compound D and Compounds C and E will also fail to meet their acceptance limits. The cutting knives should be returned are found to the supplier for resharpening (Note 2).

NOTE 2—Scanning electron microscopy may be used to identify within the width of the “flats.” This procedure will eliminate any false measurements due to reflected light that may be encountered with a normal light microscope.

9.6 When previously enumerated dimensional specifications and tests performed on the five calibration compounds are do not fall within their tolerance ranges, the tester is respective “Permissible Range” enumerated in calibration. The frequency for checking Table 1, the calibration compounds shall be remixed, recured and re-tested.

9.7 When the procedure described in 9.6 fails to yield the desired results within a laboratory, between laboratories, between customer and supplier, or for the purposes of ry-30eferee testing, the following shall occur:

9.7.1 Prepared test specimens, obtained from third party commercial sources (see Note 2), shall be tested following the procedures in Sections 9 and 10 using new or resharpened cutting knives adhering to the specifications in 6.1.7 (see Fig. 2).

10. Procedure

10.1 Although it is common practice in ASTM standards to present a section on

~~10.1 The calibration before a section on procedure, the order is reversed because the calibration of test necessitates the knowledge of operation. Both procedure procedures and calibration frequency described in Section 9 are required to be performed prior to conducting tests on experimental subject materials.~~

~~10.2 *Preparing the Test Specimen by:*~~

~~10.2.1 Buffing the surface of the specimen on a surface grinder as specified described in 6.2.~~

~~10.2.1.1 Set the abrasive wheel for an abraded depth of 0.13 mm (0.005 in.) to remove the mold skin.~~

~~10.2.1.2 Follow with a finishing depth of 0.025 mm (0.001 in.). W, then without changing the micrometer setting, give the surface of the specimen a finish grind.~~

~~10.2.1.3 The dwell time per pass under the abrasive wheel should be approximately 1 s.~~

~~10.2.1.4 Remove all loose rubber particles and debris from the buffed specimen.~~

~~10.2.2 When successive tests are made on a previously tested specimen (see 8.2), buff off the abrasion pattern produced on the specimen as given in 10.2.~~

~~10.3 Brush the buffed specimen to remove all loose rubber particles.~~

~~10.4 Different volume losses will be obtained depending on the time specimens are allowed to equilibrate after buffing. To obtain consistent results, specimens, after buffing, should be allowed to condition for 24 h in an atmosphere having a temperature of $23 \pm 2^\circ\text{C}$ ($73.4 \pm 3.6^\circ\text{F}$) and a relative humidity of $50 \pm 5\%$ in accordance with Practice D 1349.~~

~~10.5 Determine the mass of the specimen on a balance, recording results to the nearest 0.0001 g.~~

~~10.6 Close the power switch for the motor control unit. Let the thymatrol tubes warm up for two minutes before starting the motor. After the motor warms up, it can be started, stopped, and reversed at will (Note 3).~~

~~NOTE 3—A direct-current driven motor with SCR control may be used. If so, 10.6 can be disregarded.~~

~~10.2.3 Different volume losses will be obtained depending on the time specimens are allowed to equilibrate, that is, to remain in the atmospheric condition specified, after buffing, for specimen conditioning (see Section 11). NOTE 4—A tachometer may be mounted permanently to~~

~~10.3 *Preparing the apparatus to give a direct indication of rotational frequency.*~~

~~10.8 The force on Apparatus:~~

~~10.3.1 Close the knives power switch for standard conditions shall be 44 N (Note 5). The tare of the shaft, chuck, knife holder, knives, and box is stamped on each tester. Add a sufficient amount of lead or steel motor control unit. Allow the thymatrol tubes to attain 44 N.~~

~~NOTE 5—The warm up for two minutes before starting the motor. After the motor warms up, it can be started, stopped and reversed as needed.~~

~~10.3.2 Direct current drive motors with SCR controls may be used, in which case 10.3.1 can be disregarded.~~

