

Designation: D3376 - 88(Reapproved 2009)

# Standard Test Methods of Sampling and Testing Pulps to be Used in the Manufacture of Electrical Insulation<sup>1</sup>

This standard is issued under the fixed designation D3376; 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 ( $\varepsilon$ ) indicates an editorial change since the last revision or reapproval.

#### 1. Scope

1.1 These test methods cover the sampling and testing of cellulosic pulps for use in the manufacture of electrical insulating papers and boards or in the direct application of pulp fibers as insulation to electrical conductors.

NOTE 1-The significance of any one pulp property test method, as set forth herein, should be considered with discretion depending on the product made from the pulp.

1.2 Sections on Reagents, Sampling, and Report are integral parts of each of the individual test methods that follow.

1.3 Each test method is described as being a measure of either a bulk property of the pulp or a property of a handsheet formed from the pulp.

1.3.1 Bulk characteristics determinable by these procedures appear in the following sections:

Kappa Number/Permanganate Number (Substances Oxi dizable by Permangan- ate)	28 and 29	T 236, UM 251				
Laboratory Processing of Pulp (Beater Method)	54 and 55	T 200				
Moisture in Pulp	32 – 34	T 210				
Neutral Aqueous Extractable Hardness in Pulp	16 - 23 D202, D1126, and D2576					
Pentosan Content of Pulp	30 and 31	T 223				
Resistance of Pulp to Dis- inte gration (Standard RPG)	46 – 53	T 239, UM 252				
Shive Count	35 – 41					
Solvent-Soluble Matter in Pulp	26 and D202 27					
Tensile Properties	68 – 72 D202					
Water-Extractable Chlorides	14 and D202					
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1.3.2 Handsheet characteristics determinable by these procedures appear in the following sections:

		ASTM	TAPPI	cedures appear in the following sections.			
Procedure	Sec- tions	Method	Method		Sec-	ASTM Method	TAPPI Method
Flocedule	10115	neletetice	Relefence	Procedure	tion	Reference	Reference
Aqueous Extract Conductivity Aqueous Extract pH	8 and 9 10 and		ÄSTM D2276 8	Air Resistance (Porosity)	60 and	D202	T 205
Aqueous Extract pri	10 anu	D202	<u>MOTNI DJJ/0-0</u>		61	2202	. 200
Aqueous Extractable Acid-	12 and 13	0202 and ards/s	i <u>st</u> /a64f5173-42fa	Apparent Density b6cab139	66 and 67	5 <b>D202</b> d3376-88	20 <b>T</b> 205
	73 – 81	D1193 and D2576		Bursting Strength	62 and 63	D202 and D774/ D774M	T 205
photometry				Folding Endurance (M.I.T.)	54 and	D202 and D2176	T 205
Ash Content	82 - 85	D202	T 413		55		
Dirt in Pulp	42 and 43		T 213	Forming Handsheets for Physical Tests of Pulp	58 and 59		T 205
Fiber Analysis	24 and 25	D202 and D1030		Tensile Strength	68 and 69	D202 and D828	T 205
Fiber Length of Pulp	44 and 45		T 232, T 233	Note 2-Methods for Ash,	Silica,	selected cations fro	om Ash, Heat

NOTE 2-Methods for Ash, Silica, selected cations from Ash, Heat Stability,  $\alpha$ ,  $\beta$ , and  $\gamma$  Cellulose, Viscosity, Total Chlorine, Tear, and Dissipation Factor and Relative Permittivity, will be considered for addition as methods are developed.

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

Current edition approved Oct. 1, 2009. Published February 2010. Originally approved in 1975. Last previous edition approved in 2005 as D3776-88 (2005). DOI: 10.1520/D3376-88R09.

<sup>1</sup> These test methods are under the jurisdiction of ASTM Committee D09 on

Electrical and Electronic Insulating Materials and are the direct responsibility of

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Subcommittee D09.19 on Dielectric Sheet and Roll Products.

Freeness (Canadian Standard 56 and ...

