

Designation: D1442 - 06 (Reapproved 2012)

Standard Test Method for Maturity of Cotton Fibers (Sodium Hydroxide Swelling and Polarized Light Procedures)¹

This standard is issued under the fixed designation D1442; 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 covers the determination of the percentage of mature fibers in a sample of loose, chemically untreated cotton fibers, whether taken before processing or unravelled from a textile product.

1.2 This test method gives two optional procedures for determining maturity, as follows:

1.2.1 Procedure 1-Sodium Hydroxide Swelling.

1.2.2 Procedure 2-Polarized Light.

NOTE 1—For other test methods for the determination of maturity of cotton fibers refer to Test Methods D1464 and D2480.

1.3 The values stated in SI units are to be regarded as standard. No other units of measure are included in this 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 and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

- 2.1 ASTM Standards:²
- D123 Terminology Relating to Textiles
- D1440 Test Method for Length and Length Distribution of Cotton Fibers (Array Method)
- D1447 Test Method for Length and Length Uniformity of Cotton Fibers by Photoelectric Measurement

D1464 Practice for Differential Dyeing Behavior of Cotton

D1776 Practice for Conditioning and Testing Textiles

D2480 Test Method for Maturity Index and Linear Density of Cotton Fibers by the Causticaire Method (Withdrawn $1992)^3$

D7139 Terminology for Cotton Fibers

3. Terminology

3.1 For all terminology relating to D13.11, Cotton Fibers, refer to Terminology D7139.

3.1.1 The following terms are relevant to this standard: cotton fiber maturity, immature fibers, in testing with sodium hydroxide solutions (See Fig. 1 and Fig. 2), immature fibers, observed under polarized light, lumen, mature fibers, in testing with sodium hydroxide solutions (see Fig. 3), mature fibers, observed under polarized light (see Table 1), micronaire reading, test specimen, in cotton maturity test.

3.2 For all other terminology related to textiles, refer to Terminology D123.

4. Summary of Test Method

4.1 Fibers are laid parallel on a microscope slide, covered with a cover glass, treated with a mounting medium, and the magnified images are then classified as mature or immature fibers.

4.2 The method offers two procedures for classifying the fibers as mature or immature:

4.2.1 *Procedure 1, Sodium Hydroxide Swelling,* which uses an 18% solution of sodium hydroxide as the mounting medium and a laboratory microscope for viewing the fibers at a magnification of 400×.

4.2.2 *Procedure 2, Polarized Light,* which uses clear mineral oil as the mounting medium and requires a polarizing microscope giving a magnification of 100×. Fibers are classified according to their second-order interference colors, using a first-order (or full wave) retardation plate (Table 1).

5. Significance and Use

5.1 Information regarding the percentage of immature fibers is desirable because immature fibers: (1) break easily during

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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.

 $^{^{3}\,\}mathrm{The}$ last approved version of this historical standard is referenced on www.astm.org.

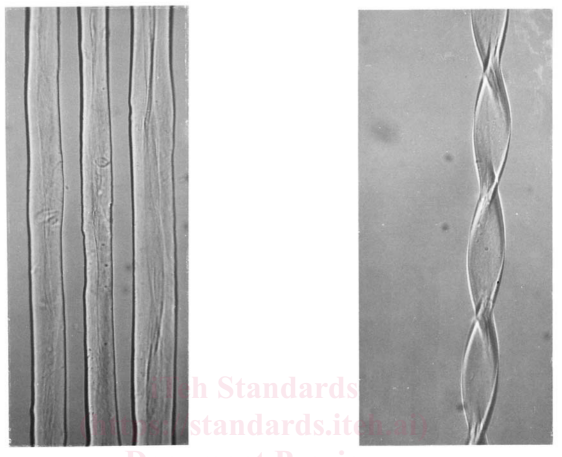


FIG. 1 Mature Fiber CUM ent Preview FIG. 2 Immature Fiber (Type A)

processing; (2) have a tendency to form neps; (3) have a tendency to become entangled around particles of trash and leaf, thus making cleaning more difficult and increasing the amount of fiber removed with foreign matter; (4) adversely affect yarn and fabric appearance; and (5) may appear differently after dyeing.

5.2 Maturity has a high positive correlation with linear density, but genetic differences and differences in wall thickness caused by plant diseases, soil, and water conditions during the growing season interfere with this relationship. Thus two cottons having the same linear density, or having the same average wall thickness as indicated by air-flow instruments, may vary greatly in maturity, that is, a cotton having extremely variable wall thickness may contain more immature fibers than another cotton of the same Micronaire reading composed of fibers having very uniform wall thickness.

5.3 The Sodium Hydroxide Swelling (Procedure 1) has been used in judging other maturity tests such as the Causticaire and the differential dye methods, in which the individual fibers are not examined.

5.4 Finer distinctions between different degrees of fiber wall development can be made with the Polarized Light procedure than with the Sodium Hydroxide Swelling procedure. The Polarized Light procedure gives a view of the fiber in its natural state so that fibrillar structure, striations, reversals, etc.,

are clearly visible as are growth abnormalities and variations in wall thickness. This method may be preferred by botanists, geneticists, and plant physiologists, while the Sodium Hydroxide Swelling procedure may be preferred for routine testing of large numbers of samples. Technicians are more easily trained for the latter method. Arbitrary classification as to maturity must be made with both methods.

