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The 2007 edition of the Materials Manual has been revised. It continues to use AASHTO, ASTM, WAQTC, and WSDOT test methods. The strategic directions for the materials laboratory is to continue to expand the use of AASHTO and ASTM standards whenever possible.

The manual has retained its dual unit format. However, English units predominate with metric units in parenthesis. WSDOT has returned to using English units, but some contracts that are in the pipeline are still in metric units.

This manual reflects the Quality System concerns of an AASHTO accredited organization. The format of this manual adopted in 2000 has been well received. The manual continues to be organized by numerical test order. It also features two contents and an index.

The manual reflects a continuing policy of adopting “Consensus” standards wherever practical. Adoption of these, in the form of AASHTO, ASTM, WAQTC, or other nationally recognized standards eliminates much of the previous text, which merely recopied the national documents. By adopting these standards, we provide a common standard that can be used by neighboring states and other laboratories or organizations. Contractors, who work in more than one state, also benefit by having to conform with fewer unique tests.

The concept of “Field Operating Procedures” (FOP) is continued to support the work of Materials Testers at the Field or Project level. Full procedures are provided when WSDOT Test Methods apply, or we have adopted the full AASHTO procedure. Other cases, when a consensus standard (AASHTO, ASTM, or WAQTC) has been adopted, the FOP provides the essential performance elements for the field technician.

The WSDOT Materials Laboratory is responsible for establishing and managing all test procedures. For technical information or suggested changes to test methods or procedures, contact the WSDOT Materials Laboratory Quality Systems Manager through the departmental mail system at MS 47365; by mail at PO Box 47365, Olympia, WA 98504-7365; by telephone at (360) 709-5497; or by fax at (360) 709-5588, physically located at 1655 South Second Avenue, Tumwater, WA 98512. Please use this physical address for all communications other than US Postal Service mail.

This manual will be updated in the future as needed. Contact the Directional Documents and Engineering Publications office at (360) 705-7430 or http://www.wsdot.wa.gov/FASC/EngineeringPublications/ to order manuals or view electronic documents.

Thomas E. Baker
State Materials Engineer
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**Bituminous Cement**

<p>| D         | 4      | ASTM  |           |           | Standard Test Method for Bitumen Content |
| R         | 28     | AASHTO|           |           | Practice of Accelerated Aging of Asphalt Binder Using a Pressurized Aging Vessel |
| R         | 29     | AASHTO|           |           | Practice for Grading or Verifying the Performance Grade of an Asphalt Binder |
| T         | 40     | AASHTO|           |           | Sampling Bituminous Materials |
| T         | 40     | WSDOT | ✔         | ✔         | FOP for WAQTC/AASHTO for Sampling Bituminous Materials |
| T         | 44     | AASHTO|           |           | Solubility of Bituminous Materials |
| T         | 47     | AASHTO|           |           | Loss on Heating of Oil and Asphaltic Compounds |
| T         | 48     | AASHTO|           |           | Flash and Fire Points by Cleveland Cup |
| T         | 49     | AASHTO|           |           | Penetration of Bituminous Materials |
| T         | 50     | AASHTO|           |           | Float Test for Bituminous Materials |
| T         | 51     | AASHTO|           |           | Ductility of Bituminous Materials |
| T         | 53     | AASHTO|           |           | Softening point of Bituminous (Ring and Ball Apparatus) |
| T         | 55     | AASHTO|           |           | Water in Petroleum Products and Bituminous Materials by Distillation |
| T         | 59     | AASHTO|           |           | Testing Emulsified Asphalts |
| T         | 72     | AASHTO|           |           | Saybolt Viscosity |
| T         | 78     | AASHTO|           |           | Distillation of CUT Back Asphaltic (Bituminous) Products |
| T         | 79     | AASHTO|           |           | Flash Point with Tag Open-Cup Apparatus for Use with Materials Having a Flash Less Than 93.3°C (200°F) 207 |
| T         | 111    | AASHTO|           |           | Inorganic Matter or Ash in Bituminous Materials |
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**Chemical**

<p>| T         | 65     | AASHTO|           |           | Mass (Weight) of Coating on Iron and Steel Articles with Zinc or Zinc-Alloy Coatings |
| T         | 267    | AASHTO|           |           | Determination of Organic Content in Soils by Loss on Ignition |
| T         | 405    | WSDOT | ✓         |           | Method of Test for Diagonal Shear Strength of Epoxy Resins (Arizona Composite Cylinder Test) |
| T         | 408    | WSDOT | ✓         |           | Method of Test for Quality of Water to be Used in Mixing Concrete |
| T         | 411    | WSDOT | ✓         |           | Method of Test for Water Absorption and Moisture Vapor Transpiration |
| T         | 412    | WSDOT | ✓         |           | Bond Test for Joint Sealants |
| T         | 414    | WSDOT | ✓         |           | Method of Test for Total Chloride Ion in Concrete |
| T         | 415    | WSDOT | ✓         |           | Method of Test for Fertilizer |
| T         | 418    | WSDOT | ✓         |           | Method of test for Corrosion of Deicing Materials |</p>
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**Concrete**

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| T         | 423    | WSDOT  |           | ✓         | Traffic Controller Conflict Monitor Testing |
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**Paint**

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**Standard Practice**

<p>| QC 1 | WSDOT | ✓ | Standard Practice for Cement Producers That Certify Portland Cement |
| QC 2 | WSDOT | ✓ | Standard Practice for Asphalt Suppliers That Certify Performance Graded Asphalts |</p>
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WSDOT Standard Practice QC 1

Standard Practice for Cement Producers

That Certify Portland Cement

1. SCOPE

This standard specifies requirements and procedures for a certification system that shall be applicable to all producers of Portland Cement.

This standard may involve hazardous materials, operations and equipment. It does not address all of the safety problems associated with their use. It is the responsibility of those using this standard to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:
   2.1.1 M-85 Standard Specifications for Portland Cement
   2.1.2 M-240 Standard Specifications for Blended Hydraulic Cement
   2.1.3 R-18 Establishing and Implementing a Quality System for Construction Materials Testing Laboratories

2.2 Agency’s Standard Specifications

3. TERMINOLOGY

3.1 AASHTO — American Association of State Highway and Transportation Officials

3.2 CCRL — Cement and Concrete Reference Laboratory

3.3 NIST — National Institute of Standards and Technology

3.4 Production Facility — A facility that has the capacity for grinding cement.

3.5 Supplier — A supplier stores and then delivers cement produced by another entity to a concrete plant or another supplier.

3.6 Supplier Certification — If a supplier makes any modifications to the cement or blended cement, then the supplier must provide the certification.

3.7 Agency — State highway agency or other agency responsible for the final acceptance of cement or blended cement.

3.8 Specification Compliance Testing — Complete testing in accordance with the specification requirements.

3.9 Quality Control Testing — The quality control testing shall be described in the Production Facility’s quality control plan. The Production Facility’s quality control plan must be approved by the Agency.

3.10 CAP — Cement Acceptance Program
4. SIGNIFICANCE AND USE

4.1 This standard specifies procedures for accepting Portland Cement and Blended Cement. This is accomplished by a certification system that evaluates quality control and specification compliance tests performed by the Production Facility according to their quality control plan.

5. LABORATORY AND TESTER REQUIREMENTS

5.1 Laboratories shall be AASHTO accredited in all tests required by specification compliance testing or meet the following requirements:

5.1.1 Laboratory facilities shall adequately house and allow proper operation of all required equipment in accordance with the applicable test procedures.

5.1.2 The laboratory shall use personnel qualified in accordance with the appropriate sections of AASHTO R-18.

5.1.3 The laboratory shall use testing equipment that has been calibrated/verified to meet the requirements of each test procedure in accordance with the appropriate sections of AASHTO R-18.

5.1.4 Documentation of personnel qualifications and the equipment certification/verification records shall be maintained.

5.1.5 The agency at their discretion may review the laboratory facility, testing equipment, personnel doing the testing, and review all qualification and calibration and verification records.

5.1.6 The laboratory must participate in the NIST’s CCRL proficiency sample program.

6. PRODUCTION FACILITY QUALIFICATION

6.1 The Production Facility shall submit a written request for acceptance into the Cement Acceptance Program to the Agency along with a copy of the Production Facility’s Quality Control Plan.

6.2 The Production Facility shall submit one sample with its Mill Test Report for the initial lot for each type of cement or blended cement it intends to provide to the Agency.

6.3 Initial lots shall be tested for conformance to Agency Standard Specifications and both physical and chemical requirements of AASHTO M-85 and/or M-240.

6.4 The Production Facility shall allow the Agency to visit and observe the quality control activities and obtain samples for test.

7. PRODUCTION FACILITY QUALITY CONTROL PLAN

7.1 The quality control plan, as a minimum, shall identify the following:

7.1.1 Facility type

7.1.2 Facility location.

7.1.3 Name and telephone number of the contact person responsible for the quality control of the facility.

7.1.4 The quality control tests to be performed on each type of cement or blended cement.

7.1.5 Name of the laboratory performing quality control tests on the cement if independent of the Production Facility.
7.1.6 Declaration stating that if a test result indicates that a lot of cement or blended cement is not in compliance with the specifications, the facility shall immediately notify the Agency of the lot in question.

7.1.7 Description of the method and frequency for sampling, quality control testing, and specification compliance testing.

7.1.8 Type of cement and/or blended cement the Production Facility intends to provide to the Agency.

7.1.9 Show compliance with Section 5.

7.2 The Quality Control Plan shall be submitted to the Agency annually for review.

8. DOCUMENTATION REQUIREMENTS

8.1 Each Production Facility shall document its conformance to the Agency’s Standard Specifications and both physical and chemical requirements of AASHTO M-85 and/or M-240 by means of a Mill Test Report certifying analysis.

8.2 The Mill Test Reports shall be provided to the agency on a continuous basis for AASHTO M-85 or M-240 cement production.

8.3 Separate sequences of Mill Test Reports shall be provided for each individual production facility and a unique lot number traceable to a production run on cement shall identify each report.

8.4 Mill Test Reports shall show the applicable test results and the applicable specifications of AASHTO M-85 or M-240 for each component or property tested and shall show the test requirements specified by the Agency.

8.5 Mill Test Reports of all cements shall be submitted on a monthly basis to the Agency. Negative reports (i.e., reports indicating no production for the month) are requested to insure that a continuous flow of documentation is maintained.

9. AGENCY REQUIREMENTS

The Agency will review the Production Facility’s quality control plan listed in Section 6 and respond to the Production Facility within 30 days.

The Agency may perform quality assurance or acceptance sampling and testing in accordance with the agency standards.

10. REQUIREMENTS FOR SHIPPING CEMENT

10.1 The Production Facility’s quality control plan as approved by the Agency (see Section 9) shall be implemented.

10.2 Each cement shipment shall identify the applicable Mill Test Report. This may be included on the Bill of Lading for the shipment or provided by other means as long as each shipment can be traced to the applicable Mill Test Report.

11. SPLIT SAMPLE TESTING

11.1 All test samples required by this standard shall be obtained as provided in the applicable standard specification or the Production Facility’s quality control plan.

11.2 Production Facilities, on a quarterly basis, shall split a production sample into two portions (10 pounds each).

11.3 The Production Facility or an independent test facility meeting the requirements specified in Section 5 shall conduct chemical and physical testing on one portion.
11.4 The other portion, along with accompanying chemical and physical analysis, shall be submitted to the Agency. This sample will include the Mill Test Report for the lot number that is traceable to this production run of cement.

11.5 The Production Facility shall submit, at least once each calendar year, a quarterly sample of each type of cement anticipated for inclusion in this program.

12. COMPARISON OF SPLIT SAMPLE TEST RESULTS

12.1 The Agency may elect not to test their portion, but when the Agency does elect to test, the Agency may conduct chemical and/or physical tests.

12.2 When deviation between the Production Facility’s test results and the Agency’s test results is excessive (defined by Table 1 for chemical analysis and the AASHTO or ASTM multilaboratory precision and bias for the physical tests), the Agency will provide its results to the Production Facility for comment.

12.3 The Production Facility shall respond within 30 days and address the following points concerning their results:
   a. Did the results reported accurately reflect the results obtained?
   b. Were the test results obtained, properly transferred to the data sheet submitted?
   c. Were all the calculations leading to the test result, correct?
   d. Did the equipment used to perform the test meet specification requirements?
   e. Did the procedures followed when performing the test conform to specification requirements?
   f. Was corrective action taken to repair or replace defective equipment, or the technician instructed of the correct procedure to follow?

12.4 The Production Facility shall prepare a response to the Agency, summarizing the results of the investigation, identifying the cause of the excessive deviation, if determined, and describing any corrective action taken. Comments may include the test facility’s data from CCRL Proficiency Tests.

13. REVOCATION OF CERTIFICATION STATUS

13.1 A Production Facility may have its certification status with the Agency revoked if found in nonconformance with the Standard Specifications or this Standard Practice.

13.2 The following criteria will be used to judge the conditions of nonconformance:
   13.2.1 Failure to follow the Production Facility’s approved quality control plan as required in Section 8.
   13.2.2 Failure to declare that test results indicated that a lot of cement or blended cement was not in compliance with the specifications as required under Section 8.2.
   13.2.3 When a test report shows nonconformance to the applicable specification, the results will be referred for comment and action to the Production Facility.
       13.2.3.1 The Production Facility shall submit one sample for retest from the next two available production runs.
       13.2.3.2 If two of three successive samples shows nonconformance the Agency will revoke certification status.

13.3 A Production Facility that has had its certification status revoked may seek reinstatement by demonstrating conformance to the qualification criteria shown in Section 7.
WSDOT Standard Practice QC 2

Standard Practice for Asphalt Suppliers

That Certify Performance Graded Asphalts

1. SCOPE

1.1 This standard specifies requirements and procedures for a certification system that shall be applicable to all suppliers of performance graded asphalt binder (PGAB). The requirements and procedures cover materials manufactured at refineries, materials mixed at terminals, in-line blended materials, and materials blended at the hot mix plant.

1.2 This standard may involve hazardous materials, operations and equipment. It does not address all of the safety problems associated with their use. It is the responsibility of whoever uses this standard to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:

MP1 Specifications for Performance Graded Asphalt Binders
PP6 Practice for Grading or Verifying the Performance Grade of an Asphalt Binder
T 40 Method of Sampling Bituminous Materials
R-18 Establishing and Implementing a Quality System for Construction Materials Testing Laboratories

2.2 ASTM Standards

D 8 Definitions of Terms Relating to Materials for Roads and Pavements
D 3665 Random Sampling of Construction Materials

3. TERMINOLOGY

3.1 AAP — AASHTO Accreditation Program

3.2 Asphalt Binder — An asphalt-based cement that is produced from petroleum residue either with or without the addition of modifiers.

3.3 ASC — Approved Supplier Certification

3.4 Modification — Any manufacturing process which alters the properties of a single asphalt binder for the purposes of meeting the requirements of a PGAB.

3.5 PGAB — Performance Graded Asphalt Binder

3.6 Supplier — A supplier shall be defined as one who produces the final PGAB product, or who makes the blend or modifications that alters properties of the PGAB. This process can occur at the refinery, terminal, the HMA Plant, or in a transport vehicle.

If no modifications are made to the PGAB after its initial production at the refinery, the refinery may be the supplier and can provide the certification.

If any modifications are made to the PGAB at the terminal or in the transport vehicle, then the terminal or transporter shall be the supplier and must provide the certification.
If any modifications, blending or commingling of PGAB from different sources is made at the HMA Plant, then the HMA producer shall provide the certification for the PGAB.

3.7 Agency — Agency shall be defined as a state highway agency or other agency responsible for the final acceptance of the PGAB.

3.8 Specification Compliance Testing – Complete testing in accordance with the specification requirements.

3.9 Quality Control Testing – The quality control testing shall be described in the Supplier’s quality control plan. The Supplier’s quality control plan shall be approved by the Agency.

3.10 HMA- Hot Mix Asphalt

Note 1 – Definitions for many terms common to asphalt binder are found in ASTM D8.

4. SIGNIFICANCE AND USE

4.1 This standard specifies procedures for minimizing disruption PGAB shipments. This is accomplished by a certification system that evaluates quality control and specification compliance tests performed by the Supplier according to their quality control plan.

5. SAMPLING

5.1 All test samples required by this standard shall be obtained in accordance with AASHTO T 40. The use of a random sampling procedure similar to ASTM D3665 is important to the establishment of a valid certification program.

6. LABORATORY AND TESTER REQUIREMENTS

6.1 AASHTO accreditation in any test required by this standard is applicable. Laboratories which are not AASHTO accredited must meet the following requirements:

6.1.1 Laboratory facilities shall adequately house and allow proper operation of all required equipment in accordance with the applicable test procedures.

6.1.2 The laboratory shall use personnel qualified in accordance with the appropriate sections of AASHTO R-18.

6.1.3 The laboratory shall use testing equipment that has been calibrated/verified meet the requirements of each test procedure in accordance with the appropriate sections of AASHTO R-18.

6.1.4 Documentation of personnel qualifications and the equipment verification records shall be maintained.

6.1.5 The agency at their discretion may review the laboratory facility, testing equipment, personnel doing the testing, and review all qualification and calibration and verification records.

7. SUPPLIER REQUIREMENTS

7.1 The Supplier shall submit a written request to the Agency for authorization to supply PGAB.

7.1.1 The request shall include copies of their preliminary test reports for the proposed PGAB with the appropriate documentation. If requested by the agency, a sample of the PGAB shall be provided to the agency for testing.