~~10.3.3 The rotational frequency of the turntable shall be 1.00 ± 0.03 Hz (rps) with a force of 44 N is a standard condition. (rps).~~

~~10.3.4 The conditions described in 10.3.3 and 10.3.6 shall be regarded as standard. If varying severity tests are desired, it is recommended that the following alternative conditions be employed:~~

	Rotational Frequency, Hz (rps)	Force, N
Low severity	0.50	24.5
High severity	1.83	88.2
Low severity	0.50	24.50
High severity	1.83	88.20

~~For each condition, the~~

~~10.3.5 The number of revolutions remains the same, that is, a total of 80.~~

~~10.9 Clamp 80 (see 10.4.10).~~

~~10.3.6 The force on the knives shall be $44 \text{ N} \pm 0.44 \text{ N}$. The tare of the shaft, chuck, knife holder, knives, and box is stamped on each tester, add the appropriate mass to attain the specified force.~~

~~10.3.7 These conditions may be modified if agreed upon between laboratories or between supplier and user and are so reported (see Section 12).~~

~~10.4 *Performing the Test:*~~

~~10.4.1 Tests shall be conducted in accordance with the conditions in Section 11.~~

~~10.4.2 Determine the mass of the specimen on a balance, recording results to the nearest 0.0001 g.~~

~~10.4.3 Mount the specimen onto the turntable.~~

~~10.4.4 Start the duster and adjust so that an even flow (5 mg/s) of 5 mg/s of dusting powder is established.~~

~~10.4.5 Push the forward button, and at button.~~

~~10.4.6 Lower the first click of the counter, lower the knives gently onto the specimen within the first quarter revolution following revolution.~~

~~10.4.7 After the click. On the twentieth click, revolution, stop the tester within one quarter of a revolution.~~

~~10.4.8 Stop the duster and lift the knives from the specimen. Vacuum the dusting powder and the abraded rubber particles from the specimen. Restart the duster and repeat these operations, reversing the direction of rotation for three more increments of 20 revolutions, to a total of 80 revolutions, 40 revolutions in each direction.~~

~~10.11~~ A specimen.

~~10.4.8.1~~ A cylindrical receptacle may be placed around the test specimen and filled with a sufficient supply of dusting powder so that the knives do not have to be lifted upon change of rotational direction.

~~10.4.8.2~~ Volume losses obtained by this test method ~~will~~ may not be as great, but as long as all test specimens and calibration compounds are tested similarly, similar index results will be attained.

~~10.4.8.3~~ If this procedure is ~~is~~ employed, it shall be reported (see Section 12).

~~10.4.9~~ Vacuum ~~the~~ dusting powder and the abraded rubber particles from the specimen.

~~10.4.10~~ Repeat operations 10.4.4 through 10.4.8, reversing the direction of rotation of the turntable each time, for three more increments of 20 revolutions (test cycle), for a total of 80 ~~revolutions~~ automans (one test), 40 revolutions in each direction.

~~10.4.10.1~~ Electronic controls, designed to automate operations 10.4.10.2 through 10.4.10.7, are permissible.

~~10.4.12~~ Remove the specimen, ~~brush away~~ remove all loose powder and abraded rubber with a stiff brush.

~~10.5~~ Calculation ~~d:~~

~~10.5.1~~ Determine the mass of the specimen on a balance, recording ~~the result~~ results to the nearest 0.0001 g.

~~10.5.2~~ Calculate the volume loss from the mass loss and density of the tested material. As done in 9.2, 9.3, and 9.4, express the result in an index number obtained by dividing the standard volume loss of the reference compounds, tested in the same series, by the volume loss of the tested material and multiplying the quotient by 100.

~~10.5.3~~ Calculate the index of the tested material against any one of the reference compounds as described in 10.5.2. This method is particularly useful when employing non-standard severity conditions.

11. Calculation

~~11.1~~ Calculate Laboratory Atmosphere, Instrument and Test Specimen Conditioning

~~11.1~~ Tests shall be conducted in the ~~volume loss from~~ standard laboratory atmosphere, as defined in Practice D 618.