Freeness)

1.5 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the

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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:<sup>2</sup>
- D202 Test Methods for Sampling and Testing Untreated Paper Used for Electrical Insulation
- D774/D774M Test Method for Bursting Strength of Paper (Withdrawn 2010)<sup>3</sup>
- D828 Test Method for Tensile Properties of Paper and Paperboard Using Constant-Rate-of-Elongation Apparatus (Withdrawn 2009)<sup>3</sup>
- D1030 Test Method for Fiber Analysis of Paper and Paperboard
- D1126 Test Method for Hardness in Water
- D1193 Specification for Reagent Water
- D2176 Test Method for Folding Endurance of Paper by the M.I.T. Tester (Withdrawn 2010)<sup>3</sup>
- D2576 Method of Test for Metals in Water and Waste Water by Atomic Absorption Spectrophotometry (Withdrawn 1979)<sup>3</sup>
- D3376 Test Methods of Sampling and Testing Pulps to be Used in the Manufacture of Electrical Insulation
- E29 Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications
- 2.2 TAPPI Standards:<sup>4</sup>
- T 200 Laboratory Processing of Pulp (Beater Method)
- T 205 Forming Handsheets for Physical Tests of Pulp
- T 210 Weighing, Sampling, and Testing Pulp for Moisture
- T 213 Dirt in Pulp
- T 221 Drainage Time of Pulp
- T 223 Pentosans in Wood and Pulp
- T 227 Freeness of Pulp
- htt T 232 Fiber Length of Pulp by Projection/sist/a6
  - T 233 Fiber Length of Pulp by Classification
  - T 236 Kappa Number of Pulp
  - T 413 Ash in Paper and Paperboard
  - T 445 Identification of Specks and Spots in Paper
  - T 1002 Drainage Time for Insulating Board
  - UM 203 Freeness of Pulp (William Tester)
  - UM 251 Permanganate Number of Pulp
  - UM 252 Resistance of Pulp and Paper Stock to Disintegration

# 3. Terminology

3.1 Definitions of Terms Specific to This Standard:

3.1.1 *pulp*, *n*—a fibrous material that is made by chemical or mechanical treatment, or both, of wood, cotton, hemp, or other

cellulosic fiber to achieve substantially separate fibers that are suitable for a sheet-forming process.

Note 3—Electrical insulation made from pulp may be papers or boards used for capacitors, transformer coils, creped papers, etc. It may also be pulp applied directly onto electrical conductors.

#### 4. Summary of Test Methods

4.1 These test methods describe the specific procedures for testing the properties of pulp, both in its original bulk form and after it has been formed into a handsheet in the testing laboratory.

#### 5. Reagents

5.1 *Purity of Reagents*—Use reagent grade chemicals 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, where such specifications are available.<sup>5</sup> 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.

5.2 *Purity of Water*—Unless otherwise indicated, references to water shall be understood to mean water conforming to Specification D1193, Type III.

# 6. Sampling

6.1 Terminology regarding sampling and evaluation terminology shall conform to those in the sampling sections of Test Methods D202.

6.2 Obtain the sample of pulp from the lot to be evaluated in a manner that will maximize the probability that a representative sample is collected. Where practicable, use one of the sampling plans shown in Test Methods D202. Protect the material sample from contamination during handling and transporting to a laboratory for testing. The instructions for preparation of specimens are given in the sections pertaining to the individual property tests. Take the sample for moisture content in accordance with TAPPI T 210.

6.3 Condition samples in a container suitable for preventing moisture variation over the period of testing. When test specimens are drawn, determine the moisture content of the material to allow correction of weights to moisture-free equivalent weight.

#### 7. Report

7.1 At the completion of any or all of the following tests, report the test results (as defined in 6.1) of the pulp properties with identifying units as follows:

7.1.1 Identification of the pulp sampled and tested by lot number, type, grade, etc.,

7.1.2 Dates of testing,

<sup>&</sup>lt;sup>2</sup> 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.

<sup>&</sup>lt;sup>3</sup> The last approved version of this historical standard is referenced on www.astm.org.

<sup>&</sup>lt;sup>4</sup> Available from Technical Association of the Pulp and Paper Industry (TAPPI), 15 Technology Parkway South, Norcross, GA 30092, http://www.tappi.org.

<sup>&</sup>lt;sup>5</sup> 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. Pharmaceutical Convention, Inc. (USPC), Rockville, MD.

7.1.3 Location of the testing laboratory and the person responsible for the testing,

7.1.4 Remarks indicating method or procedures used and the deviation, if any, from the standard test procedures,

7.1.5 Indication of the variance in test measurements (as defined in 6.1) such as high, low, standard deviation, etc., and 7.1.6 Any information particular to the cited procedure.

7.2 Report the test results (as defined in 6.1) as calculated or observed values rounded to the nearest unit in the last right-hand place of figures used in the material specification to express the limiting value. (See the rounding method of Practice E29.)