7.2 The Supplier shall allow the Agency to visit the production and/or shipping site to observe the Supplier’s quality control activities, and to obtain samples for test.
7.3 The Supplier shall submit to the Agency for approval a complete quality control plan that complies with the requirements of Section 8.

7.4 The Supplier shall follow the procedures described in the approved quality control plan.

7.5 The Supplier shall establish a continuing test record for each test required on each PGAB.

8. SUPPLIER QUALITY CONTROL PLAN (MINIMUM REQUIREMENTS)

8.1 The Supplier’s quality control plan shall identify the following:

8.1.1 Facility type (refinery, terminal, in-line blending or HMA plant).

8.1.2 Facility location.

8.1.3 Name and telephone number of the contact person responsible for the quality control of the PGAB at the facility.

8.1.4 The quality control tests to be performed on each PGAB.

8.1.5 Name the laboratory performing quality control tests on the PGAB that is shipped.

8.2 The Supplier’s quality control plan shall include a declaration stating that if a test result indicates that a shipment of PGAB is not in compliance with the purchase specifications, the Supplier shall (1) immediately notify the Agency of the shipment in question, (2) identify the material, (3) cease shipment until material complies with the specification, (4) notify the Agency prior to resuming shipment.

8.3 The Supplier’s quality control plan shall describe method and frequency for, sampling, quality control testing and specification compliance testing.

8.3.1 Specification Compliance Testing shall be performed on at least 3 random samples at the beginning of supply of a production PGAB. After 3 samples have met the required specifications, the quality control plan shall identify the frequency for specification compliance testing. If tests indicate non compliance, resume full specification compliance testing as stated above.

8.3.2 Quality Control Testing as identified in the quality control plan can be specification compliance testing or non specification compliance testing. The quality control testing does not preclude the need to meet the agency specifications.

8.4 The Supplier’s quality control plan shall include a statement that the Supplier will prepare reports for all quality control and specification compliance tests performed during a given period and submit them to the Agency on request.

8.5 The Supplier’s quality control plan shall provide the procedure to be followed for checking transport vehicles before loading to prevent contamination of shipments.

9. AGENCY REQUIREMENTS

9.1 The Agency shall review the Supplier’s quality control plan and respond to the supplier within 30 days.

9.2 The Agency may perform quality assurance or acceptance sampling and testing in accordance with the agency standards.

10. REQUIREMENTS FOR SHIPPING PGAB BY AN APPROVED SUPPLIER

10.1 The Supplier’s quality control plan as approved by the Agency (see Section 9) shall be implemented.
10.2 Each shipment shall be accompanied by two copies of the bill of lading, which shall include (1) the name and location of the Supplier, (2) the performance grade of material, (3) the quantity of material shipped, (4) the date of shipment, (5) a statement certifying the material meets specification requirements and (6) a statement certifying that the transport vehicle was inspected before loading and was found acceptable for the material shipped.

11. SPLIT SAMPLE TESTING

11.1 The Agency or the supplier may request split sample testing. The test results will be provided immediately to both parties.

11.2 If the split sample test data is not within the precision specified for that particular test, a review of both sampling and testing procedures will be conducted by both the supplier and the agency.

12. DECERTIFICATION

12.1 A supplier may have its authorization to certify and supply a Specific PGAB revoked by the Agency if it is found not to conform to the specifications and standards as established under this standard.

12.2 The following criteria shall be used to judge the conditions of non-conformance:

12.2.1 Failure to control the quality of the PGAB by failing to follow the procedures described in the Supplier’s approved quality control plan as required under Section 8.4.

12.2.2 Failure to cease shipment of PGAB as required under Section 9.2 when a test result indicates that the PGAB is not in compliance with the agency specifications.

12.3 A Supplier that has had its authorization to certify and supply revoked partially or entirely may seek reinstatement by demonstrating conformance to Agency certification criteria.
WSDOT FOP for AASHTO T 2¹

Standard Practice for Sampling Aggregates

1. Scope

1.1 This practice covers sampling of coarse and fine aggregates for the following purposes:
   
   1.1.1 Preliminary investigation of the potential source of supply,

   1.1.2 Control of the product at the source of supply,

   1.1.3 Control of the operations at the site of use, and

   1.1.4 Acceptance or rejection of the materials.

   *Note 1:* Sampling plans and acceptance and control tests vary with the type of construction in which the material is used. Attention is directed to Practices E 105 and D 3665.

1.2 The values stated in English inch-pounds units are to be regarded as the standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 AASHTO Standards:

   T 248 Reducing Samples of Aggregate to Testing Size

2.2 ASTM Standards:

   C 702 Practice for Reducing Field Samples of Aggregate to Testing Size
   D 2234 Test Method for Collection of a Gross Sample of Coal
   D 3665 Practice for Random Sampling of Construction Materials
   E 105 Practice for Probability Sampling of Materials
   E 122 Practice for Choice of Sample Size to Estimate the Average Quality of a Lot or Process
   E 141 Practice for Acceptance of Evidence Based on the Results of Probability Sampling

3. Significance and Use

3.1 Sampling is equally as important as the testing, and the sampler shall use every precaution to obtain samples that will show the nature and condition of the materials which they represent.

3.2 When sampling of aggregate sources for preliminary testing, the sampling must be witnessed or taken by a designated representative of the Regional Materials Engineer. The Acceptance samples will be taken by a qualified tester employed by the contracting agency or their designated qualified representative.

   Samples for preliminary investigation tests are obtained by the agency guidelines (Note 2). Samples of materials for control of the production at the source or control of the work at the site of use are obtained by the manufacturer, contractor, or other parties responsible for accomplishing the work. Samples for tests to be used in acceptance or rejection decisions by the purchaser are obtained by the purchaser or his authorized representative.

This Procedure is based on AASHTO T 2-91 (2000).
Note 2: The preliminary investigation and sampling of potential aggregate sources and types occupies a very important place in determining the availability and suitability of the largest single constituent entering into the construction. It influences the type of construction from the standpoint of economics and governs the necessary material control to ensure durability of the resulting structure, from the aggregate standpoint. This investigation should be done only by agency guidelines a responsible trained and experienced person. For more comprehensive guidance, see the Appendix.

4. SECURING SAMPLES

4.1 General — Where practicable, samples to be tested for quality shall be obtained from the finished product. Samples from the finished product to be tested for abrasion loss shall not be subject to further crushing or manual reduction in particle size in preparation for the abrasion test unless the size of the finished product is such that it requires further reduction for testing purposes.

Native soils within the contract limits to be used for embankment construction and/or backfill material do not require the sampling by a qualified tester. For material that requires gradation testing such as but not limited to manufactured aggregates and Gravel Borrow, a qualified tester shall be required for sampling.

4.2 Inspection — The material shall be inspected to determine discernible variations. The seller shall provide suitable equipment needed for proper inspection and sampling.

4.3 Procedure

4.3.1 Sampling from a Flowing Aggregate Stream (Bins or Belt Discharge) — Select units to be sampled by a random method, such as Practice D3665, from the production. Obtain at least three approximately equal increments, selected at random from the unit being sampled, and combine to form a field sample whose mass equals or exceeds the minimum recommended in 4.4.2. Take the sample each increment from the entire cross section of the material as it is being discharged. The Standard Specifications require an mechanical, automatic or semi-automatic sampling device be used for processed materials. It is usually necessary to have a special device constructed for use at each plant. This device consists of a pan of sufficient size to intercept the entire cross section of the discharge stream and hold the required quantity of material without overflowing. A set of rails may be necessary to support the pan as it is passed under the discharge stream. Insofar as is possible, keep bins continuously full or nearly full to reduce segregation.

Note 3: Sampling the initial discharge or the final few tons from a bin or conveyor belt increases the chances of obtaining segregated material and should be avoided.

4.3.2 Sampling from the Conveyor Belt (Stopped) — Select units to be sampled by a random method, such as Practice D3665, from the production. Obtain a field sample at least three approximately equal increments, selected at random, from the unit being sampled and combine to form a field sample whose mass equals or exceeds the minimum recommended in 4.4.2. Stop the conveyor belt while the sample increments are being obtained. Insert two templates, the shape of which conforms to the shape of the belt in the aggregate stream on the belt, and space them such that the material contained between them will yield an increment of the required weight. Carefully scoop all material between the templates into a suitable container and collect the fines on the belt with a brush and dust pan and add to the container.
4.3.3 Sampling from Stockpiles or Transportation Units — Avoid sampling coarse aggregate or mixed coarse and fine aggregate from stockpiles or transportation units whenever possible, particularly when the sampling is done for the purpose of determining aggregate properties that may be dependent upon the grading of the sample. If circumstances make it necessary to obtain samples from a stockpile of coarse aggregate or a stockpile of combined coarse and fine aggregate, design a sampling plan for the specific case under consideration. This approach will allow the sampling agency to use a sampling plan that will give a confidence in results obtained there from that is agreed upon by all parties concerned to be acceptable for the particular situation. The sampling plan shall define the number of samples necessary to represent lots and sublots of specific sizes. General principles for sampling from stockpiles are applicable to sampling from trucks, rail cars, barges or other transportation units. For general guidance in sampling from stockpiles, see the Appendix.

4.3.4 Sampling from Roadway (Bases and Subbases) — WSDOT has deleted this section.

4.4 Number and Masses of Field Samples

4.4.1 The number of field samples (obtained by one of the methods described in 4.3) required depends on the criticality of, and variation in, the properties to be measured. Designate each unit from which a field sample is to be obtained prior to sampling. The number of field samples from the production should be sufficient to give the desired confidence in test results.

Note 4: Guidance for determining the number of samples required to obtain the desired level of confidence in test results may be found in Test Method D 2234, Practice E 105, Practice E 122, and Practice E 141.

4.4.2 The field sample masses cited are tentative. The masses must be predicated on the type and number of tests to which the material is to be subjected and sufficient material obtained to provide for the proper execution of these tests. Standard acceptance and control tests are covered by ASTM standards and specify the portion of the field sample required for each specific test. Generally speaking, the amounts specified in Table 1 will provide adequate material for routine grading and quality analysis. Extract test portions from the field sample according to T 248 or as required by other applicable test methods.

5. SHIPPING SAMPLES

5.1 Transport aggregates in bags or other containers so constructed as to preclude loss or contamination of any part of the sample, or damage to the contents from mishandling during shipment. The weight limit for each bag of aggregate is 30 pounds maximum.

5.2 Shipping containers for aggregate samples shall have suitable individual identification attached and enclosed so that field reporting, laboratory logging, and test reporting may be facilitated. All samples submitted for testing to the Region or State Materials Laboratories shall be accompanied by completed sample transmittal (WSDOT Form 350-056) or equivalent.
### Table 1

**Size of Samples**

<table>
<thead>
<tr>
<th>Maximum Nominal Size of Aggregates</th>
<th>Approximate Minimum Mass of Field Samples, kg</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Fine Aggregate</strong></td>
<td></td>
</tr>
<tr>
<td>2.36 mm</td>
<td>10</td>
</tr>
<tr>
<td>4.75 mm</td>
<td>10</td>
</tr>
<tr>
<td><strong>Coarse Aggregate</strong></td>
<td></td>
</tr>
<tr>
<td>9.5 mm</td>
<td>10</td>
</tr>
<tr>
<td>12.5 mm</td>
<td>15</td>
</tr>
<tr>
<td>19.0 mm</td>
<td>25</td>
</tr>
<tr>
<td>25.0 mm</td>
<td>50</td>
</tr>
<tr>
<td>37.5 mm</td>
<td>75</td>
</tr>
<tr>
<td>50 mm</td>
<td>100</td>
</tr>
<tr>
<td>63 mm</td>
<td>125</td>
</tr>
<tr>
<td>75 mm</td>
<td>150</td>
</tr>
<tr>
<td>90 mm</td>
<td>175</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Nominal Maximum Size A* in (mm)</th>
<th>Minimum Mass B lb (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>US No. 4 (4.75)</td>
<td>5 (2)</td>
</tr>
<tr>
<td>1/4 (6.3)</td>
<td>10 (4)</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td>10 (4)</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>20 (8)</td>
</tr>
<tr>
<td>5/8 (16.0)</td>
<td>20 (8)</td>
</tr>
<tr>
<td>3/4 (19.0)</td>
<td>30 (12)</td>
</tr>
<tr>
<td>1 (25.0)</td>
<td>55 (25)</td>
</tr>
<tr>
<td>1 1/4 (31.5)</td>
<td>70 (30)</td>
</tr>
<tr>
<td>1 1/2 (37.5)</td>
<td>80 (36)</td>
</tr>
<tr>
<td>2 (50)</td>
<td>90 (40)</td>
</tr>
<tr>
<td>2 1/2 (63)</td>
<td>110 (50)</td>
</tr>
<tr>
<td>3 (75)</td>
<td>140 (60)</td>
</tr>
<tr>
<td>3 1/2 (90)</td>
<td>180 (80)</td>
</tr>
</tbody>
</table>

*For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.*

*For processed aggregate the nominal maximum size of particles is the largest sieve size listed in the applicable specification, upon which any material is permitted to be retained.*

*For combined coarse and fine aggregates (for example, base or subbase) minimum weight shall be coarse aggregate minimum plus 10 kg.*

**Note 5:** For an aggregate specification having a generally unrestrictive gradation (i.e. wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.
APPENDIXES

XI. SAMPLING AGGREGATE FROM STOCKPILES OR TRANSPORTATION UNITS

X1.1 Scope

X1.1.1 In some situations it is mandatory to sample aggregates that have been stored in stockpiles or loaded into rail cars, barges, or trucks. In such cases the procedure should ensure that segregation does not introduce a serious bias in the results.

X1.2 Sampling From Stockpiles

X1.2.1 In sampling material from stockpiles it is very difficult to ensure unbiased samples, due to the segregation which often occurs when material is stockpiles, with coarser particles rolling to the outside base of the pile. For coarse or mixed coarse and fine aggregate, every effort should be made to enlist the services of power equipment, such as a front end loader, to develop a separate, small sampling pile composed of materials drawn from various levels and locations in the main pile after which several increments may be combined to compose the field sample. If necessary to indicate the degree of variability existing within the main pile, separate samples should be drawn from separate areas of the pile.

X1.2.2 Where power equipment is not available, samples from stockpiles should be made up of at least three increments taken from the top third, at the mid-point, and at the bottom third of the volume of the pile. A board shoved vertically into the pile just above the sampling point aids in preventing further segregation. In sampling stockpiles of fine aggregate the outer layer, which may have become segregated, should be removed and the sample taken from the material beneath. Sampling tubes approximately 1 ½ in. (30-mm) min by 6 ft. (2-m) min in length may be inserted into the pile at random locations to extract a minimum of five increments of material to form the sample.

X1.3 Sampling From Transportation Units

X1.3.1 In sampling coarse aggregates from railroad cars or barges, effort should be made to enlist the services of power equipment capable of exposing the material at various levels and random locations. Where power equipment is not available, a common procedure requires excavation of three or more trenches across the unit at points that will, from visual appearance, give a reasonable estimate of the characteristics of the load. The trench bottom should be approximately level, at least 1 ft. (0.3 m) in width and in depth below the surface. A minimum of three increments from approximately equally spaced points along each trench should be taken by pushing a shovel downward into the material. Coarse aggregate in trucks should be sampled in essentially the same manner as for rail car or barges, except for adjusting the number of increments according to the size of the truck. For fine aggregate in transportation units, sampling tubes as described in X1.2 may be used to extract an appropriate number of increments to form the sample.
X2. EXPLORATION OF POTENTIAL AGGREGATE SOURCES

X2.1 Scope

X2.1.1 Sampling for evaluation of potential aggregate sources should be performed by a responsible trained and experienced person. Because of the wide variety of conditions under which sampling may have to be done it is not possible to describe detailed procedures applicable to all circumstances. This appendix is intended to provide general guidance and list more comprehensive references.

X2.2 Sampling Stone from Quarries of Ledges

X2.2.1 Inspection — The ledge or quarry face should be inspected to determine discernible variations or strata. Differences in color and structure should be recorded.

X2.2.2 Sampling and Size of Sample — Separate samples having a mass of at least 55 lbs (25 kg) should be obtained from each discernible stratum. The sample should not include material weathered to such an extent that it is no longer suitable for the purpose intended. One or more pieces in each sample should be at least 6 X 6 X 4 inch (150 by 150 by 100 mm) in size with the bedding plane plainly marked, and this piece should be free of seams or fractures.

X2.2.3 Record — In addition to the general information accompanying all samples the following information should accompany samples taken from ledges or quarry faces:

X2.2.3.1 Approximate quantity available. (If quantities is very large this may be recorded as practically unlimited.)

X2.2.3.2 Quantity and character of overburden.

X2.2.3.3 A detailed record showing boundaries and location of material represented by each sample.

Note X2.1: A sketch, plan, and elevation, showing the thickness and location of the different layers is recommended for this purpose.

X2.3 Sampling Roadside or Bank Run Sand and Gravel Deposits

X2.3.1 Inspection — Potential sources of bank run sand and gravel may include previously worked pits from which there is an exposed face or potential deposits discovered through air-photo interpretation, geophysical exploration, or other types of terrain investigation.

