

Standard Test Method for Absorption Coefficient of Ethylene Polymer Material Pigmented with Carbon Black¹

This standard is issued under the fixed designation D 3349; 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. Scope

1.1 This test method measures the amount of light transmitted through a film of carbon black pigmented ethylene polymer.

1.2 Taking into account the amount of light and film thickness, an absorption coefficient is calculated.

1.3 Whenever two sets of values are presented, in different units, the values in the first set are the standard, while those in parentheses are for information only.

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:

E 60 Practice for Molecular Absorption Spectrometric Methods for Chemical Analysis of Metals, Ores, and Related Materials²

D 1248 Specification for Polyethylene Plastics Extrusion Materials for Wire and Cable³

3. Summary of Test Method

3.1 The transmission of light of a specified wavelength through thin ethylene polymer film is measured. The result is used to calculate an absorption coefficient that is a reciprocal function of transmission after correction for thickness.

4. Significance and Use

4.1 The absorption coefficient of polyolefin polymer pigmented with carbon black is useful to judge the degree and uniformity of dispersion of the pigment, and the adequacy of the quantitative level of pigment addition. These factors are used to predict the performance of the polymer material in

response to prolonged exposure to ultraviolet light as evidenced by minimal changes in specific properties.

NOTE 1—This test method was developed to evaluate ethylene polymer materials pigmented with small particle size carbon blacks suitable for UV protection. It is not known how accurate and reproducible the test would be with larger particle size blacks.

5. Apparatus

5.1 *Spectrophotometer*—An instrument in accordance with Practice E 60 is required.

NOTE 2—Consult the spectrophotometer's operation manual or consult the manufacturer of the instrument for the calibration procedure.

5.2 *Mold*—A mold in accordance with Figs. 1 and 2. Make the mold of Ketos steel (or equivalent) hardened to Rockwell C45, with mold surfaces chromium plated to 0.005-mm (0.0002-in.) minimum thickness.

5.3 *Specimen Holder*—Cut two concentric rings from phenolic-paper laminate or other suitable material to the dimensions shown in Fig. 3. These rings should make a snug slip-fit one within the other.

5.4 *Reference Material*—A reference material is required having an absorbance value of 1.0 to 1.2 at 375 nm as measured by the instrument used for testing.

5.4.1 *Neutral Density Filter*,⁴ M-type carbon 0.85 density, laminated between cellulose acetate sheets has been found satisfactory. Since this material is furnished as a rigid sheet, it cannot be mounted as described in Section 7. Instead, a special mount must be prepared by cutting Ring 1 of Fig. 3 in half around its circumference to give two rings about 2 mm (0.08 in.) thick. Cut a 22-mm (0.87-in.) circle from the laminated filter, sandwich this between the two 2-mm rings, and slide the outer ring (Ring 2 of Fig. 3) over this composite.

5.4.2 An alternative reference material can be prepared by any user of this test method. An alternative reference material consisting of a highly uniform sample of ASTM Type I polyethylene (see Specification D 1248) containing about 1 % of a 20-nm carbon black (either channel or furnace type) may be prepared on a laboratory mill, and a uniform film of this material pressed out and mounted as in Section 8 to meet the

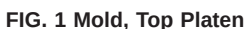
¹ This test method is under the jurisdiction of ASTM Committee D-9 on Electrical and Electronic Insulating Materials and is the direct responsibility of Subcommittee D09.18 on Solid Insulations, Non-Metallic Shieldings, and Coverings for Electrical and Telecommunications Wires and Cables.

Current edition approved March 10, 1999. Published June 1999. Originally published as D 3349 – 74. Last previous edition D 3349 – 93.

² Annual Book of ASTM Standards, Vol 03.05.

³ Annual Book of ASTM Standards, Vol 08.01.

⁴ The Kodak neutral density filter, No. 5 is available from the Eastman Kodak Co.



6. Calibration and Standardization

6.2 Insert the mounted reference material in the spectrophotometer and place it into the light beam. Read and record the absorbance value using the conditions that bring the value nearest to zero on the absorbance scale. With the reference material still in the beam, readjust the slit width to bring the reading back to zero. Recheck the infinity value.

7.1 No special conditioning is required.

8.1 Prepare three test specimens from each lot of material by hot-pressing the ethylene polymer at a suitable temperature between highly polished plates, such as those shown in Figs. 1 and 2, using a charge sufficient to yield a specimen 40 to 50 mm in diameter and approximately 0.01 mm in thickness.