5.5 This method is not considered satisfactory for acceptance testing because between laboratory precision can be poor. In some cases the purchaser and seller may have to test a commercial shipment of one or more specific material by an appropriate method even though the method has not been recommended for acceptance testing of commercial shipments. In such a case, if there are differences of practical significance between reported test results for two laboratories (or more), comparative tests should be performed to determine if there is a statistical bias between them, using competent statistical assistance. As a minimum, ensure the test samples to be used are as homogeneous as possible, are drawn from the material from which the disparate test result were obtained, and randomly assigned in equal numbers to each laboratory for testing. The test results from the two laboratories should be compared using statistical test for unpaired data, at a probability level chosen prior to the testing series. If a bias is found,

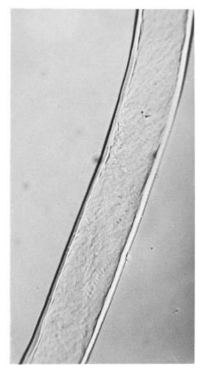


FIG. 3 Immature Fiber (Type B)

TABLE 1 Colors of Cotton Fit	ers Viewed with	Polarized Light ^A
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	Without Retardation Plate	With Retardation Plate	
Fiber Classification	First Order	Additive Colors	Subtractive
			Colors
		Second Order	First Order
Mature	light yellow	yellow	light yellow
	white	green	yellow
Immature	gray-blue	blue	orange-yellow
	gray	purple 🛛 🛆	orange D1442

^A Classified according to Mary Anna Grimes, "Polarized Light Preferred for Maturity Tests," *Textile World*, February, 1945.

either its cause must be found and corrected, or future test results for that material must be adjusted in consideration of the known bias.

6. Apparatus and Reagents

6.1 Procedure 1:

6.1.1 *Microscope or Microprojector*, which will give a magnification of approximately 400×, equipped with a mechanical stage, microscope lamp, and viewing aid such as a Euscope or projection screen.

6.1.2 Metal Comb, rake-type.

6.1.3 *Microscope Slides*, 2 by 3 in. (50 by 75 mm), and appropriate cover glasses.

6.1.4 Forceps, Dissecting Needles, and Tweezers.

6.1.5 *Multiple Counter* with totalizer or a pair of *Single Counters*.

6.1.6 *Balance*, with a capacity of 3 mg and a sensitivity of 0.005 mg (needed for specimens taken from array length groups only).

6.1.7 *Mounting Medium*, sodium hydroxide (NaOH) solution, 18 %, sp gr 1.197 ± 0.002 at 60 to 70°F (16 to 20°C) in a dropping bottle.

6.2 Procedure 2:

6.2.1 *Polarizing Microscope* equipped with a polarizer, an analyzer, a first-order retardation plate, a cross-hair eyepiece mounted so that the hairs make a 45° angle with the plane of polarization, a rotatable, mechanical stage, and a microscope lamp. The magnification must be at least $100 \times$.

6.2.2 *Mounting Medium*, clear mineral oil in a dropping bottle.

6.2.3 *Other Apparatus* as specified in 6.1.2-6.1.6 for Procedure 1.

7. Safety Precaution

7.1 The sodium hydroxide solution used in Procedure 1 is caustic and corrosive. Use care in its preparation and application to avoid contact with the skin or with equipment, especially the microscope objective, which may be permanently damaged if the solution is not removed immediately following contact. Clear water and a soft tissue will remove the solution.

8. Sampling and Preparation of Specimens

8.1 Three sources of specimens may be used with either procedure. If Suter-Webb array length groups are not available, either of the other two sources of specimens may be used.

8.1.1 Option A—Suter-Webb Array Length Groups—Prepare the array length groups as directed in Method D1440. From one array discard the $\frac{1}{16}$ -in. (1.6-mm) and $\frac{3}{16}$ -in. (4.8-mm) length groups and any other length groups containing less than 1 mg of fibers. From each length group remaining, remove a bundle of approximately 100 fibers by lengthwise separation beginning with the longest group. Place the fibers on a microscope slide, spread them carefully to a width of 30 to 40 mm. Cover the fibers with a cover glass and apply a drop of the mounting medium to one corner. Tap the cover glass to cause the mounting medium to spread more rapidly and help prevent air bubbles. Mark the slide with the length group identification. The series of slides shall constitute a test specimen. Have a second operator prepare a second test specimen from a second array of the sample.

Note 2—The sampling method described in 8.1.1 has been used for a longer period of time and given slightly more reliable results than the other sampling methods.

8.1.2 Option B, Laboratory Blended Samples—Take a subsample consisting of a section of sliver approximately 2 in. (50 mm) long from the blended laboratory sliver. Twist one end of the subsample, hold it firmly and place the loose ends near the edge of a microscope slide. By means of a second slide held perpendicularly, grip a few fibers, hold them lightly and pull the subsample away gently. Repeat the process until approximately 200 fibers have been extracted. Pull the fibers from the entire width of the subsample and do not purposely discard any fibers. Spread the extracted fibers and separate them as evenly as possible, keeping them nearly parallel. A dissecting needle may be used to move the fibers while holding them lightly with a second slide or a cover glass. A minimum amount of overlapping will greatly facilitate fiber classification. Cover the