X2.3.2 Sampling — Samples should be so chosen from each different stratum in the deposit discernible to the sampler. An estimate of the quantity of the different materials should be made. If the deposit is worked as an open-face bank or pit, samples should be taken by channeling the face vertically, bottom to top, so as to represent the materials proposed for use. Overburdened or disturbed material should not be included in the sample. Test holes should be excavated or drilled at numerous locations in the deposit to determine the quality of the material and the extent of the deposit beyond the exposed face, if any. The number and depth of test holes will depend upon the quantity of the material needed, topography of the area, nature of the deposit, character of the material, and potential value of the material in the deposit. If visual inspection indicates that there is considerable variation in the material, individual samples should be selected from the material in each well defined stratum. Each sample should be thoroughly mixed and quartered if necessary so that the field sample thus obtained will be at least 25 lb (12 kg) for sand and 75 lb (35 kg) if the deposit contains an appreciable amount of coarse aggregate.
X2.3.3 Record — In addition to the general information accompanying all samples the following information should accompany samples of bank run sand and gravel:

X2.3.3.1 Location of supply.
X2.3.3.2 Estimate of approximate quantity available.
X2.3.3.3 Quantity and character of overburden.
X2.3.3.4 Length of haul to proposed site of work.
X2.3.3.5 Character of haul (kind of road, maximum grades, etc.)
X2.3.3.6 Details as to extent and location of material represented by each sample.
Performance Exam Checklist

Sampling of Aggregates
FOP for AASHTO T 2

Participant Name ________________________________ Exam Date ______________

Procedure Element

1. The tester has a copy of the current procedure on hand?  Yes ☐ No ☐

Conveyor Belts – Stopped

2. Belt stopped? ☐ ☐
3. Sampling device set on belt, avoiding intrusion of adjacent material? ☐ ☐
4. Sample, including all fines, scooped off? ☐ ☐

Flowing Aggregate Sampler

5. Container passed through full stream of material as it runs off end of belt? (Mechanical, Automatic or Semi Automatic Sampler Only) ☐ ☐ ☐

Transport Units

6. Three or more trenches cut across the unit? ☐ ☐
7. Trench bottom level and approximate 1 foot wide and 1 foot below surface of material in unit? ☐ ☐
8. Three samples taken at equal spacing along each trench? ☐ ☐

Stockpiles

9. Create vertical face, if one does not exist, or use mechanical equipment to build a small sampling pile? ☐ ☐
10. At least three increments taken, at various locations? ☐ ☐

Procedure Element

11. If vertical face cannot be created, increment taken from at least three locations from top, middle, and bottom? ☐ ☐
12. When sampling sand, outer layer removed and increments taken from at least five locations? ☐ ☐

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________
SAMPLING FRESHLY MIXED CONCRETE
FOP FOR WAQTC TM 2

SIGNIFICANCE

Testing fresh concrete in the field begins with obtaining and preparing the sample to be tested. Standardized procedures for obtaining a representative sample from various types of mixing and/or agitating equipment have been established. Specific time limits regarding when tests for temperature, slump, and air content must be started and for when the molding of test specimens must begin are also established.

Technicians must be patient and refrain from obtaining the sample too quickly. Doing so would be a violation of the specifications under which the concrete is being supplied and it may result in a nonrepresentative sample of concrete. If one considers that the specifications may require strength tests to be made only once every 50 cy (40 m³), the need for a truly representative sample is apparent. The minimum 1 ft³ (0.03 m³) sample from which the compressive strength test specimens will be made represents only 0.07 to 0.08 percent of the total quantity of concrete placed. For this reason, every precaution must be taken to obtain a sample that is truly representative of the entire batch and then to protect that sample from the effects of evaporation, contamination, and physical damage.

SCOPE

This procedure provides instruction for obtaining samples of fresh concrete in accordance with WAQTC TM 2. Sources covered include stationary and paving mixers, revolving drum truck mixers or agitators, open-top truck mixers and the discharge of pump or conveyor placement systems.

APPARATUS

- Wheelbarrow
- Cover for wheelbarrow (plastic, canvas, or burlap)
- Shovel
- 5 gal (19 L) bucket for water

PROCEDURE

Use every precaution in order to obtain samples representative of the true nature and condition of the concrete being placed being careful not to obtain samples form the very first or very last portions of the batch. The size of the sample will be 1.5 times the volume of concrete required for the specified testing, but not less than 1 ft³ (0.03 m³) after wet-sieving, if required.

RANDOM SAMPLE SELECTION

Concrete samples other than initial load samples or samples for questioned acceptance will be taken from each subplot by a random selection. Sublots are determined by the designated sampling frequency in the Standard Specifications. Random selection will be accomplished by using WSDOT Test Method T716, Method of Random Sampling for Locations of Testing and Sampling Sites.
• **Sampling from stationary mixers, except paving mixers**

Sample the concrete after a minimum of 1/2 yd$^3$ (1/2 m$^3$) of concrete has been discharged. Perform sampling by passing a receptacle completely through the discharge stream, or by completely diverting the discharge into a sample container. If discharge of the concrete is too rapid to divert the complete discharge stream, discharge the concrete into a container or transportation unit sufficiently large to accommodate the entire batch and then accomplish the sampling in the same manner as given for paving mixers. Take care not to restrict the flow of concrete from the mixer, container, or transportation unit so as to cause segregation. These requirements apply to both tilting and nontilting mixers.

• **Sampling from paving mixers**

Sample after the contents of the paving mixer have been discharged. Obtain material from at least five different locations in the pile and combine into one test sample. Avoid contamination with subgrade material or prolonged contact with absorptive subgrade. To preclude contamination or absorption by the subgrade, sample the concrete by placing a shallow container on the subgrade and discharging the concrete across the container. The container shall be of a size sufficient to provide a sample size that is in agreement with the nominal maximum aggregate size.

• **Sampling from revolving drum truck mixers or agitators**

Sample the concrete after a minimum of 1/2 yd$^3$ (1/2 m$^3$) of concrete has been discharged. Do not obtain samples until after all of the water has been added to the mixer. Do not obtain samples from the very first or last portions of the batch discharge. Sample by repeatedly passing a receptacle through the entire discharge stream or by completely diverting the discharge into a sample container. Regulate the rate of discharge of the batch by the rate of revolution of the drum and not by the size of the gate opening.

• **Sampling from open-top truck mixers, agitators, non-agitating equipment or other types of open-top containers**

Sample by whichever of the procedures described above is most applicable under the given conditions.

• **Sampling from pump or conveyor placement systems**

Sample after a minimum of 1/2 yd$^3$ (1/2 m$^3$) of concrete has been discharged. Do not obtain samples until after all of the pump slurry has been eliminated. Sample by repeatedly passing a receptacle through the entire discharge system or by completely diverting the discharge into a sample container. Do not lower the pump arm from the placement position to ground level for ease of sampling, as it may modify the air content of the concrete being sampled. Do not obtain samples from the very first or last portions of the batch discharge.

Transport samples to the place where fresh concrete tests are to be performed and specimens are to be molded.

Combine and remix the sample minimum amount necessary to ensure uniformity. Protect the sample from direct sunlight, wind, rain, and sources of contamination.

Complete test for temperature and start tests for slump and air content within 5 minutes of obtaining the sample. Complete tests as expeditiously as possible. Start molding specimens for strength tests within 15 minutes of obtaining the sample.
WET SIEVING

When required for slump testing, air content testing or molding test specimens, the concrete sample shall be wet-sieved, prior to remixing, by the following:

1. Place the sieve designated by the test procedure over dampened sample container.

2. Pass the concrete over the designated sieve. Do not overload the sieve (one particle thick.)

3. Shake or vibrate the sieve until no more material passes the sieve.

4. Discard oversize material including all adherent mortar.

5. Repeat until sample of sufficient size is obtained.

6. Mortar adhering to the wet-sieving equipment shall be included with the sample.

**Note**: Wet-sieving is not allowed for samples being utilized for density determinations according to the FOP for AASHTO T 121.
Performance Exam Checklist

Sampling Freshly Mixed Concrete
FOP for WAQTC TM 2

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Obtain a representative sample:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Sample the concrete after 1/2 cy (1/2 m³) discharged?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>b. Pass receptacle through entire discharge stream or completely divert discharge stream into sampling container?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>c. Transport samples to place of testing?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>d. Sample remixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>e. Sample protected?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>f. Minimum size of sample used for strength tests 1 ft³ (0.03 m³)?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Start tests for slump and air within 5 minutes of sample being obtained?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Start molding cylinders within 15 minutes of sample being obtained?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Protect sample against rapid evaporation and contamination?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐  Fail ☐  Second attempt: Pass ☐  Fail ☐

Signature of Examiner __________________________________________

This checklist is derived, in part, from copyrighted material printed in ACI CP-1, published by the American Concrete Institute.

Comments:

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
IN-PLACE DENSITY OF HOT MIX ASPHALT USING THE NUCLEAR MOISTURE-DENSITY GAUGE
FOP FOR WAQTC TM 8

Significance

The final in-place density of roadway pavement is critical to the quality and longevity of a highway project. Low density material will lead to excessive deflection under load and/or permanent deformation.

This procedure provides a rapid, nondestructive technique for determining the in-place density of compacted bituminous mixes. It can be used to establish the proper rolling effort and pattern to achieve the required density. The non-destructive nature of the test allows repetitive measurements to be made at a single test location between roller passes.

Scope

This test method describes a test procedure for determining the density of Hot Mix Asphalt (HMA) by means of a nuclear gauge employing either direct transmission or backscatter methods. Correlation with densities determined under the FOP for AASHTO T 166 is required by some agencies.

This test method involves potentially hazardous materials, operations and equipment. This standard does not purport to address all of the safety problems, 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. See Hazards. All operators will be trained in radiation safety prior to operating nuclear density gauges. Some agencies WSDOT requires the use of personal monitoring devices such as a thermoluminescent dosimeter or film badge which is monitored on a quarterly basis.

REFERENCE DOCUMENTS

AASHTO Standards

- T 166 Standard Method of Test for Bulk Specific Gravity of Compacted Hot-Mix Asphalt Using Saturated Surface-Dry Specimens
- T168 Standard Method of Test for Sampling Bituminous Paving Mixtures
- T 209 Standard Method of Test for Theoretical Maximum Specific Gravity and Density of Hot-Mix Asphalt
- Paving Mixtures
  T 275 Standard Method of Test for Bulk Specific Gravity of Compacted Bituminous Mixtures Using Paraffin-Coated Specimens

ASTM Standards:

- D 1188 Test Method for Bulk Specific Gravity and Density of Compacted Bituminous Mixtures Using Paraffin-Coated Specimens
- D 2041 Test Method for Theoretical Maximum Specific Gravity of Bituminous Paving Mixtures
• D 2726 Test Method for Bulk Specific Gravity and Density of Non-Absorptive Compacted Bituminous Mixtures
• D 3665 Practice for Random Sampling of Construction Materials

WSDOT Standards
• FOP for WAQTC/AASHTO T 166 Standard Method of Test for Bulk Specific Gravity of Compacted Hot-Mix Asphalt Using Saturated Surface-Dry Specimens
• FOP for AASHTO T 168 Standard Method of Test for Sampling Bituminous Paving Mixtures
• FOP for AASHTO T 209 Standard Method of Test for Theoretical Maximum Specific Gravity and Density of Hot-Mix Asphalt Paving Mixtures
• SOP 729 In Place Density of Bituminous Mixes Using the Nuclear Moisture-Density Gauge FOP for WAQTC TM 8
• SOP 730 Correlation of Nuclear Gauge Densities with Hot Mix Asphalt (HMA) Cores
• SOP 733 Determination of Pavement Density Differentials Using the Nuclear Density Gauge
• SOP 735 Standard Operating Procedure for Longitudinal Joint Density

Radiation Safety
This method does not purport to address all of the safety problems associated with its use. This test method involves potentially hazardous materials. The gauge utilizes radioactive materials that may be hazardous to the health of the user unless proper precautions are taken. Users of this gauge must become familiar with the applicable safety procedures and governmental regulations. All operators will be trained in radiation safety prior to operating nuclear density gauges. Some agencies require the use of personal monitoring devices such as a thermoluminescent dosimeter or film badge. Effective instructions together with routine safety procedures such as source leak tests, recording and evaluation of personal monitoring device data, etc., are a recommended part of the operation and storage of this gauge.

Apparatus
• Nuclear density gauge with the factory matched standard reference block.
• Drive pin, guide, scraper plate, and hammer for testing in direct transmission mode.
• Transport case for properly shipping and housing the gauge and tools.
• Instruction manual for the specific make and model of gauge.
• Radioactive materials information and calibration packet containing:
  – Daily Standard Count Log
  – Factory and/or Laboratory Calibration Data Sheet
  – Leak Test Certificate
  – Shippers Declaration for Dangerous Goods
  – Procedure Memo for Storing, Transporting and Handling Nuclear Testing Equipment
  – Other radioactive materials documentation as required by local regulatory requirements.
Material

- **Filler material:** Fine graded sand from the source used to produce the asphalt pavement or other agency approved materials.

Calibration

1. **WSDOT has deleted this section. WSDOT performs calibrations according to the manufacturer’s Operators Manual.**

Standardization

1. Turn the gauge on and allow it to stabilize (approximately 10 to 20 minutes) prior to standardization. Leave the power on during the day’s testing.

2. Standardize the nuclear gauge at the construction site at the start of each day’s work and as often as deemed necessary by the operator or agency. Daily variations in standard count shall not exceed the daily variations established by the manufacturer of the gauge. If the daily variations are exceeded after repeating the standardization procedure, the gauge should be repaired and or recalibrated.

3. Record the standard count for both density and moisture in the Daily Standard Count Log. The exact procedure for standard count is listed in the manufacturer’s Operators Manual.

Test Site Location

1. Select a test location(s) randomly and in accordance with WSDOT Test Method T 716. Test sites should be relatively smooth and flat and meet the following conditions:
   a. At least 33 30 ft (10 m) away from other sources of radioactivity
   b. At least 10 ft (3 m) away from large objects
   c. No closer than 18 in. (450 mm) 600 mm (24 in.) to any vertical mass, or less than 18 in. (450 mm) 300 mm (12 in.) from a vertical pavement edge.

Overview

There are two methods for determining in-place density of HMA. See agency requirements for method selection.

- **Direct Transmission** - The standard for WSDOT is to run density tests in “Direct Transmission mode.”

- **Backscatter** - When the depth of Hot Mix Asphalt is less than 0.11 foot or when the driving of the drive pin is not possible to achieve the required depth for the gauge probe (i.e., underlying concrete) then a “Thin Lift Density gauge” or a Moisture Density Gauge in the “Thin Layer mode” will be allowed.
Procedure

Direct Transmission Mode

1. **Maintain maximum contact between the base of the gauge and the surface of the material under test.** Maintaining maximum contact between the base of the gauge and the surface of the material under test is critical. *This mode can not be used where the depth of the Hot Mix Asphalt (HMA) is less than 33 mm (0.11').*
   a. Use the guide and scraper plate as a template and drill a hole to a depth of at least 1/4 in. (7 mm) deeper than the measurement depth required for the gauge.
   b. Place the gauge on the prepared surface so the source rod can enter the hole. Insert the probe in the hole and lower the source rod to the desired test depth using the handle and trigger mechanism. Position the gauge with the long axis of the gauge parallel to the direction of paving. Pull the gauge by pulling it towards the sealer/detector so that the probe is firmly against the side of the hole.

   **WSDOT Note:** For alignment purposes, the user may expose the source rod for a maximum of ten seconds.

   c. Take two one four-minute tests and record the wet density (WD) readings. Rotate the gauge 90 degrees. Reset the gauge by gently moving it side to side while pulling back it towards the sealer / detector so that the probe is firmly against the side of the hole. Take another one-minute test and record the wet density reading. If the two density readings are not within 3 lbs/ft³ (50 kg/m³), rotate the gauge 180 degrees and repeat the test in the same hole until they disagree. If not, then rotate the gauge another 90 degrees. Reset the gauge by gently moving it side to side while pulling it towards the sealer / detector so that the probe is firmly against the side of the hole. Then take another one-minute test and record the wet density reading. Again compare the readings. If difference between the second and third one minute tests is not less than 50 kg/m³ (3 lb/ft³) move to another test location.

Backscatter (THIN LIFT) MODE

**WSDOT has removed this section and replaced it with the following**

a. Place the gauge on the test site and extend the probe to the backscatter position.

b. Take tests in accordance with manufacturer’s recommendation. Contact the materials laboratory for direction.

Calculation of Results

**See WSDOT SOP 729 to determine the percent compaction. It should be stressed that the numbers obtained with the nuclear gauge are simply in-place densities and tell the operator nothing in regard to relative compaction. In-place densities are to be compared with theoretical maximum density as determined by the FOP for AASHTO T 209.**

The density reported for each test site shall be the average of the two individual one-minute tests.

Percent compaction is determined by comparing the in-place wet density as determined by this method to the appropriate agency density standard. See appropriate agency policy for use of density standards.
Correlation with Cores

WSDOT has deleted this section, refer to WSDOT SOP 730.

Report

Report the test results for each subplot on WSDOT Form 350-092 or other report approved by the State Materials Engineer.