9. Procedure

9.2 Close the shutter and recheck the reference material. Rotate the specimen to bring the next 120° mark to the top and

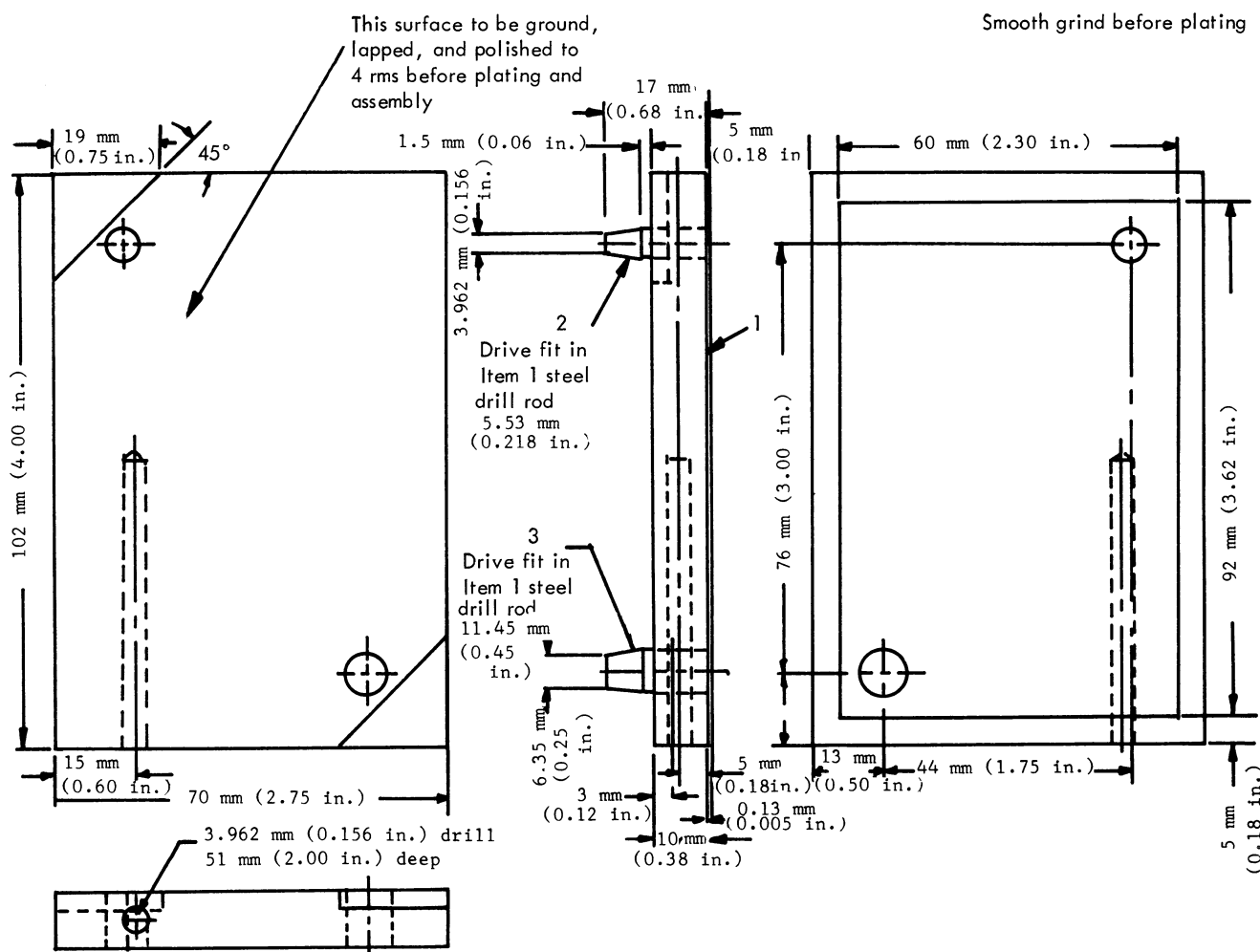


FIG. 2 Mold, Bottom Platen

obtain the absorbance value v_2 . Repeat the procedure with the third 120° mark at the top to obtain the value v_3 . Average the values v_1 , v_2 , and v_3 to obtain the average absorbance $\bar{v}$. If any value of v differs from $\bar{v}$ by more than 10 %, discard the specimen and replace it.

9.3 Remove the film from the specimen mounting rings by cutting it carefully along the boundary between the two rings and weigh it to the nearest 0.0001 g. The No. 11 Bard-Parker scalpel blade⁵ in a suitable handle has been found very effective for the cutting operation. Measure the diameter and calculate the area of the specimen. Calculate average specimen thickness as follows:

$$t = (10^{-2} \cdot W) / (D \cdot A) \quad (1)$$

where:

t = thickness, m,
 W = weight, g,
 D = density, g/cm³, and
 A = area, cm².

NOTE 3—The nominal density value for black low density ethylene polymer is 0.932 g/cm³.

⁵ Available from most laboratory supply houses.