### AQUEOUS EXTRACT CONDUCTIVITY

#### 8. Significance and Use

8.1 The conductivity of the water extract of electrical grade pulp results from electrolytic impurities in the pulp which may be present as ionizable acids, bases, salts, or a combination of these. The presence of electrolytic impurities in electrical insulation is undesirable as they tend to lower insulation resistance and have corrosion-producing tendencies under conditions of applied potential. When comparing test data it should be noted that the extract conductivity of pulps, especially those of high purity, may change with time after manufacturing. This test is useful for routine acceptance testing, the comparison of different pulps, and research work.<sup>6</sup>

#### 9. Procedure

9.1 Follow Test Methods D202 except use a specimen weight equivalent to 1 g of moisture-free pulp.

# AQUEOUS EXTRACT pH

#### 10. Significance and Use atalog/standards/sist/a641517

10.1 The extract pH determination measures the degree to which a pulp alters the hydrogen-hydroxyl equilibrium of pure water. The test gives a measure of the active acidity or alkalinity of the pulp extract. The presence of active acidic or alkaline contaminants in a pulp may result in their being incorporated into the electrical insulation made from the pulp, and can lead to a deterioration of the insulation in service. This test is useful for routine acceptance testing, the comparison of different pulps, and research work.<sup>2</sup>

#### 11. Procedure

11.1 Follow Test Methods D202 except use a specimen weight equivalent to 1 g of moisture-free pulp.

# AQUEOUS EXTRACTABLE ACIDITY-ALKALINITY

# 12. Significance and Use

12.1 The extract acidity-alkalinity determination for a pulp measures the quantity of extracted ionizable material, which

alters the hydrogen-hydroxyl equilibrium of pure water. The presence of active acidic or alkaline contaminants in a pulp may result in their being incorporated into the electrical insulation made from the pulp, and can lead to a deterioration of the insulation in service. This test is useful for routine acceptance testing, the comparison of different pulps, and research.<sup>4</sup>

#### 13. Procedure

13.1 Follow Test Methods D202 except use a specimen weight equivalent to 1 g of moisture-free pulp.

# WATER-EXTRACTABLE CHLORIDES

#### 14. Significance and Use

14.1 The occurrence of significant amounts of chloride ion in a pulp may lead to the incorporation of the ion in the electrical insulation made from the pulp. The presence of chloride ions may adversely affect the electrical properties and service life of the insulation. This test is useful for routine acceptance testing, the comparison of different pulps, and research testing.

#### 15. Procedure

15.1 Follow Test Methods D202 except use a specimen weight equivalent to 4 g of moisture-free pulp. For pulps with higher levels of chloride (greater than 30 ppm) 10 min of masceration as in the above method for aqueous extract conductivity may be used to hasten the extraction followed by 1 h refluxing as in Test Methods D202. When the chloride content is less than 30 ppm, masceration is not permitted. The appropriate extraction time must be determined to give complete extraction of the chloride for each pulp type. Times greater than 1 h may be necessary.

#### NEUTRAL AQUEOUS EXTRACTABLE HARDNESS PULP

# 16. Terminology

16.1 Definitions of Terms Specific to This Standard:

16.1.1 *aqueous extractable hardness, n*—the amount of calcium and magnesium present in pulp and which may be extracted by hot neutral water under prescribed conditions.

16.1.2 *hardness*, n—a characteristic of water that represents the total concentration of calcium and magnesium in the water expressed as parts per million (ppm) CaCO<sub>3</sub>.

#### 17. Significance and Use

17.1 Cellulose pulps may contain varying amounts of aqueous extractable hardness as supplied to the purchaser. The dissolved hardness from the pulp may accumulate in process water used in wet-forming methods and may interfere with the action of process additives and affect product quality adversely.

17.2 Method A is the preferred method and shall be used for reference purposes.

# METHOD A

#### 18. Procedure

18.1 Extraction:

<sup>&</sup>lt;sup>6</sup> For more detailed information see *Paper and Paperboard—Characteristics*, *Nomenclature, and Significance of Tests, ASTM STP 60 B*, Am. Soc. Testing Mats., 1963, pp. 59–61.

18.1.1 Prepare extracts of the pulp specimens in accordance with the Test Methods D202 method for aqueous extract conductivity, except:

18.1.2 Use a specimen weight equivalent to 2.0 g of moisture-free pulp. Determine the moisture content of the pulp sample on a separate specimen taken at the same time as the test specimen.