~~11.2~~ The instrument shall be maintained in the standard laboratory atmosphere, as defined in D 618, for 24 h prior to performing calibration or tests.

~~11.3~~ The specimen shall be conditioned in accordance with Condition 24/23/50 as described in D 618, and ~~density of~~ within the ~~tested material~~. As ~~done~~ tolerances specified in 9.2, 9.3, D 618, Section 3, and 9.4, ~~express the result~~ stated as an index Condition 24/23/50, T-23-50.

where:

~~24~~ ≡ the number obtained by dividing the standard volume loss of the reference compounds, tested in the same series, by the volume loss hours of the tested material and multiplying the quotient by 100.

~~11.2~~ Calculate the index of the conditioning,

~~23~~ ≡ the temperature in degrees Celsius, $\pm 2^{\circ}\text{C}$ ($73.4 \pm 3.6^{\circ}\text{F}$), and

~~50~~ ≡ the relative humidity in %, ± 5 %.

~~and tested material against any one of~~ under the reference compounds as same conditions, described in D 618 as T-23-50,

where:

~~T~~ ≡ test conditions of $23^{\circ} \pm 2^{\circ}\text{C}$, and 50 % relative humidity ± 5 %.

~~11.4~~ No conclusive evaluation has been made on Pico Abrasion calibrations or tests under conditions other than those stated in 11.1. ~~F~~ Conditioning or testing at temperatures other than the above may cause variations in calibration or test method is particularly useful when employing nonstandard severity conditions. ~~results.~~

~~11.5~~ These procedures may be modified if agreed upon between laboratories or between supplier and user and are in accordance with alternative procedures identified in D 618.

12. Report

12.1 The report shall include the following:

~~12.1.1~~ Identification

~~12.1.1~~ Type of specimen, standard or alternative, as described in Section 8,

~~12.1.2~~ Identification of the tested specimens,

~~12.1.23~~ Cure time and temperature,

~~12.1.3~~ Density,

~~12.1.4~~ Force on knives used, in N, as described in 10.3.4 or 10.3.6,

~~12.1.5~~ Rotational frequency used, in MHz, as described in 10.3.4 or 10.3.6,

~~12.1.6~~ Type of dusting, as described in 10.4.8.1, Standard Feed or Cylinder Contained,

~~12.1.7~~ Initial specimen mass, M_i , as calculated in 9.2, Eq 1,

~~12.1.78~~ Final specimen mass, M_f , as calculated in 9.2, Eq 1,

~~12.1.89~~ Specimen mass loss, $M_i - M_f$, as calculated in 9.2, Eq 1,

~~12.1.10~~ Density, D , as calculated in 9.2, Eq 1,

12.1.11 Specimen volume loss, L_v , and calculated in 9.2, Eq 1,

12.1.10² Abrasion index with the test method used to calculate the index as described in ~~H.1~~ 10.5.2 or ~~H.1~~ 10.5.3 (see 9.3, Eq 2, and 9.4, Example 1).

13. Precision and Bias ¹¹

13.1 These precision statements have been prepared in accordance with Practice D 4483. Please refer to this practice for terminology and other testing and statistical concept explanations.

13.2 *Type I*—Precision data have been compiled which excludes the compounding variation among laboratories (see Table 2).

13.3 The statements were developed from interlaboratory testing of the five new ϵ calibration ϵ compounds (A–E). These five compounds were mixed in a B Banbury internal mixer with curatives added on a mill. All specimens of the same compound were cured from the same mix.

13.4 The precision statements are based on the testing of five samples by six laboratories on two days.

13.5 A test result is defined to be the average of two separately cured specimens.

13.6 Precision statements were prepared for both volume loss and abrasion index. Volume loss is the measure of the physical differences among the interlaboratory testers and index is a measure of the test method in general.

13.7 The precision of this test method may be expressed in the format of the following statements that use an appropriate value of r , R , (r), or (R), that is, that value to be used in decisions about test results (obtained with this test method). The *appropriate value* is that value of r or R associated with a mean level in the precision table closest to the mean level under consideration (at any given time, for any given material) in routine testing operations.