Results shall be reported on standard forms approved by the agency. Include the following information:

• Location of test and thickness of layer tested
• Mixture type
• Make, model and serial number of the nuclear moisture-density gauge
• Mode of measurement, depth, calculated wet density of each measurement and any adjustment data
• Standard density
• Percent compaction and/or percent air voids
• Name and signature of operator
Tester Qualification Practical Exam Checklist

In-place Density of Hot Mix Asphalt (HMA) Using the Nuclear Moisture-Density Gauge FOP for WAQTC TM 8

Participant Name __________________________________________ Exam Date ______________

Procedure Element Yes No
1. The tester has a copy of the current procedure on hand? ☐ ☐
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? ☐ ☐
3. Gauge turned on? ☐ ☐
4. Gauge calibrated and standard count recorded? ☐ ☐
5. Test location selected appropriately? ☐ ☐
6. Direct Transmission Mode:
   a. Hole made a minimum of ¼ inch deeper than measurement depth? ☐ ☐
   b. Gauge placed parallel to direction of paving, probe extended, gauge pulled back so probe against hole? ☐ ☐
   c. For alignment purposes did not expose the source rod for more than 10 seconds.
   d. Two one-four-minute test made? ☐ ☐
   e. Wet density recorded is averaged? ☐ ☐
7. Backscatter Mode (Thin Lift):
   a. Gauge placed, probe extended to backscatter position? ☐ ☐
   b. 4-one minute test made; gauge placed as described in the manufacture recommendations? ☐ ☐
   c. Wet Densities averaged? ☐ ☐
8. If difference greater than 3 lb/ft³, retest made? ☐ ☐
9. All calculations performed correctly? ☐ ☐
10. Nuclear Gauge secured in a manner consistent with current DOH requirements? ☐ ☐

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________
Comments:

_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
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_________________________________________________________________________________
_________________________________________________________________________________
WSDOT FOP for AASHTO T 22¹

Compressive Strength of Cylindrical Concrete Specimens

1. SCOPE

1.1 This test method covers determination of compressive strength of cylindrical concrete specimens such as molded cylinders and drilled cores. It is limited to concrete having a unit weight in excess of 50 lb/ft³ [800 kg/m³].

1.2 The values stated in English units are the standard.

1.3 This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety problems associated with its use. It is the responsibility of whoever uses this standard to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

Warning: Means should be provided to contain concrete fragments during sudden rupture of specimens. Tendency for sudden rupture increases with increasing concrete strength. (Note 1)

Note 1 - The safety precautions given in the Manual of Aggregate and Concrete Testing, located in the Related Materials section of Volume 04.02 of the Annual Book of ASTM Standards, are recommended.

1.4 The text of this standard reference notes which provide explanatory material. These notes shall not be considered as requirements of the standard.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards

T 23 Making and Curing Concrete Test Specimens in the Field
T 24 Obtaining and Testing Drilled Cores and Sawed Beams of Concrete
T 67 Standard Practices for Force Verification of Testing Machines
T 126 Making and Curing Concrete Test Specimens in the Laboratory
T 231 Capping Cylindrical Concrete Specimens

2.2 ASTM STANDARDS

C 683 Test Method for Compressive and Flexural Strength of Concrete Under Field Conditions
C 873 Test Method for Compressive Strength of Concrete Cylinders Cast in Place in Cylindrical Molds
C 1231 Practice for Use of Unbonded Caps in Determination of Compressive Strength of Hardened Concrete Cylinders
E 74 Practice for Calibration of Force-Measuring Instruments for Verifying the Load Indication of Testing Machines.

¹This Test Method is based on AASHTO T 22-05
3. SUMMARY OF TEST METHOD

3.1 This test method consists of applying a compressive axial load to molded cylinders or cores at a rate which is within a prescribed range until failure occurs. The compressive strength of the specimen is calculated by dividing the maximum load attained during the test by the cross-sectional area of the specimen.

4. SIGNIFICANCE AND USE

4.1 Care must be exercised in the interpretation of the significance of compressive strength determinations by this test method since strength is not a fundamental or intrinsic property of concrete made from given materials. Values obtained will depend on the size and shape of the specimen, batching, mixing procedures, the methods of sampling, molding, and fabrication and the age, temperature, and moisture conditions during curing.

4.2 This test method is used to determine compressive strength of cylindrical specimens prepared and cured in accordance with Methods T23, T24, T126, T231, and ASTM C873.

4.3 The results of this test method are used as a basis for quality control of concrete proportioning, mixing, and placing operations; determination of compliance with specifications; control for evaluating effectiveness of admixtures and similar uses.

5. APPARATUS

5.1 Testing Machine — The testing machine shall be of a type having sufficient capacity and capable of providing the rates of loading prescribed in Section 7.5. As a minimum, the capacity should be capable of achieving 170% of the design strength.

5.1.1 Verify calibration of the testing machines in accordance with Method T 67 except that the verified loading range shall be as required in 5.3.2. Verification is required under the following conditions:

5.1.1.1 At least annually, but not to exceed 13 months;

5.1.1.2 On original installation or immediately after relocation;

5.1.1.3 Immediately after making repairs or adjustments that affect the operation of the force applying system or the values displayed on the load indicating system, except for zero adjustments that compensate for the mass (weight) of tooling, or specimen, or both; or

5.1.1.4 Whenever there is reason to suspect the accuracy of the indicated loads.

5.1.2 Design — The design of the machine must include the following features:

5.1.2.1 The machine must be power operated and must apply the load continuously rather than intermittently, and without shock. If it has only one loading rate (meeting the requirements of Section 7.5), it must be provided with a supplemental means for loading at a rate suitable for verification. This supplemental means of loading may be power or hand operated.

Note 1 — High-strength concrete cylinders rupture more intensely than normal strength cylinders. As a safety precaution, it is recommended that the testing machines should be equipped with protective fragment guards.

5.1.2.2 The space provided for test specimens shall be large enough to accommodate, in a readable position, an elastic calibration device which is of sufficient capacity to cover the potential loading range of the testing machine and which complies with the requirements of Practice E 74.
Note 2: The types of elastic calibration devices most generally available and most commonly used for this purpose are the circular proving ring or load cell.

5.1.3 Accuracy — The accuracy of the testing machine shall be in accordance with the following provisions:

5.1.3.1 The percentage of error for the loads within the proposed range of use of the testing machine shall not exceed ± 1.0 % of the indicated load.

5.1.3.2 The accuracy of the testing machine shall be verified by applying five test loads in four approximately equal increments in ascending order. The difference between any two successive test loads shall not exceed one third of the difference between the maximum and minimum test loads.

5.1.3.3 The test load as indicated by the testing machine and the applied load computed from the readings of the verification device shall be recorded at each test point. Calculate the error, E, and the percentage of error, Ep, for each point from these data as follows:

\[ E = A - B \]
\[ Ep = \frac{100(A - B)}{B} \]

where:
\( A \) = load, lbf [kN] indicated by the machine being verified, and
\( B \) = applied load, lbf [kN] as determined by the calibrating device.

5.1.3.4 The report on the verification of a testing machine shall state within what loading range it was found to conform to specification requirements rather than reporting a blanket acceptance or rejection. In no case shall the loading range be stated as including loads below the value which is 100 times the smallest change of load that can be estimated on the load-indicating mechanism of the testing machine or loads within that portion of the range below 10% of the maximum range capacity.

5.1.3.5 In no case shall the loading range be stated as including loads outside the range of loads applied during the verification test.

5.1.3.6 The indicated load of a testing machine shall not be corrected either by calculation or by the use of a calibration diagram to obtain values within the required permissible variation.

5.2 The testing machine shall be equipped with two steel bearing blocks with hardened faces (Note 3), one of which is a spherically seated block that will bear on the upper surface of the specimen, and the other a solid block on which the specimen shall rest. Bearing faces of the blocks shall have a minimum dimension at least 3 % greater than the diameter of the specimen to be tested. Except for the concentric circles described below, the bearing faces shall not depart from a plane by more than 0.001 in. [0.025 mm] in any 6 in. [150 mm] of blocks 6 in. [150 mm] in diameter or larger, or by more than 0.001 in. [0.025 mm] in the diameter of any smaller block; and new blocks shall be manufactured within one half of this tolerance. When the diameter of the bearing face of the spherically seated block exceeds the diameter of the specimen by more than 0.5 in. [13 mm], concentric circles not more than 0.031 in. [0.8 mm] deep and not more than 0.047 in. [1 mm] wide shall be inscribed to facilitate proper centering.

Note 3: It is desirable that the bearing faces of blocks used for compression testing of concrete have a Rockwell hardness of not less than 55 HRC.
5.2.1 Bottom bearing blocks shall conform to the following requirements:

5.2.1.1 The bottom bearing block is specified for the purpose of providing a readily machinable surface for maintenance of the specified surface conditions (Note 4). The top and bottom surfaces shall be parallel to each other. Its least horizontal dimension shall be at least 3 % greater than the diameter of the specimen to be tested. Concentric circles as described in Section 5.2 are optional on the bottom block.

**Note 4**—The block may be fastened to the platen of the testing machine.

5.2.1.2 Final centering must be made with reference to the upper spherical block when the lower bearing block is used to assist in centering the specimen. The center of the concentric rings, when provided, or the center of the block itself must be directly below the center of the spherical head. Provision shall be made on the platen of the machine to assure such a position.

5.2.1.3 The bottom bearing block shall be at least 1 in. [25 mm] thick when new, and at least 0.9 in. [22.5 mm] thick after any resurfacing operations, except when the block is in full and intimate contact with the lower platen of the testing machine, the thickness may be reduced to 0.38 in. (10 mm).

**Note 5:** If the testing machine is so designed that the platen itself can be readily maintained in the specified surface condition, a bottom block is not required.

5.2.2 The spherically seated bearing block shall conform to the following requirements:

5.2.2.1 The maximum diameter of the bearing face of the suspended spherically seated block shall not exceed the values given below:

<table>
<thead>
<tr>
<th>Diameter of Test Specimens in. [mm]</th>
<th>Maximum Diameter of Bearing Face in. [mm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 [50]</td>
<td>4 [105]</td>
</tr>
<tr>
<td>3 [75]</td>
<td>5 [130]</td>
</tr>
<tr>
<td>4 [100]</td>
<td>6.5 [165]</td>
</tr>
<tr>
<td>6 [150]</td>
<td>10 [255]</td>
</tr>
<tr>
<td>8 [200]</td>
<td>11 [280]</td>
</tr>
</tbody>
</table>

**Note 6:** Square bearing faces are permissible, provided the diameter of the largest possible inscribed circle does not exceed the above diameter.
5.2.2.2 The center of the sphere shall coincide with the surface of the bearing face within a tolerance of ± 5 % of the radius of the sphere. The diameter of the sphere shall be at least 75 % of the diameter of the specimen to be tested.

5.2.2.3 The ball and the socket shall be designed so that the steel in the contact area does not permanently deform when loaded to the capacity of the test machine. (Note 7).

![Schematic Sketch of a Typical Spherical Bearing Block](image)

**Figure 1**

*Note:* Provision shall be made for holding the ball in the socket and for holding the entire unit in the testing machine.

*Note 7:* The preferred contact area is in the form of a ring (described as preferred "bearing" area) as shown on Figure 1.

5.2.2.4 The curved surfaces of the socket and of the spherical portion shall be kept clean and shall be lubricated with a petroleum-type oil such as conventional motor oil, not with a pressure type grease. After contacting the specimen and application of small initial load, further tilting of the spherically seated block is not intended and is undesirable.

5.2.2.5 If the radius of the sphere is smaller than the radius of the largest specimen to be tested, the portion of the bearing face extending beyond the sphere shall have a thickness not less than the difference between the radius of the sphere and radius of the specimen. The least dimension of the bearing face shall be at least as great as the diameter of the sphere (see Figure 1).  

5.2.2.6 The movable portion of the bearing block shall be held closely in the spherical seat, but the design shall be such that the bearing face can be rotated freely and tilted at least 4 degrees in any direction.
5.3 Load Indication:

5.3.1 If the load of a compression machine used in concrete testing is registered on a dial, the dial shall be provided with a graduated scale that is readable to at least the nearest 0.1 % of the full scale load (Note 8). The dial shall be readable within 1 % of the indicated load at any given load level within the loading range. In no case shall the loading range of a dial be considered to include loads below the value that is 100 times the smallest change of load that can be read on the scale. The scale shall be provided with a graduation line equal to zero and so numbered. The dial pointer shall be of sufficient length to reach the graduation marks; the width of the end of the pointer shall not exceed the clear distance between the smallest graduations. Each dial shall be equipped with a zero adjustment which is easily accessible from the outside of the dial case, while observing the zero mark and dial pointer, and with a suitable device that at all times until reset will indicate to within one percent accuracy the maximum load applied to the specimen.

Note 8: As close as can reasonably be read is considered to be 0.02 in. [0.5 mm] along the arc described by the end of the pointer. Also, one half of a scale interval is close as can reasonably be read when the spacing on the load indicating mechanism is between 0.04 in. [1 mm] and 0.06 in. [2 mm]. When the spacing is between 0.06 and 0.12 in. [2 and 3 mm], one third of a scale interval can be read with reasonable certainty. When the spacing is 0.12 in. [3 mm] or more, one fourth of a scale interval can be read with reasonable certainty.

5.3.2 If the testing machine load is indicated in digital form, the numerical display must be large enough to be easily read. The numerical increment must be equal to or less than 0.10 % of the full scale load of a given loading range. In no case shall the verified loading range include loads less than the minimum numerical increment multiplied by 100. The accuracy of the indicated load must be within 1.0 % for any value displayed within the verified loading range. Provision must be made for adjusting to indicate true zero at zero load. There shall be provided a maximum load indicator that at all times until reset will indicate within 1.0 % system accuracy the maximum load applied to the specimen.

5.4 Provide a means for containing fragments in the event of explosive rupture of the cylinders during testing.

6. SPECIMENS

6.1 Specimens shall not be tested if any individual diameter of a cylinder differs from any other diameter of the same cylinder by more than 2 %. (Note 9).

Note 9: This may occur when single use molds are damaged or deformed during shipment, when flexible single use molds are deformed during molding or when a core drill deflects or shifts during drilling.

6.2 Neither end of compressive test specimens when tested shall depart from perpendicularity to the axis by more than 0.5 degrees (approximately equivalent to 0.12 in. in 12 in. (3 mm in 300 mm). The ends of compression test specimens that are not plane within 0.002 in. [0.050 mm] shall be sawed, or ground or capped in accordance with T 231 to meet that tolerance or capped in accordance with either T 231 or, when permitted, Practice C 1231. The diameter used for calculating the cross-sectional area of the test specimen shall be determined to the nearest 0.01 in. [0.25 mm] by averaging two diameters measured at right angles to each other at about mid-height of the specimen. The height of the cylinder shall be determined to 0.01 in.
6.3 The number of individual cylinders measured for determination of average diameter may be reduced to one for each ten specimens or three specimens per day, whichever is greater, if all cylinders are known to have been made from a single lot of reusable or single-use molds which consistently produce specimens with average diameters within a range of 0.02 in. [0.5 mm]. When the average diameters do not fall within the range 0.02 in. (0.5 mm) or when the cylinders are not made from a single lot of molds, each cylinder tested must be measured and the value used in calculation of the unit compressive strength of that specimen. When the diameters are measured at the reduced frequency, the cross-sectional areas of all cylinders tested on that day shall be computed from the average of the diameters of the three or more cylinders representing the group tested that day.

6.4 If the purchaser of the testing services requests measurement of density of the test specimen, determine the mass of the specimen before capping. Remove any surface moisture with a towel and measure the mass of the specimen using a balance or scale that is accurate to within 0.3% of the mass being measured. Measure the length of the specimen to the nearest 1 mm (0.05 in.) at three locations spaced evenly around the circumference. Compute the average length and record to the nearest 1 mm (0.05 in.). Alternatively, determine the cylinder density by weighing the cylinder in air and then submerged under water at 23.0 ±2.0 °C (73.5 ±3.5 °F), and computing the volume according to 8.3.1.

6.5 When density determination is not required and the length to diameter ratio is less than 1.8 or more than 2.2, measure the length of the specimen to the nearest 0.05 D. The length shall be measured to the nearest 0.05 D when the length to diameter ratio is less than 1.8, or more than 2.2, or when the volume of the cylinder is determined from measured dimensions.

7. PROCEDURE

7.1 Compression tests of moist-cured specimens shall be made as soon as practicable after removal from moist storage.

7.2 Test specimens shall be kept moist by any convenient method during the period between removal from moist storage and testing. They shall be tested in the moist condition.

7.3 All test specimens for a given test age shall be broken within the permissible time tolerances prescribed as follows:

```
<table>
<thead>
<tr>
<th>Test Age</th>
<th>Permissible Tolerance</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 h</td>
<td>± 0.25 h or 2.1 %</td>
</tr>
<tr>
<td>24 h</td>
<td>± 0.5 h or 2.1 %</td>
</tr>
<tr>
<td>3 days</td>
<td>2 h or 2.8 %</td>
</tr>
<tr>
<td>7 days</td>
<td>6 h or 3.6 %</td>
</tr>
<tr>
<td>28 days</td>
<td>20 h or 3.0 %</td>
</tr>
<tr>
<td>56 days</td>
<td>40 h or 3.0 %</td>
</tr>
<tr>
<td>90 days</td>
<td>2 days 2.2 %</td>
</tr>
</tbody>
</table>
```

7.4 Placing the Specimen Place the plain (lower) bearing block, with its hardened face up, on the table or platen of the testing machine directly under the spherically seated (upper) bearing block. Wipe clean the bearing faces of the upper and lower bearing blocks and of the test specimen and place the test specimen on the lower bearing block.
7.4.1. Zero Verification and Block Seating—Prior to testing the specimen, verify that the load indicator is set to zero. In cases where the indicator is not properly set to zero, adjust the indicator (Note 10). As Prior to the spherically-seated block is being brought to bear on the specimen, rotate its movable portion gently by hand so that uniform seating is obtained.

Note 10—The technique used to verify and adjust load indicator to zero will vary depending on the machine manufacturer. Consult your owner’s manual or compression machine calibrator for the proper technique.

7.5 Rate of Loading-Apply the load continuously and without shock.

7.5.1 The load shall be applied at a rate of movement (platen to crosshead measurement) corresponding to a stress rate on the specimen of 35 +/- 7 psi/s (0.25 +/- 0.05 MPa/s) (Note 11). The designated rate of movement shall be maintained at least during the latter half of the anticipated loading phase.