18.1.3 The extraction volume shall be 200 mL.

18.1.4 Run a blank determination concurrently with the test specimen determination.

18.1.5 Following extraction and filtration, collect the clear filtrate and adjust the volume to exactly 200 mL.

18.2 Determine the calcium and magnesium concentration of the extract in accordance with Test Method D2576.

## **19.** Calculation

19.1 Calculate the hardness of the extracts as follows:

Hardness, ppm = 
$$100[2.497(P_1 - P_a) + 4.117(P_2 - P_b)]$$
 (1)

where:

 $P_1$  = ppm calcium in the pulp extract,

 $P_a$  = ppm calcium in the blank,

 $P_2$  = ppm magnesium in the pulp extract, and

 $P_{h}$  = ppm magnesium in the blank.

# METHOD B iTeh Sta

#### **20.** Procedure

20.1 Follow the procedure of Method A for the preparation of the extract.

20.2 Take two 100-mL aliquots of the extract and titrate for total hardness following the "low total hardness" procedure of the nonreferee volumetric method of Test Method D1126.

# 21. Calculation, iteh ai/catalog/standards/sist/a64f5173-4

21.1 Calculate the hardness of the specimen extract as follows:

Hardness, ppm = 500 
$$(V_1 + V_2 - V_a - V_b)$$
 (2)

where:

- $V_1$  = standard EDTA solution for titration of first aliquot of extract, mL
- $V_2$  = standard EDTA solution for titration of second aliquot of extract, mL
- $V_a$  = standard EDTA solution for titration of first blank aliquot, mL, and
- $V_b$  = standard EDTA solution for titration of second blank aliquot, mL.

# 22. Report

22.1 Report the results as neutral aqueous extractable hardness, ppm, expressed as calcium carbonate according to the appropriate method of Test Methods D3376.

# 23. Precision and Bias

23.1 The precision of this test has not been determined. No statement can be made about the bias of this test since standard material is not available.

# FIBER ANALYSIS

#### 24. Significance and Use

24.1 The fiber composition of a pulp (fiber source and pulping treatment) strongly affects the ultimate product characteristics. Fiber analysis is useful both as a specification and as a control test, and may be used in referee testing or research.

Note 4—For accurate results the analyst should have considerable training and experience. The analyst should make frequent use of standard fiber sources of known composition, or of authentic fiber samples, and be thoroughly familiar with different fibers and their behavior when treated with the various stains.

# 25. Procedure

25.1 Follow Test Methods D202 and D1030 using a specimen weight of 0.2 g of pulp drawn from a composite sample of pulp equivalent to 30 g of moisture-free pulp disintegrated in 2 L of water as in Test Method D1030 and the section on Fiber Analysis of Test Methods D202.

# SOLVENT-SOLUBLE MATTER IN PULP

#### 26. Significance and Use

26.1 Since pulping processes usually remove most watersoluble and volatile compounds that are also soluble in organic solvents, the solvent extractives in pulp may be considered to consist primarily of resin and fatty acids, their esters, waxes, and unsaponifiable substances. No single organic solvent is capable of removing all these substances, and different solvents remove different combinations. The mixture of 1 part by (volume) of 95 % ethanol and 2 parts of benzene appears to provide the most complete removal of all solvent-extractable substances in pulp.

26.2 Solvent-extractable materials, if present in sufficient quantity in electrical insulation, may lower the quality of the insulation or have a deleterious effect on the liquid compounds used in contact with insulation in various types of apparatus. Ethanol-soluble materials in capacitor paper have been found to increase the conductivity of chlorinated organic compounds, which are used as impregnants in capacitors. A combination of ether and alcohol-benzene extractives is reported to represent the quantity of pitch in wood pulp. Pitch in pulp may cause operating problems in the paper mill, such as the plugging of felts, wires, etc.

26.3 This method, with solvent or solvents specified, may be used for routine acceptance testing, comparing different pulps, and research tests.

# 27. Procedure

27.1 Test for solvent-soluble matter in accordance with Test Methods D202.

# KAPPA NUMBER/PERMANGANATE NUMBER (SUBSTANCES OXIDIZABLE BY PERMANGANATE)

#### 28. Significance and Use

28.1 The permanganate consumption of a pulp is a measure of its lignin content and may be used as a measure of lot-to-lot

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uniformity. For a given species of wood or other fibers, the permanganate consumption is a measure of the degree of cooking to which the fiber has been subjected, and can relate to the ease of refining of the pulp produced from the cooked fiber. This test is useful for control purposes, specifications, and the comparison of different pulps.