13.8 *Repeatability*— The repeatability r , of this test method has been established as the *appropriate value* tabulated in the precision tables. Two single test results, obtained under normal test method procedures, that differ by more than this tabulated r (for any given level) must be considered as derived from different or non-identical sample populations.

13.9 *Reproducibility*— The reproducibility R , of this test method has been established as the *appropriate value* tabulated in the precision tables. Two single test results obtained in two different laboratories, under normal test method procedures, that differ by more than the tabulated R (for any given level) must be considered to have come from different or non-identical sample populations.

13.10 Repeatability and reproducibility expressed as a percentage of the mean level, (r) and (R), have equivalent application statements as 13.8 and 13.9 for r and R . For the (r) and (R) statements, the difference in the two single test results is expressed as a percentage of the arithmetic mean of the two test results.

13.11 *Bias*—In test method terminology, bias is the difference between an average test value and the reference (or true) test property value. Reference values do not exist for this test method since the value (of the test property) is exclusively defined by the test method. Bias, therefore, cannot be determined.

14. Keywords

~~14.1~~ knife

14.1 abrasion; abrasion resistance; knife abrasion; Pico abrader; Pico abraser; rubber articles; rubber products

¹¹ This test method is under the jurisdiction of ASTM Committee D11 on Rubber and is the direct responsibility of Subcommittee D11.15 on Degradation Tests . Current edition approved June 10, 2002. Published September 2002. Originally published as D 2228 – 63 T. Last previous edition D 2228 – 88 (2001).

ANNEX

(Mandatory Information)

A1. FORMULAS AND MIXING AND CURING SPECIFICATIONS FOR THE CALIBRATION COMPOUNDS

A1.1 Calibration Compounds

A1.1.1 The formulas for the calibration compounds are given in Table A1.1.

A1.2 Methods of Mixing

A1.2.1 The compounds are mixed in a B Banbury internal mixer in accordance with Practice D 3182. Initial temperature of the mixing chamber is 50°C (120°F). Batch size is 70 % of total chamber volume capacity. The curatives are added on a mixing mill having rolls between 250 and 258-mm (9.8 to 10.2-in.) outside diameter and an operating temperature of 65°C (150°F).

NOTE A1.1—The standard mill has rolls between 150 and 155-mm (5.9 to 6.1-in.) outside diameter. If this mill is used to add curatives, the batch may be divided into three equal portions. The mixing cycle may have to be adjusted to obtain comparable results.

A1.2.2 The mixing cycles for the calibration compounds are given in Table A1.2. Mixing is accomplished by following the time