Note 11 – For a screw driven or displacement-controlled testing machine, preliminary testing will be necessary to establish the required rate of movement to achieve the specified stress rate. The required rate of movement will depend on the size of the test specimen, the elastic modulus of the concrete, and the stiffness of the testing machine.

7.5.2 During application of the first half of the anticipated Loading phase a higher rate of loading shall be permitted. The higher loading rate shall be applied in a controlled manner so that the specimen is not subjected to shock loading.

7.5.3 Make no adjustment in the rate of movement (platen to crosshead) as the ultimate load is being approached and the stress rate decreases due to cracking in the specimen.

7.6 Apply the compressive load until the load indicator shows that the load is decreasing steadily and the specimen displays a well-defined fracture pattern (Fig. 2). For a testing machine equipped with a specimen break detector, automatic shut-off of the testing machine is prohibited until the load has dropped to a value that is less than 95 % of the peak load. When testing with unbonded caps, a corner fracture may occur before the ultimate capacity of the specimen has been attained. Continue compressing the specimen until the user is certain that the ultimate capacity has been attained. Record the maximum load carried by the specimen during the test, and note the type of fracture pattern according to Fig. 2. If the fracture pattern is not one of the typical patterns shown in Fig. 2, sketch and describe briefly the fracture pattern. If the measured strength is lower than expected, examine the fractured concrete and note the presence of large air voids, evidence of segregation, whether fractures pass predominantly around or through the coarse aggregate particles, and verify end preparations were in accordance with Practice T 231 or Practice C1231.

Note WSDOT 1: The test loading should be stopped when 80% of the loading capacity of the testing machine has been reached. Record the maximum load achieved and note that the sample was not taken to failure as it exceeded the safe working limits of the testing machine.
8. CALCULATION

8.1 Calculate the compressive strength of the specimen by dividing the maximum load carried by the specimen during the test by the average cross-sectional area determined as described in Section 6 and express the result to the nearest 10 psi [0.1 MPa].

8.2 If the specimen length to diameter ratio is 1.75 or less, correct the result obtained in Section 8.1 by multiplying by the appropriate correction factor shown in the following table Note 11:

<table>
<thead>
<tr>
<th>L/D:</th>
<th>1.75</th>
<th>1.50</th>
<th>1.25</th>
<th>1.00</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factor:</td>
<td>0.98</td>
<td>0.96</td>
<td>0.93</td>
<td>0.87</td>
</tr>
</tbody>
</table>

(Note 11) Use interpolation to determine correction factors for L/D values between those given in the table.

Note 11—Correction factors depend on various conditions such as moisture condition, strength level, and elastic modulus. Average values are given in the table. These correction factors apply to lightweight concrete weighing between 100 and 120 lb/ft³ (1,600 and 1,920 kg/m³) and to normal weight concrete. They are applicable to concrete dry or soaked at the time of loading and for nominal concrete strengths from 2000 to 6000 psi (15 to 45 MPa). For strengths higher than 6000 psi (45 MPa) correction factors may be larger than the values listed above.

9. Report

9.1 Report the following information:

9.1.1 Identification number;

9.1.2 Diameter (and length, if outside the range of 1.8D to 2.2D), in inches or millimeters;

9.1.3 Cross-sectional area, in square inches or centimeters;

9.1.4 Maximum load, in pounds-force or newtons;

9.1.5 Compressive strength calculated to the nearest 10 psi or 0.1MPa;

9.1.6 Type of fracture, if other than the usual cone (see Figure 2);

9.1.7 Defects in either specimen or caps; and

9.1.8 Age of specimen.

9.1.9 When determined, record the density to the nearest 10 kg/m³ (1 lb/ft³).
10. PRECISION AND BIAS

See AASHTO T 22 for Precision and bias

WSDOT has added Appendix A and it is an excerpt of ASTM C1231-00 sections 1 through 7

A1. Scope

A1.1 This practice covers requirements for a capping system using unbonded caps for testing concrete cylinders molded in accordance with Practice C 31/C 31M or C 192/C 192M. Unbonded neoprene caps of a defined hardness are permitted to be used for testing for a specified maximum number of reuses without qualification testing up to a certain concrete compressive strength level. Above that strength, level neoprene caps will require qualification testing. Qualification testing is required for all elastomeric materials other than neoprene regardless of the concrete strength.

A1.2 Unbonded caps are not to be used for acceptance testing of concrete with compressive strength below 1500 psi [10 MPa] or above 12 000 psi [85 MPa].

A1.3 The values stated in either inch-pound or SI units shall be regarded as standard. SI units are shown in brackets. That values stated in each system may not be exact equivalents; therefore, each system must be used independently of the other, without combining the values in any way.

A1.4 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. For a specific hazard statement, see Note 4.

A2. Referenced Documents

A2.1 ASTM Standards:
C 31/C 31M Practice for Making and Curing Concrete Test Specimens in the Field
C 39 Test Method for Compressive Strength of Cylindrical Concrete Specimens
C 192/C 192M Practice for Making and Curing Concrete Test Specimens in the Laboratory
C 617 Practice for Capping Cylindrical Concrete Specimens
D 2000 Classification System for Rubber products in Automotive Applications
D 2240 Test Method for Rubber Property—Durometer Hardness

A3. Terminology

A3.1 Definitions of Terms Specific to This Standard:
A3.1.1 pad, n—an unbonded elastomeric pad.
A3.1.2 unbonded cap, n—a metal retainer and an elastomeric pad.
A4. Significance and Use

A4.1 This practice provides for using an unbonded capping system in testing hardened concrete cylinders made in accordance with Practices C 31/C 31M or C 192/C 192M in lieu of the capping systems described in Practice C 617.

A4.2 The elastomeric pads deform in initial loading to conform to the contour of the ends of the cylinder and are restrained from excessive lateral spreading by plates and metal rings to provide a uniform distribution of load from the bearing blocks of the testing machine to the ends of the concrete or mortar cylinders.

A5. Materials and Apparatus

A5.1 Materials and equipment necessary to produce ends of the reference cylinders that conform to planeness requirements of Test Method C 39 and the requirements of Practice C 617. This may include grinding equipment or capping materials and equipment to produce neat cement paste, high strength gypsum plaster, or sulfur mortar caps.

A5.2 Elastomeric Pads:

A5.2.1 Pads shall be 1/2 ± 1/16 in. [13 ± 2 mm] thick and the diameter shall not be more than 1/16 in. [2 mm] smaller than the inside diameter of the retaining ring.

1 This practice is under the jurisdiction of ASTM Committee C09 on Concrete and Concrete Aggregate and is the direct responsibility of Subcommittee C09.61 on Testing Concrete for Strength. Current edition approved Jan. 10, 2000. Published April 2000. Originally published as C 1231–93. Last previous edition C 1231–99.


3 Annual Book of ASTM Standards, Vol 09.02.

4 Annual Book of ASTM Standards, Vol 09.01.

A5.2.2 Pads shall be made from polychloroprene (neoprene) meeting the requirements of Classification D 2000 as follows:

<table>
<thead>
<tr>
<th>Shore A Durometer</th>
<th>Classification D 2000 Line Call-Out</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>M2BC514</td>
</tr>
<tr>
<td>60</td>
<td>M2BC614</td>
</tr>
<tr>
<td>70</td>
<td>M2BC714</td>
</tr>
</tbody>
</table>

The tolerance on Shore A durometer hardness is ± 5. Table 1 provides requirements for use of caps made from material meeting the requirements of Classification D 2000, above.

A5.2.3 Other elastomeric materials that meet the performance requirements of qualification tests in Section 8 are permitted.
A5.2.4 Elastomeric pads shall be supplied with the following information:

A5.2.4.1 The manufacturer’s or supplier’s name,

A5.2.4.2 The Shore A hardness, and

A5.2.4.3 The applicable range of concrete compressive strength from Table 1 or from qualification testing.

A5.2.5 The user shall maintain a record indicating the date the pads are placed in service, the pad durometer, and the number of uses to which they have been subjected.

A5.3 Retainers, shall be made of metal that will prove durable in repeated use (Note 1). The cavity in the metal retainers shall have a depth at least twice the thickness of the pad. The inside diameter of the retaining rings shall not be less than 102 % or greater than 107 % of the diameter of the cylinder. The surfaces of the metal retainer which contact the bearing blocks of the testing machine shall be plane to within 0.002 in. [0.05 mm].

The bearing surfaces of the retainers shall not have gouges, grooves, or indentations greater than 0.010 in. [0.25 mm] deep or greater than 0.05 in.2 [32 mm2] in surface area.

NOTE 1—Retainers made from steel and some aluminum alloys have been found acceptable. Steel retaining rings that have been used successfully with 1/2-in. [13-mm] neoprene pads are shown in Fig. 1. Retainer design and metals used are subject to the performance and acceptance requirements of Section 8.

A6. Test Specimens

A6.1 The specimens shall be either 6 by 2 in. [150 by 300 mm] or 4 by 8 in. [100 by 200 mm] cylinders made in accordance with Practices C 31/C 31M or C 92/C 92M.

Neither end of a cylinder shall depart from perpendicularity to the axis by more than 0.5° (approximately equivalent to 1/8 in. in 12 in. [3 mm in 300 mm]). No individual diameter of a cylinder may differ from any other diameter by more than 2 %.

NOTE 2—One method of measuring the perpendicularity of ends of cylinders is to place a try square across any diameter and measure the departure of the longer blade from an element of the cylindrical surface. An alternative method is to place the end of the cylinder on a plane surface and support the try square on that surface.

A6.2 Depressions under a straight edge measured with a round wire gage across any diameter shall not exceed 0.20 in. [5 mm]. If cylinder ends do not meet this tolerance, the cylinder shall not be tested unless irregularities are corrected by sawing or grinding.

### TABLE 1 Requirements for Use of Polychloroprene (Neoprene) Pads

<table>
<thead>
<tr>
<th>Cylinder Compressive Strength, psi [MPa]</th>
<th>Shore A Durometer Hardness</th>
<th>Qualification Tests Required</th>
<th>Maximum ReusesA</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 500 to 6 000 [10 to 40]</td>
<td>50</td>
<td>none</td>
<td>100</td>
</tr>
<tr>
<td>2 500 to 7 000 [17 to 50]</td>
<td>60</td>
<td>none</td>
<td>100</td>
</tr>
<tr>
<td>4 000 to 7 000 [28 to 50]</td>
<td>70</td>
<td>None</td>
<td>100</td>
</tr>
<tr>
<td>7 000 to 12 000 [50 to 80]</td>
<td>70</td>
<td>Required</td>
<td>50</td>
</tr>
<tr>
<td>Greater than 12,000 [80]</td>
<td></td>
<td>not permitted</td>
<td></td>
</tr>
</tbody>
</table>

A Maximum number of reuses. Will be less if pads wear, crack or split. See Note 6.
A7. Procedure

A7.1 Unbonded caps are permitted to be used on one or both ends of a cylinder in lieu of a cap or caps meeting Practice C 617, provided they meet the requirements of Section 5.

A7.2 Examine the pads for excessive wear or damage (Note 6). Replace pads which have cracks or splits exceeding 3/8 in. [10 mm] in length regardless of depth. Insert the pads in the retainers before they are placed on the cylinder (Note 3).

NOTE 3—Some manufacturers recommend dusting the pads and the ends of the cylinders with corn starch or talcum powder prior to testing.

NOTE 4—Caution: Concrete cylinders tested with unbonded caps rupture more violently than comparable cylinders tested with bonded caps. As a safety precaution the cylinder testing machine must be equipped with a protective cage. In addition, some users have reported damage to testing machines from the sudden release of energy stored in the elastomeric pads.
A7.3 Center the unbonded cap or caps on the cylinder and place the cylinder on the lower bearing block of the testing machine. Carefully align the axis of the cylinder with the center of thrust of the testing machine by centering the upper retaining ring on the spherically seated bearing block. As the spherically seated block is brought to bear on the upper retaining ring, rotate its movable portion gently by hand so that uniform seating is obtained. After application of load, but before reaching 10 % of the anticipated specimen strength, check to see that the axis of the cylinder is vertical within a tolerance of 1⁄8 in. [3.2 mm in 300 mm] and that the ends of the cylinder are centered within the retaining rings. If the cylinder alignment does not meet these requirements, release the load, check compliance with 6.1, and carefully recenter the specimen. Reapply load and recheck specimen centering and alignment. A pause in load application to check cylinder alignment is permissible.

A7.4 Complete the load application, testing, calculation, and reporting of results in accordance with Test Method C 39.

NOTE 5—Because of the violent release of energy stored in pads, the broken cylinder rarely exhibits conical fracture typical of capped cylinders and the sketches of types of fracture in Test Method C 39 are not descriptive. Occasionally, unbonded capped cylinders may develop early cracking, but continue to carry increasing load. For this reason cylinders must be tested to complete failure.
Performance Exam Checklist

Compressive Strength of Cylindrical Concrete Specimens
FOP for AASHTO T 22

Participant Name __________________________________________ Exam Date __________________________

Procedure Element
1. The tester has a copy of the current procedure on hand? □ Yes □ No
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? □ Yes □ No
3. Is the diameter of the cylinder record to the nearest 0.01 inch by averaging two diameters taken at about mid-height? □ Yes □ No
4. Are lower and upper bearing surface wiped clean? □ Yes □ No
5. Is the axis of the cylinder aligned with center of the spherical block? □ Yes □ No
6. Is the spherical block rotated prior to it contacts with the cylinder? □ Yes □ No
7. Is the load applied continuously and without shock? □ Yes □ No
8. Is the load applied at the specified rate and maintain for the latter half of the anticipated load. □ Yes □ No
9. Is no rate adjustment made while the cylinder is yielding? □ Yes □ No
10. Is the maximum load recorded? □ Yes □ No
11. Are cylinders tested to failure, or to 80% capacity of machine, and the type of fracture recorded? □ Yes □ No
12. Breaking Cylinders at 28 days ± 20 hours? □ Yes □ No
13. All calculations performed correctly? □ Yes □ No

Pad Cap-concrete Cylinders AASHTO-22 Appendix A
1. Ends of cylinders checked for perpendicularity to axis? □ Yes □ No
2. Ends of cylinders checked for depressions greater than 0.2 inch? □ Yes □ No
3. Pads examined for splits or cracks? □ Yes □ No
4. Cylinders centered in retaining rings? □ Yes □ No
5. Is cylinders checked for alignment with a small load applied? □ Yes □ No

First attempt: Pass □ Fail □ Second attempt: Pass □ Fail □

Signature of Examiner __________________________________________
WSDOT FOP for AASHTO T 23

Making and Curing Concrete Test Specimens in the Field

1. SCOPE

1.1 This method covers procedures for making and curing cylinder and beam specimens from representative samples of fresh concrete for a construction project.

1.2 The concrete used to make the molded specimens shall be sampled after all on-site adjustments have been made to the mixture proportions, including the addition of mix water and admixtures, except as modified in Section 5.1. This practice is not satisfactory for making specimens from concrete not having measurable slump or requiring other sizes or shapes of specimens.

1.3 The values stated in SI units are to be regarded as the standard.

1.4 This standard does not purport to address the safety problems 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 AASHTO Standards

M 195, Lightweight Aggregates for Structural Concrete
M 201 Moist Cabinets, Moist Rooms, and Water Storage Tanks Used in the Testing of Hydraulic Cements and Concretes
M 205 Molds for Forming Concrete Test Cylinders Vertically
T 119 Slump of Hydraulic Cement Concrete
T 126 Making and Curing Concrete Test Specimens in the Laboratory
T 141 Sampling Freshly Mixed Concrete
T 152 Air Content of Freshly Mixed Concrete by the Pressure Method
T 196 Air Content of Freshly Mixed Concrete by the Volumetric Method
T 197 Time of Setting of Concrete Mixtures by Penetration Resistance
T 231 Capping Cylindrical Concrete Specimens
T 309 Temperature of Freshly Mixed Portland-Cement Concrete

ASTM Standards:
C 125, Terminology Related to Concrete and Concrete Aggregates

ACI Standards:
309 R, Guide for Consolidation of Concrete

This FOP is based on AASHTO T 23-04
3. Terminology

For definitions of terms used in this practice, refer to Terminology ASTM C 125.

4. SIGNIFICANCE AND USE

4.1 This practice provides standardized requirements for making, curing, protecting, and transporting concrete test specimens under field conditions.

4.2 If the specimens are made and standard cured, as stipulated here, the resulting strength test data where the specimens are tested are able to be used for the following purposes:

4.2.1 Acceptance testing for specified strength,
4.2.2 Checking the adequacy of mixture proportions for strength.
4.2.3 Quality control.

4.3 If the specimens are made and field cured, as stipulated herein, the resulting strength test data when the specimens are tested are able to be used for the following purposes:

4.3.1 Determination of whether a structure is capable of being put in service.
4.3.2 Comparison with test results of standard cured specimens or with test results from various in-place test methods,
4.3.4 Adequacy of curing and protection of concrete in the structure, or,
4.3.5 Form or shoring removal time requirements,

5. APPARATUS

5.1 Molds, General — Molds for specimens or fastenings thereto in contact with the concrete shall be made of steel, cast iron, or other nonabsorbent material, non-reactive with concrete containing portland or other hydraulic cements. Molds shall hold their dimensions and shape under all conditions of use. Molds shall be watertight during use as judged by their ability to hold water poured into them. Provisions for tests of watertightness are given in 6 of Specification M 205. A suitable sealant, such as heavy grease, modeling clay, or microcrystalline wax shall be used where necessary to prevent leakage through the joints. Positive means shall be provided to hold base plates firmly to the molds. Molds shall be lightly coated with mineral oil or a suitable non-reactive form release material before use.