# 29. Procedure

29.1 Kappa Number-Follow TAPPI T 236.

29.2 Permanganate Number—Follow TAPPI UM 251.

# PENTOSAN CONTENT OF PULP

### 30. Significance and Use

30.1 The pentosan content of a pulp strongly affects the dissipation factor—temperature relationship of electrical insulation made from it. It also may be an indicator of lot-to-lot uniformity and is one of several factors related to the bonding power of a pulp, and the amount of energy required to refine the pulp.

#### **31. Procedure**

31.1 Follow TAPPI T 223.

#### **MOISTURE IN PULP**

# 32. Significance and Use

32.1 Pulp is purchased on the basis of moisture content. In addition, the moisture content may be used for consistency control, and it may affect the energy of repulping and the biological degradation of the pulp. This test is useful for control purposes and specifications.

#### **33.** Procedure

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htt 33.1 Follow TAPPI T 210. log/standards/sist/a6415173-4

#### 34. Report

34.1 This method gives the percentage moisture content of the pulp. Report percent water and percent moisture-free fiber (equal to 100 minus percent water).

#### SHIVE COUNT

#### 35. Terminology

35.1 Definitions of Terms Specific to This Standard:

35.1.1 *shive*, *n*—a particle in pulp or paper that is a bundle of cellulosic fibers bonded together in a parallel arrangement.

Note 5—Dark single fibers are not to be counted as shives. Count only bundles of fibers regardless of color.

35.1.2 *shive count, n*—the quantitative expression of the concentration of shives in a quantity of pulp or paper. For this method the shive count is restricted to the number of shives that exceed 1.5 mm in length that are present after a specified processing of the pulp to form handsheets for evaluation.

#### 36. Significance and Use

36.1 Several grades of electrical insulating paper are most effectively manufactured using pulps having a low shive count.

Shives in wood pulp to be used for direct application to electrical conductors can be detrimental to the insulating characteristics and strength of the insulating wall. This test is useful for control purposes, specifications, and the comparison of different pulps.

### **37.** Apparatus

37.1 Disintegrator, Sheet Machine, Press, Blotters, etc., in accordance with TAPPI T 205.

37.2 Steel Rule, graduated in 0.5 mm.

37.3 Balance, to weigh up to 100 g with 0.1-g accuracy.

37.4 *Specimen Viewer* with white opaque glass and a fluorescent or incandescent light source.

37.5 Transparent Cylinder, 0.6 L.

#### 38. Procedure

38.1 From the sample obtained as specified in Section 6, take a quantity of pulp equivalent to 30 g of moisture-free pulp. Soak this pulp in 0.5 L of water for 4 h at a temperature of 20 to  $30^{\circ}$ C.

38.2 Tear the pulp into smaller pieces (approximately 1 in. or 25 mm square) and dilute to 2.0 L. **Warning**—*Tear*, do not cut the pulp.

38.3 Using the TAPPI disintegrator, disintegrate for 10 minute minimum. This time should be sufficient to disperse the pulp completely. A technique for checking the dispersion is as follows:

38.3.1 Take a small sample of the slurry from the disintegrator (about 2 or 3 mL) and dilute to 0.6 L in a clear cylinder. Stopper the cylinder and mix the suspension by rotating the cylinder end over end. Observe the suspension by looking through it toward a strong light source. The suspension should be free of clumps or agglomerates of fibers, but may contain shives. If the suspension contains clumps or agglomerates, subject the pulp to additional 5-min periods of disintegration until it is free of clumps and agglomerates.

38.4 Dilute the disintegrated pulp with water to result in a consistency of 0.3 % or 3 g/L.

38.5 Clean the sheet machine thoroughly.

38.6 Form handsheets in accordance with TAPPI T 205 and couch but do not dry the sheets.

38.7 Make at least five handsheets for viewing.

38.8 On each of five handsheets, mark out six viewing areas. Each viewing area shall be 625 mm<sup>2</sup>. This step can be facilitated by having previously made a transparent plastic overlay grid with these areas cut out.

38.9 With the viewer and the steel rule, count the shives that exceed 1.5 mm in length that are viewed within each of the six areas. Record the total number of shives in each handsheet.

### **39.** Calculation

39.1 Add the total shives found in all five handsheets. This sum multiplied by 53 yields the shive count expressed as shives per square metre.

39.2 The shive count may be calculated for shives per kilogram if the handsheet area is 0.02 m<sup>2</sup>. Shives per kilogram is the product of total shives counted times 888. This multiplier is valid only for the standard handsheets in accordance with TAPPI T 205 with a grammage of  $60 \pm 1.2 \text{ g/m}^2$  and an area of  $0.02 \text{ m}^2$ .