TABLE A1.1 Formulas for Calibration Compounds

Ingredient	NBS Industry Trade Name or Reference Supplier	Designation				
		A	B	C	D	E
Natural rubber	385			100.0		
Natural rubber ^A	SMR L	0.00	0.00	100.0	0.00	0.00
GB-441					68.75	68.75
Polybutadiene ^B	TAKTENE 1220	0.00	0.00	0.00	50.00	50.00
SBR 1502			100.0			
SBR 1502 ^C	SBR 1502	0.00	100.00	0.00	0.00	0.00
SBR 1712		137.5			68.75	68.75
SBR 1712 ^C	SBR 1712	137.50	0.00	0.00	68.75	68.75
Stearic acid	372	-1.5	-1.5	-2.0	-1.5	-1.5
Stearic acid ^D	Stearic Acid IRM 021a	1.5	1.5	2.0	1.5	1.5
Zinc oxide	370	-5.0	-5.0	-5.0	-5.0	-5.0
Zinc oxide ^E	Zinc Oxide IRM 91a	5.0	5.0	5.0	5.0	5.0
IRB ^A		-60.0	-40.0	-45.0	80.0	
IRB ^F	IRB #7	60.0	40.0	45.0	80.00	0.00
ISAF black ^B						80.0
ISAF black ^G	N220	0.00	0.00	0.00	0.00	80.00
Process oil ^C		-5.0	-5.0	-5.0	-8.75	-8.75
Process oil ^{E,H}	SUNDEX 8125	5.0	5.0	5.0	8.75	8.75
Dimethyl butylphenyl phenylenediamine						
Dimethyl butylphenyl phenylenediamine	SANTOFLEX 13	1.50	1.50	1.00	1.50	1.50
Trimethyldihydro- quinoline		-1.5	-1.5	-1.0	-1.5	-1.5
Trimethyldihydro- quinoline ^I	FLECTOL H	1.5	1.5	1.0	1.5	1.5
TBBS ^D		-1.5	-1.5	-1.5	-1.5	-1.5
TBBS ^{J,K}	TBBS IRM 003	1.5	1.5	1.5	1.5	1.5
Sulfur	384	-1.0	-1.0	-0.6	-1.2	-1.2
Sulfur ^D	Sulfur IRM 031a	1.0	1.0	0.6	1.2	1.2
Specific gravity	374	-2.0	-2.0	-2.5	-2.0	-2.0
	Totals	2.0	2.0	2.5	2.0	2.0
		-1.13	-1.12	-1.13	-1.16	-1.16

^AIndustry Reference Black is available in 22.68 kg (50 lb) bags from ea H.A. Astlett, Toron-bto, Ontaek-suppliersio.

^BN234, or its equi Available from Bayer Canada, Toronto, Ontario.

^CSBR 1502 andex SBR 171260T, e are available from synthetic rubber suppliers in various quantities.

^D Available from Akron Rubber Development Laboratory, Akron, OH.

^E Available from R.E. Carroll, Akron, OH.

^F Industry Reference Black #7 is available in 22.68 kg (50 lb) bags from carbon black suppliers.

^G This replaces N234, which is no longer commercially available.

^H This replaces Sundex 7260T, which is no longer commercially available.

^I Available from Harwick, Akron, OH.

^J N- tert-butyl-2-benzothiazole sulfenamide.

^K Available from Akrochem, Akron, OH.

TABLE A1.2 Mixing Cycles for Calibration Compounds

Step	Compound, min				
	A	B	C	D	E
<i>B Banbury:</i>					
Add rubber	0:00	0:00	0:00	0:00	0:00
Add fillers	0:30	0:30	0:30	0:30	0:30
Ram sweep	2:00	1:45	2:10	2:00	2:00
Add oil	2:30	2:10	2:55	2:25	2:30
Ram scrape	3:10	3:00	3:50	3:15	3:15
Dump	4:15	3:50	5:05	4:00	4:15
Probe					
temperature at dump, °C, (°F)	143 (290)	152 (305)	160 (320)	138 (280)	138 (280)
<i>Mill:</i>					
Band material	0:00	0:00	0:00	0:00	0:00
Add curatives	0:30	0:40	0:45	0:40	0:40
End pass	2:30	2:15	3:00	2:15	2:00
Take off	6:15	6:30	6:15	5:30	5:30

specifications. No provisions are made for preconditioning of the carbon black.

A1.2.3 Sheet the compound off the mill at an approximate thickness of 2.1 mm (0.08 in.) and cool on a flat dry clean metal surface.

A1.2.4 Die cut disks of the required diameter and stack the disks to the required height.

A1.3 Recommended Cures

A1.3.1 Cures for the calibration compounds, in the form of the molded specimens required for the test, are given in Table A1.3.

A1.3.2 The size of the molded specimen is such that a lag time or incubation time of approximately five minutes is involved in the cure. It is recommended that cures of experimental materials be increased by this amount over the cure that is established for sheets approximately 2 mm (0.08 in.) thick.

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TABLE A1.3 Recommended Cures for Calibration Compounds

Compound	Time, min	Temperature, ° C (°F)
A	65	150 (302)
B	65	150 (302)
C	60	140 (284)
D	65	150 (302)
E	65	150 (302)