9.4 Repeat 9.1-9.3 for the second and third specimens.

10. Calculation

10.1 Calculate the absorption coefficient of each specimen, in turn, as follows:

$$a = (10^{-3} \cdot 2.303 \bar{v}) / (t) \quad (2)$$

where:

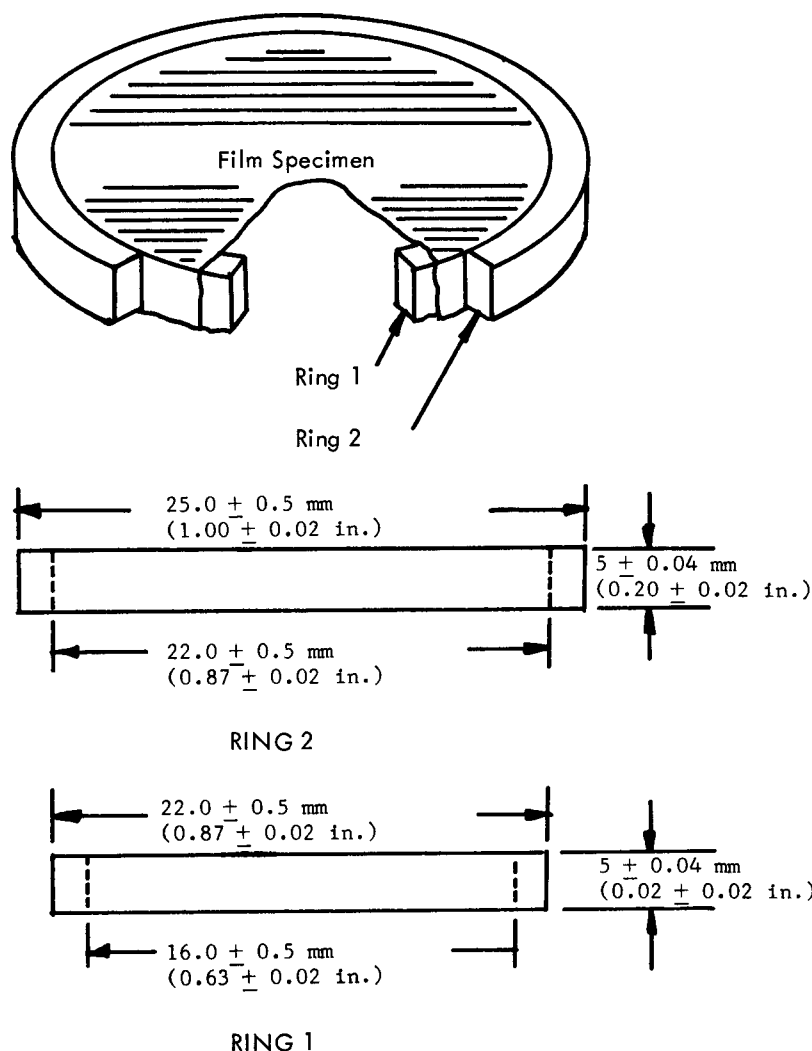
a = absorption coefficient, 1000(absorbance/metre), at 375 nm,
 $\bar{v}$ = average absorbance as found in 9.2, and
 t = thickness, m, as found in 9.3.

10.2 Average the three values, a , for the three specimens to obtain the value for the sample.

11. Report

11.1 Report the following:

- 11.1.1 Complete identification of the material tested,
- 11.1.2 Nominal density of the material tested,
- 11.1.3 Calculated absorption coefficient of each specimen tested,
- 11.1.4 Average absorption coefficient of three specimens tested, which is the value for the sample, and
- 11.1.5 Type and model of spectrophotometer used.



NOTE 1—Adjustments to the inside diameter of Ring 2 or outside diameter of Ring 1 may be necessary to obtain a snug fit of pressed film.

FIG. 3 Specimen Mounting Rings

12. Precision and Bias

12.1 Precision:

12.1.1 In the early 1970's, a round-robin study was conducted by five laboratories using one sample of carbon black-pigmented ethylene polymer and the same make and model of spectrophotometer. Each laboratory molded from two to five test specimens from the sample and determined the absorption coefficient of each. The following statistics, each expressed in the units of 1000 (absorbance/metre), were derived from the data submitted for this one sample.

Mean	= 443
S_r	= 24
S_R	= 43

12.1.2 Interpretation of the data for the intervals S_r and S_R given above are as follows.

12.1.2.1 *Repeatability*—In comparing two averages (of three specimens each) for the sample obtained in one laboratory by one operator on the same day, the averages should be judged as not equivalent if they differ by more than S_r .

12.1.2.2 *Reproducibility*—In comparing two averages (of three specimens each) for the sample obtained in different

laboratories by different operators on different days, the averages should be judged as not equivalent if they differ by more than S_R .

12.1.2.3 The judgments in 12.1.2.1 and 12.1.2.2 will be correct in approximately 95 % of such comparisons.

12.2 Bias:

12.2.1 In the round-robin study, it was found that different makes and models of spectrophotometer can produce systematically different results for the average absorbance of a specimen, with the maximum consistent difference being 6 % relative for the four types of instrument used in the study.

12.2.2 In critical applications, such as conformance of a material to a specification, it may be necessary for the buyer and seller to agree upon the instrument that is to be used or to establish a correlation between the two types of instruments in use.

13. Keywords

13.1 absorption coefficient; carbon black; ethylene polymer

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