5.2 Cylinder: Molds for casting concrete test specimens shall conform to the requirements of M 205, and Shall come from an approved shipment as verified by the WSDOT Quality Systems Manual Verification Procedure No. 2.

5.3 Beam Molds — Refer to WSDOT Test Method T 808

5.4 Tamping Rod — Two sizes are specified as indicated in Table 1. Each shall be a round, straight steel rod with at least the tamping end rounded to a hemispherical tip of the same diameter as the rod. Both ends may be rounded if preferred.
### Table 1—Tamping Rod Requirements

<table>
<thead>
<tr>
<th>Diameter of Cylinder or Width of Beam, in (mm.)</th>
<th>Rod Dimensions</th>
<th>Diameter, in (mm.)</th>
<th>Length of Rod, in (mm.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;6 (150)</td>
<td>⅜ (10)</td>
<td>12 (300)</td>
<td></td>
</tr>
<tr>
<td>6 (150)</td>
<td>⅝ (16)</td>
<td>20 (500)</td>
<td></td>
</tr>
<tr>
<td>9 (225)</td>
<td>⅝ (16)</td>
<td>26 (650)</td>
<td></td>
</tr>
</tbody>
</table>

* Rod tolerances length 100 mm (±4 in.) and diameter 2 mm (± 0.079 in.).

5.5 Vibrators — Internal vibrators shall be used. The vibrator frequency shall be at least 7,000 vibrations per minute (150 Hz) while the vibrator is operating in the concrete. The diameter of a round vibrator shall be no more than one-fourth the diameter of the cylinder mold or one-fourth the width of the beam mold. Other shaped vibrators shall have a perimeter equivalent to the circumference of an appropriate round vibrator. The combined length of the vibrator shaft and vibrating element shall exceed the depth of the section being vibrated by at least 3 in. (75 mm). The vibrator frequency shall be checked periodically.

**Note 1** — For information on size and frequency of various vibrators and a method to periodically check vibrator frequency, see ACI 309R.

5.6 Mallet — A mallet with a rubber or rawhide head weighing 1.25 ± 0.50 lb [0.57 ± 0.23 kg] shall be used.

5.7 Small Tools — Tools and items that may be required are shovels, pails, trowels, wood float, metal float, blunted trowels, straightedge, feeler gauge, scoops, and rules.

5.8 Slump Apparatus — The apparatus for measurement of slump shall conform to the requirements of T 119.

5.9 Sampling and Mixing Receptacle — The receptacle shall be a suitable heavy gage metal pan, wheelbarrow, or flat, clean non-absorbent mixing board of sufficient capacity to allow easy remixing of the entire sample with a shovel or trowel.

5.10 Air Content Apparatus — The apparatus for measuring air content shall conform to the requirements of T 196 or T 152.

5.11 Temperature Measuring Devices — The temperature measuring devices shall conform to the applicable requirements of Test Method T 309.

6. TESTING REQUIREMENTS

Testing for determining the compressive strength at 28 days shall require a set of two specimens made from the same sample.

Testing for compressive strength at ages less than 28 days, a single cylinder specimen for each test age will be sufficient.
6.1 **Compressive Strength Specimens** — Compressive strength specimens shall be cylinders cast and allowed to set in an upright position. The length shall be twice the diameter. The cylinder diameter shall be at least three times the nominal maximum size of the coarse aggregate. The standard specimen shall be the 4 by 8-in. (100 by 200-mm) cylinder when the nominal maximum size of the coarse aggregate does not exceed 1 in. (25 mm). When the nominal maximum size of the coarse aggregate exceeds 1 in. (25 mm) the specimens shall be made with 6 by 12 in. (150 by 300 mm) cylinders. **Mixing of cylinder sizes for a particular class of mix is not permitted on a project.** When the nominal maximum size of the coarse aggregate exceeds 2 in (50 mm), the concrete sample shall be treated by wet sieving through a 2 in (50 mm) sieve as described in FOP for WAQTC TM 2. Contact the Materials Laboratory for directions.

**Note 2:** The nominal maximum size is the smallest sieve opening through which the entire amount of aggregate is REQUIRED to pass.

**Note 3:** When molds in SI units are required and not available, equivalent inch-pound unit size molds should be permitted.

6.2 **Flexural Strength Specimens** — Flexural strength specimens shall be beams of concrete cast and hardened with long axes horizontal. The length shall be at least 2 in. [50 mm] greater than three times the depth as tested. The ratio of width to depth as molded shall not exceed 1.5. The standard beam shall be 6 by 6 in. [150 by 150 mm] in cross section, and shall be used for concrete with maximum size coarse aggregate up to 2 in. [50 mm]. When the nominal maximum size of the coarse aggregate exceeds 2 in. [50 mm], the smaller cross sectional dimension of the beam shall be at least three times the nominal maximum size of the coarse aggregate. Unless required by project specifications, beams made in the field shall not have a width or depth of less than 6 in. [150 mm].

Refer to WSDOT Test Method T 808

7. **SAMPLING CONCRETE**

7.1 The samples used to fabricate test specimens under this standard shall be obtained in accordance with FOP for WAQTC TM-2 unless an alternative procedure has been approved.

7.2 Record the identification of the sample with respect to the location of the concrete represented and the time of casting.

8. **SLUMP, AIR CONTENT, AND TEMPERATURE**

8.1 Slump — Measure and record the slump of each batch of concrete from which specimens are made immediately after remixing in the receptacle, as required in FOP FOR AASHTO T 119.

8.2 Air Content — Determine the air content in accordance with either FOP for AASHTO T 152 or FOP for AASHTO T 196. The concrete used in performing the air content test shall not be used in fabricating test specimens.

8.3 Temperature — Determine and record the temperature in accordance with FOP for AASHTO T 309.

**Note 4** — Some specifications may require the measurement of the unit weight of concrete. The volume of concrete produced per batch may be desired on some projects. Also, additional information on the air content measurements may be desired. Test Method T 121 is used to measure the unit weight, yield, and gravimetric air content of freshly mixed concrete.
9. MOLDING SPECIMENS

9.1 Place of Molding — Mold specimens promptly on a level, rigid horizontal surface, free of vibration and other disturbances, at a place as near as practicable to the location where they are to be stored.

9.2 Casting the Concrete — Place the concrete in the mold using a scoop, blunted trowel, or shovel. Select each scoopful, trowelful, or shovelful of concrete from the mixing pan to ensure that it is representative of the batch. Remix the concrete in the mixing pan with a shovel or trowel to prevent segregation during the molding of specimens. Move the scoop, trowel, or shovel around the perimeter of the mold opening when adding concrete so the concrete is uniformly distributed within each layer with a minimum of segregation. Further distribute the concrete by use of the tamping rod prior to the start of consolidation. In placing the final layer, the operator shall attempt to add an amount of concrete that will exactly fill the mold after consolidation. Underfilled molds shall be adjusted with representative concrete during consolidation of the top layer. Overfilled molds shall have excess concrete removed.

9.2.1 Number of Layers — Make specimens in layers as indicated in Table 2 or 3.

### Table 2—Molding Requirements by Rodding

<table>
<thead>
<tr>
<th>Specimen Type and Size</th>
<th>Number of Layers of Approximately Equal Depth</th>
<th>Number of Roddings per Layer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cylinders:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diameter, mm (in.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 (4)</td>
<td>2</td>
<td>25</td>
</tr>
<tr>
<td>150 (6)</td>
<td>3</td>
<td>25</td>
</tr>
<tr>
<td>225 (9)</td>
<td>4</td>
<td>50</td>
</tr>
<tr>
<td>Beams:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Width, mm (in.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>150 (6) to 200 (8)</td>
<td>3 or more equal depths, each not to exceed 150 mm (6 in.)</td>
<td>See 8.3.2</td>
</tr>
<tr>
<td>200 (&gt;8)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Table 3—Molding Requirements by Vibration

<table>
<thead>
<tr>
<th>Specimen Type and Size</th>
<th>Number of Layers</th>
<th>Number of Vibrator Insertions per Layer</th>
<th>Approximate Depth of Layer, mm (in.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cylinders:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diameter, mm (in.)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 (4)</td>
<td>2</td>
<td>1</td>
<td>one-half depth of specimen</td>
</tr>
<tr>
<td>150 (6)</td>
<td>2</td>
<td>2</td>
<td>one-half depth of specimen</td>
</tr>
<tr>
<td>225 (9)</td>
<td>2</td>
<td>4</td>
<td>one-half depth of specimen</td>
</tr>
<tr>
<td>Beams:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Width, mm (in.)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>150 (6) to 200 (8)</td>
<td>1</td>
<td>See 8.4.2</td>
<td>depth of specimen 200 (8) as near as practicable</td>
</tr>
<tr>
<td>over 200 (8)</td>
<td>2 or more</td>
<td>See 8.4.2</td>
<td>depth of specimen 200 (8) as near as practicable</td>
</tr>
</tbody>
</table>
9.2.2 Select the proper tamping rod from 4.4 and Table 1 or the proper vibrator from 4.5. If the method of consolidation is rodding, determine molding requirements from Table 2. If the method of consolidation is vibration, determine molding requirements from Table 3.

9.3 Consolidation:

9.3.1 Method of Consolidation — Preparation of satisfactory specimens require different methods of consolidation. The methods of consolidation are rodding and vibration. Base the selection of the method of consolidation on slump, unless the method is stated in the specifications under which the work is being performed. Rod or vibrate concretes with slumps greater than 1 in. (25 mm). Vibrate concretes with slumps less than or equal to 1 in. (25 mm). Concretes of such low water content that they cannot be properly consolidated by the method herein, or requiring other sizes and shapes of specimens to represent the product or structure, are not covered by this method. Specimens for such concretes shall be made in accordance with the requirements of Method T 126 with regards to specimen size and shape and method of consolidation.

9.3.2 Rodding — Place the concrete in the mold, in the required number of layers of approximately equal volume. Rod each layer with the rounded end of the rod using the required number of roddings specified in Table 2. The number of roddings per layer required for beams is one for each 2 in² (3 cm²) top surface area of the specimen. Rod the bottom layer throughout its depth. Distribute the strokes uniformly over the cross section of the mold. For each layer, allow the rod to penetrate through the layer being rodded and into the layer below approximately 25 mm (1 in.). After each layer is rodded, tap the outsides of the mold lightly 10 to 15 times with the open hand mallet, or rod, to close any holes left by rodding and to release any large air bubbles that may have been trapped. Use an open hand to tap light-gage single-use molds which are susceptible to damage if tapped with a mallet or rod. After tapping, spade each layer of the concrete along the sides and ends of beam molds with a trowel or other suitable tool.

9.3.3 Vibration — Maintain a uniform time period for duration of vibration for the particular kind of concrete, vibrator, and specimen mold involved. The duration of vibration required will depend upon the workability of the concrete and the effectiveness of the vibrator. Usually, sufficient vibration has been applied as soon as the surface of the concrete has become relatively smooth, and large air bubbles cease to break through the top surface. Continue vibration only long enough to achieve proper consolidation of the concrete. (See Note 5.) Fill the molds and vibrate in the required number of approximately equal layers. Place all the concrete for each layer in the mold before starting vibration of that layer. Compacting the specimen, insert the vibrator slowly and do not allow it to rest on the bottom or sides of the mold. Slowly withdraw the vibrator so that no large air pockets are left in the specimen. When placing the final layer, avoid overfilling by more than ¼ in. (6 mm).

Note 5 — Generally, no more than 5 s of vibration should be required for each insertion to adequately consolidate concrete with a slump greater than 3 in. (75 mm). Longer times may be required for lower slump concrete, but the vibration time should rarely have to exceed 10 s per insertion.
9.3.3.1 **Cylinders** — The number of insertions of a vibrator at per layer is given in Table 3. When more than one insertion per layer is required, distribute the insertion uniformly within each layer. Allow the vibration to penetrate through the layer being vibrated, and into the layer below, approximately 1 in. (25 mm). After each layer is vibrated, tap the outsides of the mold at least 10 times with the mallet, to close holes left by vibrating and to release entrapped air voids. Use an open hand to tap cardboard and single-use metal molds, that are susceptible to damage if tapped with a mallet.

9.3.3.2 **Beam** — Refer to WSDOT Test Method T 808.

9.4 **Finishing** — After consolidation, strike off excess concrete from the surface and float or trowel it as required. Perform all finishing with the minimum manipulation necessary to produce a flat even surface that is level with the rim or edge of the mold and that has no depressions or projections larger than ⅛ in. (3.2 mm).

9.4.1 **Cylinders** — After consolidation, finish the top surfaces by striking them off with the tamping rod where the consistency of the concrete permits or with a wood float or trowel. If desired, cap the top surface of freshly made cylinders with a thin layer of stiff Portland cement paste which is permitted to harden and cure with the specimen. See section on Capping Materials of T 231.

9.4.2 **Beams** — After consolidation of the concrete, strike off the top surface to the required tolerance to produce a flat even surface. A wood float may be used.

9.5 **Initial Storage** — Immediately after being struck off, the specimens shall be move to the storage place were they will remain undisturbed for the initial curing period. If specimens made in single-use mold are move, lift and support the specimens from the bottom of the molds with a large trowel or similar device.

10. **CURING**

10.1 **Standard Curing** — Standard curing is the curing method used when the specimens are made and cured for the purposes stated in 4.2.

10.1.1 **Storage** — If specimens cannot be molded at the place where they will receive initial curing, immediately after finishing, move the specimens to an initial curing place for storage. The supporting surface on which specimens are stored shall be level to within ¼ in. per ft (20 mm per m.). If cylinders in the single-use molds are moved, lift and support the cylinders from the bottom of the molds with a large trowel or similar device. If the top surface is marred during movement to place of initial storage, immediately refinish.

10.1.2 **Initial Curing** — Immediately after molding and finishing, the specimens shall be stored for a period 24 ± 8 hours, unless Contractor provides initial curing information for final set up to 48 h, in at a temperature range from 60 to 80°F (16 to 27°C), and in an environment preventing moisture loss from the specimens. For concrete mixtures with a specified strength of 6000 psi (40 Mpa) or greater, the initial curing temperature shall be between 68 and 78°F (20 and 26°C). Various procedures are capable of being used during the initial curing period to maintain the specified moisture and temperature conditions. An appropriate procedure or combination of procedures shall be used (Note 6). Shield all specimens from direct sunlight and, if used, radiant heating devices. The storage temperature shall be controlled by the use of heating and cooling devices, as necessary. Record the temperature using a maximum-minimum thermometer. If cardboard molds are used, protect the outside surface of the molds from contact with wet burlap or other sources of water.
Note 6 — A satisfactory moisture environment can be created during the initial curing of the specimens by one or more of the following procedures: (1) immediately immerse molded specimens with plastic lids in water saturated with calcium hydroxide, (2) store in properly constructed wood boxes or structures, (3) place in damp sand pits, (4) cover with removable plastic lids, (5) place inside plastic bags, or (6) cover with plastic sheets or nonabsorbent plates if provisions are made to avoid drying and damp burlap is used inside the enclosure, but the burlap is prevented from contacting the concrete surfaces.

A satisfactory temperature environment can be controlled during the initial curing of the specimens by one or more of the following procedures: approved cure boxes with capability of maintaining a temperature of 60-80 degrees F, store in properly constructed wood boxes or structures, place in damp sand pits, or by (1) use of ventilation, (2) use of ice, (3) use of thermostatically controlled heating or cooling devices, or (4) use of heating methods such as stoves or light bulbs. Other suitable methods may be used if the requirements limiting specimen storage temperature and moisture loss are met. For concrete mixtures with a specified strength of 6000 psi (40 MPa) or greater, heat generated during the early ages may raise the temperature above the required storage temperature. When specimens are to be immersed in water saturated with calcium hydroxide, specimens in cardboard molds or other molds that expand when immersed in water should not be used. Early-age strength test results may be lower when stored at 60ºF (16ºC) and higher when stored at 80ºF (27ºC). On the other hand, at later ages, test results may be lower for higher initial storage temperatures.

10.1.3 Final Curing:

10.1.3.1 Cylinders—Upon completion of initial curing and within 30 minutes after removing the molds, cure specimens with free water maintained on their surfaces at all times at a temperature of 73 ± 3ºF (23 ± 2ºC) using water storage tanks or moist rooms complying with the requirements of Specification M 201, except when capping with sulfur mortar capping compound and immediately before testing. When capping with sulfur mortar capping compounds, the ends of the cylinder shall be dry enough to preclude the formation of steam or foam pockets under or in cap larger than ¼ in (6 mm.) as described in T 231. For a period not to exceed 3 h immediately prior to test, standard curing temperature is not required provided free moisture is maintained on the cylinders and ambient temperature is between 68 to 80ºF (20 and 30ºC).

10.1.3.2 Beams—Refer to WSDOT Test Method T 808. Beams are to be cured the same as cylinders (see 9.1.3.1), except that they shall be stored in water saturated with calcium hydroxide at 73 ± 3ºF (23 ± 2ºC) at least 20 h prior to testing. Drying of the surfaces of the beam shall be prevented between removal from water storage and completion of testing (Note 7).

Note 7 — Relatively small amounts of surface drying of flexural specimens can induce tensile stresses in the extreme fibers that will markedly reduce the indicated flexural strength.
10.2 Field Curing—Field curing is the curing method used for the specimens made for the purposes stated in 4.3.