# 40. Report

40.1 Report the disintegration time if more than 10 min.

40.2 Report the shive count as shives per square metre or as shives per kilogram as agreed upon between the supplier and the purchaser.

# 41. Precision and Bias

41.1 Precision—From a round-robin test involving 3 laboratories, a coefficient of variation between laboratories in the order of 300 was obtained, at a level of 5000 to 8000 shives/m<sup>2</sup>.

41.2 Bias-No statement of bias can be made because of the unavailability of standard reference material.

#### **DIRT IN PULP**

#### 42. Significance and Use

42.1 Dirt content is one indication of the quality of the pulp. This measure gives only a visual indication of contamination. The nature of foreign particles is very significant in determining whether the contamination is detrimental to the endproduct use. TAPPI T 445 may be appropriate for identification of the particulate contamination. This test is useful for control purposes and the comparison of different pulps.

# 43. Procedure

htt 43.1 Follow TAPPI T 213 . g/standards/sist/a6415173

# **FIBER LENGTH OF PULP**

#### 44. Significance and Use

44.1 The fiber length is a means of comparing pulps. The fiber length distribution may affect the forming characteristics, which in turn influence the physical characteristics of the end-product. This method is useful for control purposes, the comparison of different pulps, and research.

# 45. Procedure

45.1 Follow TAPPI T 232, T 233.

# **RESISTANCE OF PULP TO DISINTEGRATION** (STANDARD RPG)

Note 6-The method described herein is essentially an adaptation of TAPPI UM 252, with several significant changes. The method is complete and no reference to UM 252 is required in its use.

# 46. Terminology

# 46.1 Definitions of Terms Specific to This Standard:

46.1.1 resistance to disintegration, n-the amount of work (expressed as revolutions per gram of pulp) required under standard conditions to bring a sample of pulp to a state of complete dispersion of single fibers.

# 47. Significance and Use

47.1 Resistance to disintegration is important in that it is a measure of repulpability. The method is useful for control purposes and the comparison of different pulps.

# 48. Apparatus

48.1 Disintegrator-TAPPI standard disintegrator in accordance with T 205; equipped with a 2 L standard vessel, timer/timer controlled power source, 1-s maximum time division

48.2 Handsheet Machine-British Standard sheet machine in accordance with T 205, with couch roll, couch plate, 8 by 8-in. (200 by 200-mm) blotters, and a hot plate.

48.3 Beakers-Set of twelve numbered 50-mL beakers and set of twelve numbered 600-mL beakers.

48.4 Graduated Cylinder, 500-mL, transparent.

48.5 Vessel to contain 1500 mL or more pulp, calibrated at 1500 mL.

48.6 Sample Cup, to hold 50 mL of pulp slurry.

48.7 Oven, drying, controlled at  $105 \pm 3^{\circ}$ C.

48.8 *Balance*, to weigh up to 100 g with 0.1-g accuracy.

48.9 Weighing Bottle, to hold 10 g of pulp.

48.10 Light Box-Source of uniform illumination (incandescent or fluorescent) with surface sufficiently large to hold two handsheets- 160-mm diameter disks.

# 49. Test Specimens

STM D3376-88(49.1) From the sample obtained as specified in Section 6, weigh out three specimens each equivalent to 60 g of moisturefree pulp. Determine the moisture content in accordance with Sections 32 - 34. Tear, do not cut, the test specimen from the test unit.

# 50. Procedure

50.1 Connect the disintegrator to the timed power outlet, and fill the disintegrator vessel with 2 L of water ( $25 \pm 5^{\circ}$ C). Select a disintegration period appropriate to the resistance of the pulp to disintegration.

NOTE 7-The times shown below may be used as a guide for selecting the appropriate disintegration period.

Resistance of Pulp to Dispersion **Disintegration Period** 

slight	15
moderate	30
high	45

50.1.1 Take one specimen and briefly immerse it in the water in the disintegrator vessel to wet it thoroughly (for example, for 15 s) in water at  $25 \pm 5^{\circ}$ C. Tear the specimen into pieces about 20 mm square. If the pulp is flash dried it may be split so that the specimen pieces are about 2.5 mm thick.

50.1.2 Quickly put the torn specimen into the disintegrator vessel and start the disintegration/sampling sequence. Disintegrate the specimen for one period, stop the disintegrator,