10.2.1 Cylinders — Store cylinders in or on the structure as near to the point of deposit of the concrete represented as possible. Protect all surfaces of the cylinders from the elements in as near as possible the same way as the formed work. Provide the cylinders with the same temperature and moisture environment as the structural work. Test the specimens in the moisture condition resulting from the specified curing treatment. To meet these conditions, specimens made for the purpose of determining when a structure is capable of being put in service shall be removed from the molds at the time of removal of form work.

10.2.2 Beams — Refer to WSDOT Test Method T 808.

10.3 Structural Lightweight Concrete Curing—Cure structural lightweight concrete cylinders in accordance with M 195.

To prevent evaporation of water from the unhardened concrete, cover the specimen with a nonabsorptive, nonreactive plate or sheet of tough, durable, impervious plastic or wet burlap. When wet burlap is used for covering, the burlap must be kept wet until the specimens are removed from the mold. (See Note 3 below.) Remove specimen from the mold in 24 ± 8 hours after casting and store in a moist room maintained at 73 ± 3°F (23 ± 2°C) with a relative humidity of not less than 95 percent. At the age of seven days, remove the specimen from the moist room, measure for length, and store in a curing cabinet maintained at 100 ± 2°F (37.8 ± 1.1°C) with a relative humidity of 32 ± 2 percent.

Note 3 — Placing a sheet of plastic over the burlap will facilitate keeping it wet.

11. TRANSPORTATION OF SPECIMENS TO LABORATORY

11.1 Prior to transporting, cure and protect specimens as required in Section 10. Specimens shall not be transported until at least 8 h after final set. (See Note 8.) During transporting, protect the specimen with suitably cushioning material to prevent damage from jarring. During cold weather, protect the specimens from freezing with suitable insulation material. Prevent moisture loss during transportation by wrapping the specimens in plastic, wet burlap, by surrounding them with wet sand or tight-fitting plastic caps on plastic molds. Transportation time shall not exceed 46 h.

Note 8 — Setting time may be measured by T 197. If a specimen does not attain final set within hours 24 ± 8 hours, it is to remain in place until final set is reached. After final set is reached, it can then be transported. The time of final set shall be provided by the concrete producer.

12 REPORT

12.1 Report the following information to the laboratory that will test the specimens:

12.1.1 Identification number;

12.1.2 Location of concrete represented by the samples;

12.1.3 Date, time, and name of individual molding specimens;
12.1.4 Slump, air content, and concrete temperature, test results and results of any other tests on the fresh concrete and any deviations from referenced standard test methods, and

12.1.5 Curing method. For standard curing method, report the initial curing method with maximum and minimum temperatures and final curing method. For field curing method, report the location where stored, manner of protection from the elements, temperature and moisture environment, and time of removal from molds.

Record all information required on WSDOT Form 350-009 Concrete Cylinder Transmittal.
Performance Exam Checklist

Making and Curing Concrete Test Specimens in the Field
FOP for AASHTO T 23

Participant Name ___________________________________________ Exam Date ______________

Procedure Element                                                  Yes  No
1. The tester has a copy of the current procedure on hand?             [ ]  [ ]
2. Molds placed on a level, rigid, horizontal surface free of vibration? [ ]  [ ]
3. Making of specimens begun within 15 minutes of sampling?           [ ]  [ ]
4. Concrete placed in the mold, moving a scoop or trowel around the perimeter of the mold to evenly distribute the concrete as discharged? [ ]  [ ]
5. Mold filled in correct number of layers, attempting to exactly fill the mold on the last layer? [ ]  [ ]
6. Each layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes? [ ]  [ ]
7. Bottom layer rodded throughout its depth?                         [ ]  [ ]
8. Middle and top layers rodded, each throughout their depths, and penetrate into the underlying layer? [ ]  [ ]
9. Sides of the mold tapped 10-15 times after rodding each layer?      [ ]  [ ]
   a. with mallet for reusable steel molds                          [ ]  [ ]
   b. with the open hand for flexible light-gauge molds              [ ]  [ ]
   c. with open hand or tamping rod for plastic mold               [ ]  [ ]
10. Strike off excess concrete, and finished the surface with a minimum of manipulation? [ ]  [ ]
11. Specimens covered with non-absorbent, nonreactive cap or plate?   [ ]  [ ]

First attempt: Pass [ ] Fail [ ]   Second attempt: Pass [ ] Fail [ ]

Signature of Examiner ___________________________________________

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SIEVE ANALYSIS OF FINE AND COARSE AGGREGATES

Significance

Sieve analyses are performed on aggregates used in roadway bases and in portland cement and asphalt cement concretes. Sieve analyses reveal the size makeup of aggregate particles – from the largest to the smallest. A gradation curve or chart showing how evenly or unevenly the sizes are distributed between largest and smallest is created in this test. How an aggregate is graded has a major impact on the strength of the base or on the properties and performance of concrete. In portland cement concrete (PCC), for example, gradation influences shrinkage and shrinkage cracking, pumpability, finishability, permeability, and other characteristics.

Scope

This procedure covers sieve analysis in accordance with AASHTO T 27 and materials finer than No. 200 (75 µm) in accordance with AASHTO T 11. The procedure combines the two test methods.

Sieve analyses determines the gradation or distribution of aggregate particles within a given sample in order to determine compliance with design and production standards.

Accurate determination of material smaller than No. 200 (75 µm) cannot be made with AASHTO T 27 alone. If quantifying this material is required, it is recommended that AASHTO T 27 be used in conjunction with AASHTO T 11. Following AASHTO T 11, the sample is washed through a No. 200 (75 µm) sieve. The amount of material passing this sieve is determined by comparing dry sample masses before and after the washing process.

This procedure covers sieve analysis in accordance with AASHTO T 27 and materials finer than No. 200 (75 µm) in accordance with AASHTO T 11. The procedure includes two method choices, A, and B.

Apparatus

- Balance or scale: Capacity sufficient for the masses shown in Table 1, accurate to 0.1 percent of the sample mass or better and conform to the requirements of AASHTO M 231.
- Sieves – Meeting the requirements of AASHTO M 92.
- Mechanical sieve shaker – Meeting the requirements of AASHTO T 27.
- Suitable drying equipment (see FOP for AASHTO T 255)
- Containers and utensils: A pan or vessel of a size sufficient to contain the sample covered with water and to permit vigorous agitation without loss of any part of the sample or water
- Optional Mechanical washing device

This FOP is based on T 27/T 11 and has been modified per WSDOT standards. To View the redline modifications, contact WSDOT Quality Systems Manager (360) 709-5497.
Sample Preparation

Obtain samples in accordance with the FOP for AASHTO T 2 and reduce to the size shown in Table 1 in accordance with the FOP for AASHTO T 248.

If the gradation sample is obtained from FOP for AASHTO T-308, the Ignition Furnace, proceed to Procedure Step 1 of procedure method A.

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>Sample Sizes for Aggregate Gradation Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nominal Maximum Size* in. (mm)</td>
<td>Minimum Mass lb kg</td>
</tr>
<tr>
<td>US No. 4 (4.75)</td>
<td>1 0.5</td>
</tr>
<tr>
<td>1/4 (6.3)</td>
<td>2 1</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td>2 1</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>5 2</td>
</tr>
<tr>
<td>5/8 (16.0)</td>
<td>5 2</td>
</tr>
<tr>
<td>3/4 (19.0)</td>
<td>7 3</td>
</tr>
<tr>
<td>1 (25.0)</td>
<td>13 6</td>
</tr>
<tr>
<td>1 1/4 (31.5)</td>
<td>17 7.5</td>
</tr>
<tr>
<td>1 1/2 (37.5)</td>
<td>20 9</td>
</tr>
<tr>
<td>2 (50)</td>
<td>22 10</td>
</tr>
<tr>
<td>2 1/2 (63)</td>
<td>27 12</td>
</tr>
<tr>
<td>3 (75)</td>
<td>33 15</td>
</tr>
<tr>
<td>3 1/2 (90)</td>
<td>44 20</td>
</tr>
</tbody>
</table>

* For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.

Note: For an aggregate specification having a generally unrestrictive gradation (i.e. wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.

**WSDOT Note 1:** These sample sizes are standard for aggregate testing but, due to equipment restraints, samples may need to be partitioned into several “subsamples.” See method A.

Selection of Procedure

Agencies may specify what method will be performed. If a method is not specified, Method A will be performed.
Overview

Method A

- Determine dry mass of original sample
- Wash through a No. 200 (75 µm) sieve
- Determine dry mass of washed sample
- Sieve material

Method B

- Determine dry mass of original sample
- Wash through a No. 200 (75 µm) sieve
- Determine dry mass of washed sample
- Sieve coarse material
- Determine mass of fine material
- Reduce fine portion
- Determine mass of reduced portion
- Sieve fine portion

Sample Sieving

In all procedures it is required to shake the sample over nested sieves. Sieves are selected to furnish information required by specification. The sieves are nested in order of decreasing size from the top to the bottom and the sample, or a portion of the sample, is placed on the top sieve. Additional sieves may be necessary to provide other information, such as fineness modulus, or to keep from overloading the specified sieves. The sample may also be sieved in increments.

Sieves are shaken in a mechanical shaker for the minimum time determined to provide complete separation for the sieve shaker.

Time Evaluation

Overload Determination

Additional sieves may be necessary to provide other information, such as fineness modulus, or to keep from overloading sieves. The sample may also be sieved in increments.

WSDOT Note 2: Prevent an overload of material on an individual sieve by one of the following methods (See Table 2 for maximums allowed):

A. Insert an additional sieve with opening size intermediate between the sieve that may be overloaded and the sieve immediately above that sieve in the original set of sieves.

B. Split the sample into two or more portions, sieving each portion individually. Combine the masses of the several portions retained on a specific sieve before calculating the percentage of the sample on the sieve.

C. Use sieves having a larger frame size and providing greater sieving area.
TABLE 2
Maximum Allowable Mass of Material Retained on a Sieve, kg

<table>
<thead>
<tr>
<th>Sieve Size US inches (mm)</th>
<th>8 φ (203)</th>
<th>12 φ (305)</th>
<th>12 x 12 (305 x 305)</th>
<th>14 x 14 (350 x 350)</th>
<th>16 x 24 (372 x 580)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieving Area m²</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.0285</td>
<td>0.0670</td>
<td>0.0929</td>
<td>0.1225</td>
<td>0.2158</td>
<td></td>
</tr>
<tr>
<td>3 1/2 (90)</td>
<td>*</td>
<td>15.1</td>
<td>20.9</td>
<td>27.6</td>
<td>48.5</td>
</tr>
<tr>
<td>3 (75)</td>
<td>*</td>
<td>12.6</td>
<td>17.4</td>
<td>23.0</td>
<td>40.5</td>
</tr>
<tr>
<td>2 1/2 (63)</td>
<td>*</td>
<td>10.6</td>
<td>14.6</td>
<td>19.3</td>
<td>34.0</td>
</tr>
<tr>
<td>2 (50)</td>
<td></td>
<td>3.6</td>
<td>8.4</td>
<td>11.6</td>
<td>15.3</td>
</tr>
<tr>
<td>1 1/2 (37.5)</td>
<td></td>
<td>2.7</td>
<td>6.3</td>
<td>8.7</td>
<td>11.5</td>
</tr>
<tr>
<td>1 (25.0)</td>
<td></td>
<td>1.8</td>
<td>4.2</td>
<td>5.8</td>
<td>7.7</td>
</tr>
<tr>
<td>3/4 (19.0)</td>
<td></td>
<td>1.4</td>
<td>3.2</td>
<td>4.4</td>
<td>5.8</td>
</tr>
<tr>
<td>5/8 (16.0)</td>
<td></td>
<td>1.1</td>
<td>2.7</td>
<td>3.7</td>
<td>4.9</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td></td>
<td>0.89</td>
<td>2.1</td>
<td>2.9</td>
<td>3.8</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td></td>
<td>0.67</td>
<td>1.6</td>
<td>2.2</td>
<td>2.9</td>
</tr>
<tr>
<td>1/4 (6.3)</td>
<td></td>
<td>0.44</td>
<td>1.1</td>
<td>1.5</td>
<td>1.9</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td></td>
<td>0.33</td>
<td>0.80</td>
<td>1.1</td>
<td>1.5</td>
</tr>
<tr>
<td>Less than (No. 4)</td>
<td></td>
<td>0.20</td>
<td>0.47</td>
<td>0.65</td>
<td>1.2</td>
</tr>
</tbody>
</table>

Procedure Method A

1. Dry the sample in accordance with the FOP for AASHTO T 255, and record to the nearest 0.1 percent of total mass or better.

2. When the specification requires that the amount of material finer than No. 200 (75 µm) be determined, do Step 3 through Step 9 – otherwise, skip to Step 10.

**WSDOT Note 3:** If the applicable specification requires that the amount passing the No. 200 (75 µm) sieve be determined on a portion of the sample passing a sieve smaller than the nominal maximum size of the aggregate, separate the sample on the designated sieve and determine the mass of the material passing that sieve to 0.1 percent of the mass of this portion of the test sample. Use the mass as the original dry mass of the test sample.

3. Nest a sieve, such as a No. 10 (2 mm), above the No. 200 (75 µm) sieve.

4. Place the test sample in a container and add sufficient water to cover it.

**WSDOT requires the use of a detergent, dispersing agent, or other wetting solution when washing a sample from FOP for AASHTO T308, an ignition furnace sample.**

**WSDOT Note 4:** A detergent, dispersing agent, or other wetting solution may be added to the water to assure a thorough separation of the material finer than the No. 200 (75 µm) sieve from the coarser particles. There should be enough wetting agent to produce a small amount of suds when the sample is agitated. Excessive suds may overflow the sieves and carry material away with them.

5. Agitate vigorously to ensure complete separation of the material finer than No. 200 (75 µm) from coarser particles and bring the fine material into suspension above the coarser material. When using a mechanical washing device, exercise caution to not degrade the sample.
6. Immediately pour the wash water containing the suspended and dissolved solids over the nested sieves, being careful not to pour out the coarser particles.

7. Add a second change of water to the sample remaining in the container, agitate, and repeat Step 6. Repeat the operation until the wash water is reasonably clear.

8. Return all material retained on the nested sieves to the container by flushing into the washed sample.

9. Dry the washed aggregate in accordance with the FOP for AASHTO T 255, and then cool prior to sieving. Record the dry mass.

10. Select sieves to furnish information required by the specifications. Nest the sieves in order of decreasing size from top to bottom and place the sample, or a portion of the sample, on the top sieve.

11. Place sieves in mechanical shaker and shake for a minimum of 10 minutes, or the minimum time determined to provide complete separation for the sieve shaker being used.

12. Determine the individual or cumulative mass retained on each sieve and the pan to the nearest 0.1 percent or 0.1 g.

**WSDOT Note 5:** Use coarse wire brushes to clean the No. 40 600 µm (No. 30) and larger sieves, and soft bristle brushes for smaller sieves.

**Calculations**

The total mass of material after sieving should be verified with the mass before sieving. If performing T 11 with T 27 this would be the dry mass after wash. If performing just T 27 this would be the original dry mass. When the masses before and after sieving differ by more than 0.3 percent do not use the results for acceptance purposes. When performing the gradation from HMA using T 308, the masses before and after sieving shall not differ by more than 0.2%.

Calculate the total percentages passing, individual or cumulative percentages retained, or percentages in various size fractions to the nearest 0.1 percent by dividing the masses for method A, or adjusted masses for methods B and C, on the individual sieves by the total mass of the initial dry sample. If the same test sample was first tested by T 11, use the total dry sample mass prior to washing in T 11 as the basis for calculating all percentages. Report percent passing as indicated in the “Report” section at the end of this FOP.

**Percent Retained:**

Where:

- \( IPR \) = Individual Percent Retained
- \( CPR \) = Cumulative Percent Retained
- \( M \) = Total Dry Sample mass before washing
- \( IMR \) = Individual Mass Retained OR Adjusted Individual mass from Methods B or C
- \( CMR \) = Cumulative Mass Retained OR Adjusted Individual mass From Methods B or C

OR
## Percent Passing (Calculated):

Where:

\[
PP = \text{Percent Passing}
\]

\[
PPP = \text{Previous Percent Passing}
\]

\[
PP = PPP - IPR \quad \text{OR} \quad PP = 100 - CPR
\]

Calculate cumulative percent retained on and passing each sieve on the basis of the dry mass of total sample, before washing. This will include any material finer than No. 200 (75 µm) that was washed out.

Divide the cumulative masses, or the corrected masses, on the individual sieves by the total mass of the initial dry sample (prior to washing) to determine the percent retained on and passing each sieve. Calculate the percent retained on and passing each sieve. Report percent passing as indicated in the “Report” section at the end of this FOP.

### Example

Dry mass of total sample, before washing: 3214.0g

Dry mass of sample, after washing out the No. 200 (75 µm) minus: 3085.1g

For the 1/2 sieve:

**Cumulative Mass retained on 1/2" sieve = 161.0g**

Cumulative % retained = \( \frac{161.0}{3214.0} \times 100 = 5.0\% \text{ retained} \)

% passing = 100 - 5.0 = 95% passing 1/2" sieve

### Gradation on All Screens

<table>
<thead>
<tr>
<th>Sieve Size in. (mm)</th>
<th>Cumulative Mass Retained g</th>
<th>Cumulative Percent Retained</th>
<th>Reported Percent Passing*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4 (19.0)</td>
<td>0</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>161.0</td>
<td>5.0</td>
<td>95</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td>642.0</td>
<td>20.0</td>
<td>80</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td>1118.3</td>
<td>34.8</td>
<td>65</td>
</tr>
<tr>
<td>**No. 6 (3.35)</td>
<td>1515.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No. 10 (2.0)</td>
<td>1914.7</td>
<td>59.6</td>
<td>40</td>
</tr>
<tr>
<td>No. 40 (0.425)</td>
<td>2631.6</td>
<td>81.9</td>
<td>18</td>
</tr>
<tr>
<td>No. 80 (0.210)</td>
<td>2862.7</td>
<td>89.1</td>
<td>11</td>
</tr>
<tr>
<td>No. 200 (0.075)</td>
<td>3051.1</td>
<td>94.9</td>
<td>5.1</td>
</tr>
<tr>
<td>Pan</td>
<td>3086.4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Test Validation: 3086.4 – 3085.1 / 3085.1 x 100 = 0.04 % which is within the 0.3 percent requirement and the results can be used for acceptance purposes.

* Report No. 200 (75 µm) sieve to 0.1 percent. Report all others to 1 percent.

** Intermediate sieve used to prevent overloading the U. S. No. 10 sieve.
**Procedure Method B**

1. Perform steps 1 thru 9 from the “Procedure Method A” then continue as follows:

2. Select sieves to furnish information required by the specifications. Nest the sieves in order of decreasing size from top to bottom through the No. 4 (4.75 mm) with a pan at the bottom to retain the minus No. 4 (4.75 mm).

   See WSDOT Note 2 above.

3. Place sieves in mechanical shaker and shake for a minimum of 10 minutes, or the minimum time determined to provide complete separation for the sieve shaker being used.

4. Determine the individual or cumulative mass retained on each sieve and the pan to the nearest 0.1 percent or 0.1 g. Ensure that all material trapped in the openings of the sieve are cleaned out and included in the mass retained.

   Note 4: Use coarse wire brushes to clean the No. 40 and larger sieves, and soft bristle brushes for smaller sieves.

5. Determine the mass retained on each sieve to the nearest 0.1 percent of the total mass or better.

6. Determine the mass of the material in the pan [minus No. 4 (4.75 mm)].

7. Reduce the minus No. 4 (4.75 mm) using a mechanical splitter in accordance with the FOP for AASHTO T 248 to produce a sample with a mass of 500 g minimum. Determine and record the mass of the minus No. 4 (4.75 mm) split.

8. Select sieves to furnish information required by the specifications. Nest the sieves in order of decreasing size from top to bottom through the No. 200 (75 µm) with a pan at the bottom to retain the minus No. 200 (75 µm).

9. Place sieves in mechanical shaker and shake for a minimum of 10 minutes, or the minimum time determined to provide complete separation for the sieve shaker being used.

10. Determine the individual or cumulative mass retained on each sieve and the pan to the nearest 0.1 percent or 0.1 g. Ensure that all material trapped in the openings of the sieve are cleaned out and included in the mass retained.

   Note 4: Use coarse wire brushes to clean the No. 40 and larger sieves, and soft bristle brushes for smaller sieves.

**Calculations**

Compute the “Adjusted Cumulative Mass Retained” of the size increment of the original sample as follows when determining “Cumulative Mass Retained”:

Divide the cumulative masses, or the corrected masses, on the individual sieves by the total mass of the initial dry sample (prior to washing) to determine the percent retained on and passing each sieve. Calculate the percent retained on and passing each sieve. Report percent passing as indicated in the “Report” section at the end of this FOP.

When material passing the No. 4 (4.75 mm) sieve is split and only a portion of that is tested, the proportionate share of the amount passing the No. 200 (75 µm) sieve must be added to the sample mass to obtain a corrected test mass. This corrected test mass is used to calculate the gradation of the material passing the No. 4 (4.75 mm) sieve.
C = \left( \frac{M_1 \times B}{M_2} \right) + D

where:

C = \text{Total cumulative mass retained of the size increment based on a total sample}

M_1 = \text{mass of fraction finer than No. 4 (4.75 mm) sieve in total sample}

M_2 = \text{mass of reduced portion of material finer than No. 4 (4.75 mm) sieve actually sieved}

B = \text{cumulative mass of the size increment in the reduced portion sieved.}

D = \text{cumulative mass of plus No. 4 (4.75 mm) portion of sample.}

Example:

Dry mass of total sample, before washing: 3214.0g

Dry mass of sample, after washing out the No. 200 (75 µm) minus: 3085.1g

Gradation on Coarse Screens

<table>
<thead>
<tr>
<th>Sieve Size in. (mm)</th>
<th>Cumulative Mass Retained g</th>
<th>Cumulative Percent Retained</th>
<th>Reported Percent Passing*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4 (19.0)</td>
<td>0</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>161.0</td>
<td>5.0</td>
<td>95</td>
</tr>
<tr>
<td>3/8 (9.50)</td>
<td>642.0</td>
<td>20.0</td>
<td>80</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td>1118.3</td>
<td>34.8</td>
<td>65</td>
</tr>
</tbody>
</table>

Pan = 1968.0

Test Validation: $1118.3 + 1968.0 - 3085.1 \times 100 = 0.04\%$ which is within the 0.3 percent requirement and the results can be used for acceptance purposes.

The actual mass of material passing the No. 4 (4.75 mm) sieve and retained in the pan is 1968.0 g. This is $M_1$.

The pan (1968.0 grams) was reduced in accordance with the FOP for AASHTO T 248, so that at least 500 g are available. In this case, the mass determined was 512.8 g. This is $M_2$.

Gradation on Fine Screens

<table>
<thead>
<tr>
<th>Sieve Size in. (mm)</th>
<th>Cumulative Mass Retained (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 4 (4.75)</td>
<td>0</td>
</tr>
<tr>
<td>No. 10 (2.00)</td>
<td>207.5</td>
</tr>
<tr>
<td>No. 40 (0.425)</td>
<td>394.3</td>
</tr>
<tr>
<td>No. 80 (0.210)</td>
<td>454.5</td>
</tr>
<tr>
<td>No. 200 (0.075)</td>
<td>503.6</td>
</tr>
<tr>
<td>Pan</td>
<td>512.8</td>
</tr>
</tbody>
</table>
Test Validation: $512.8 - 512.8 / 512.8 = 0.0 \%$ which is within the 0.3 percent requirement and the results can be used for acceptance purposes.

For the No. 10 sieve:

\[
\begin{align*}
M_1 &= 1968.0g \\
M_2 &= 512.8g \\
B &= 207.5g \\
D &= 1118.3g \\
C &= \left( \frac{M_1 \times B}{M_2} \right) + D = \left( \frac{1968.0g \times 207.5g}{512.8g} \right) + 1118.3g = 1914.7g \\
\end{align*}
\]

% retained $\frac{1914.7g}{3214.0g} = 59.6\%$

% passing = 100-59.6=40.4\% reported as 40\%

### Final Gradation on All Screens

<table>
<thead>
<tr>
<th>Sieve Size in. (mm)</th>
<th>Cumulative Mass Retained g</th>
<th>Adjusted Cumulative Mass Retained g</th>
<th>Cum.</th>
<th>Reported Percent Passing*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4 (19.0)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>100.0</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>161.1</td>
<td>161.1</td>
<td>5.0</td>
<td>95</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td>642.5</td>
<td>642.5</td>
<td>20.0</td>
<td>80</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td>1118.3</td>
<td>1118.3</td>
<td>34.8</td>
<td>65</td>
</tr>
<tr>
<td>No. 10 (2.0)</td>
<td>207.5 x 3.838 +</td>
<td>1914.7</td>
<td>59.6</td>
<td>40</td>
</tr>
<tr>
<td>No. 40</td>
<td>394.3 x 3.838 +</td>
<td>2631.6</td>
<td>81.6</td>
<td>18</td>
</tr>
<tr>
<td>No. 80</td>
<td>454.5 x 3.838 +</td>
<td>2862.7</td>
<td>89.1</td>
<td>11</td>
</tr>
<tr>
<td>No. 50</td>
<td>503.6 x 3.838 +</td>
<td>3051.1</td>
<td>94.9</td>
<td>5.1</td>
</tr>
<tr>
<td>Pan</td>
<td>512.8 x 3.838 +</td>
<td>3086.4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Report No. 200 (75 µm) sieve to 0.1 percent. Report all others to 1 percent.

### Alternative Method B

As an alternate method to account for the fact that only a portion of the minus No. 4 (4.75 mm) material was sieved, multiply the fine screen “Percent Passing” values by the percent passing the No. 4 (4.75 mm) sieve obtained in the coarse screen procedure, 65 percent in this case.

The mass retained in the pan must be corrected to include the proper percent of No. 200 (.075 mm) minus material washed out.

Divide the cumulative masses, or the corrected masses, on the individual sieves by the corrected pan mass of the initial dry sample (prior to washing) to determine the percent retained on and passing each sieve. Calculate the percent retained on and passing each sieve. Report percent passing as indicated in the “Report” section at the end of this FOP.

Dry mass of total sample, before washing: 3214.0g

Dry mass of sample, after washing out the No. 200 (75 µm) minus: 3085.1g

Amount of No. 200 (75 µm) minus washed out: $3214.0 \text{ g} - 3085.1 \text{ g} = 128.9g$
Gradation on Coarse Screens

<table>
<thead>
<tr>
<th>Sieve Size in. (mm)</th>
<th>Cumulative Mass Retained g</th>
<th>Cumulative Percent Retained</th>
<th>Reported Percent Passing*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4 (19.0)</td>
<td>0</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>161.0</td>
<td>5.0</td>
<td>95</td>
</tr>
<tr>
<td>3/8 (9.50)</td>
<td>642.0</td>
<td>20.0</td>
<td>80</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td>1118.3</td>
<td>34.8</td>
<td>65</td>
</tr>
</tbody>
</table>

Pan = 1968.0

Test Validation: 1118.3 + 1968.0 - 3085.1 / 3085.1 x 100 = 0.04% which is within the 0.3 percent requirement and the results can be used for acceptance purposes.

The actual mass of material passing the No. 4 (4.75 mm) sieve and retained in the pan is 1968.0 g. This is M₃.

The pan (1968.0 grams) was reduced in accordance with the FOP for AASHTO T 248, so that at least 500 g are available. In this case, the mass determined was 512.8 g. This is M₄.

Corrected pan mass = M₄ \left( \frac{M₄}{M₃} \right) \left( \frac{C₁}{M₃} \right)

Where:

M₄ = mass retained in the pan from the split of the No. 4 (4.75 mm) minus.

M₃ = mass of the No. 4 (4.75 mm) minus of entire sample, not including No. 200 (.075 mm) minus washed out.

C₁ = mass of No. 200 (.075 mm) minus washed out.

The corrected pan mass is the mass used to calculate the percent retained for the fine grading.

Example:

M₄ = 512.8g

M₃ = 1968.0g

C₁ = 128.9g

Corrected pan mass = 512.8g + \frac{512.8g \times 128.9g}{1968.0g} = 546.4g
For the No. 10 sieve:

Mass of No. 10 sieve = 207.5g

Corrected Pan Mas = 546.4g

Cumulative % retained = \( \frac{207.5g}{546.4g} \times 100 = 38.0\% \)

% passing = 100-38.0 = 62.0%

Adjusted % passing No. 10 = % passing No. 10 x % No. 4 = 62.0 x 0.65 = 40%

Final Gradation on All Screens

<table>
<thead>
<tr>
<th>Sieve Size in. (mm)</th>
<th>Adjustment</th>
<th>Reported Percent Passing*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4 (19.0)</td>
<td></td>
<td>100</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td></td>
<td>95</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td></td>
<td>80</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td>100 x .65 =</td>
<td>65</td>
</tr>
<tr>
<td>No. 10 (2.00)</td>
<td>62.0 x .65 =</td>
<td>40</td>
</tr>
<tr>
<td>No. 40 (0.425)</td>
<td>27.8 x .65 =</td>
<td>18</td>
</tr>
<tr>
<td>No. 80 (0.210)</td>
<td>16.8 x .65 =</td>
<td>11</td>
</tr>
<tr>
<td>No. 200 (0.075)</td>
<td>7.8 x .65 =</td>
<td>5.1</td>
</tr>
</tbody>
</table>

• Report No. 200 (75 µm) sieve to 0.1 percent. Report all others to 1 percent.

• SAMPLE CALCULATION FOR FINENESS MODULUS

Fineness Modulus (FM) is used in determining the degree of uniformity of aggregate gradation in PCC mix designs. It is an empirical number relating to the fineness of the aggregate. The higher the FM, the coarser the aggregate. Values of 2.40 to 3.00 are common for FA in PCC.

The FM is the sum of the percentages retained on specified sieves 150 mm (6”), 75 mm (3”), 37.5 mm (1/2”), 19.0 mm (3/4”), 9.5 mm (3/8”), No. 4 (4.75 mm), 2.36 mm (No. 8), 1.18 mm (No. 16), 0.60 mm (No. 30), 0.30 mm (No. 50), and 0.15 mm (No. 100) divided by 100 gives the FM.

The following example is for WSDOT Class 2 Sand:

<table>
<thead>
<tr>
<th>Sieve</th>
<th>Size</th>
<th>% Passing</th>
<th>% Retained on Specified Sieves</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 in.</td>
<td>150 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>3 in.</td>
<td>75 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>2-1/2 in.</td>
<td>62.5 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>2 in.</td>
<td>50 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>1-1/2 in.</td>
<td>37.5 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>1 in.</td>
<td>25 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>3/4 in.</td>
<td>19 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>½ in.</td>
<td>12.5 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>3/8 in.</td>
<td>9.5 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>No. 4</td>
<td>4.75 mm</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>No. 8</td>
<td>2.36 mm</td>
<td>87</td>
<td>13</td>
</tr>
<tr>
<td>No. 16</td>
<td>1.18 mm</td>
<td>69</td>
<td>31</td>
</tr>
<tr>
<td>No. 30</td>
<td>0.60 mm</td>
<td>44</td>
<td>56</td>
</tr>
<tr>
<td>No. 50</td>
<td>0.30 mm</td>
<td>18</td>
<td>82</td>
</tr>
<tr>
<td>No. 100</td>
<td>0.15 mm</td>
<td>4</td>
<td>96</td>
</tr>
</tbody>
</table>

= 278
FM = 2.78
Report

Results shall be reported on standard forms approved for use by the agency. Depending on the agency, this may include:

- Cumulative mass retained on each sieve*
- Cumulative percent retained on each sieve*
- Percent passing and retained on each sieve shall be reported to the nearest 1 percent except for the percent passing the U.S. No. 200 (75 µm) sieve, which shall be reported to the nearest 0.1 percent
- FM to the nearest 0.01 percent for WSDOT Class 2 Sand

Report results using WSDOT Form 422-020, or other report approved by the State Materials Engineer.
Performance Exam Checklist

**WAQTC FOP FOR AASHTO T 27/T 11**

**SIEVE ANALYSIS OF FINE AND COARSE AGGREGATES**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>and if required, has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Minimum sample mass meets requirement of Table 1 or from FOP for AASHTO T308?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Test sample dried to a constant mass by FOP for AASHTO T 255?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Test sample cooled and mass determined to nearest 0.1 percent of mass?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Sample placed in container and covered with water?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(If specification requires that the amount of material finer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>than the No. 200 sieve is to be determined.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Dispersing Agent used for HMA?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Contents of the container vigorously agitated?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Complete separation of coarse and fine particles achieved?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Wash water poured through nested sieves such as No. 10 and No. 200?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Operation continued until wash water is clear?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12. Material retained on sieves returned to washed sample?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. Washed aggregate dried to a constant mass by FOP for AASHTO T 255?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14. Washed aggregate cooled and mass determined to nearest 0.1 percent of mass?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15. Sample placed in nest of sieves specified? (Additional sieves</td>
<td></td>
<td></td>
</tr>
<tr>
<td>may be used to prevent overloading as allowed in FOP.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16. Material sieved in verified mechanical shaker for minimum of 10 minutes?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17. Mass of residue on each sieve determined to 0.1 percent of mass?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18. Total mass of material after sieving agrees with mass before sieving to</td>
<td></td>
<td></td>
</tr>
<tr>
<td>within 0.3 percent, or 0.2 percent for HMA (per FOP for AASHTO T308)?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>19. Percentages calculated to the nearest 0.1 percent and reported to the</td>
<td></td>
<td></td>
</tr>
<tr>
<td>nearest whole number, except No. 200 - reported to the nearest 0.1 percent?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20. Percentage calculations based on original dry sample mass?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>21. Calculations performed properly? If material passing No. 4 sieve is split</td>
<td></td>
<td></td>
</tr>
<tr>
<td>and only a portion is tested, calculation as noted in FOP performed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>properly?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
First attempt: Pass ☐ Fail ☐  Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

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**WSDOT FOP FOR WAQTC/ AASHTO T 40**

**SAMPLING BITUMINOUS MATERIALS**

**SIGNIFICANCE**

The quality of bituminous materials has a tremendous impact on a roadway project. The grade of binder selected is based on a number of factors, including local temperature extremes and characteristics of expected traffic. Using a grade of binder material other than that specified will have serious impacts on roadway performance and durability.

**SCOPE**

The procedure covers obtaining samples of liquid bituminous materials in accordance with AASHTo T 40. Sampling of solid and semi-solid bituminous materials (included in AASHTo T 40) is not covered here.

Agencies may be more specific on exactly who samples, where to sample, and what type of sampling device to use.

WSDOT personnel need to observe the contractor’s personnel sampling to assure that proper sampling procedures are followed.

If proper sampling procedures are not followed it shall be noted on the sample transmittal “Proper sampling procedures not followed.” See WSDOT Standard Specification 1-06.

**PROCEDURE**

1. Coordinate sampling with contractor or supplier.
2. Use appropriate safety equipment and precautions.
3. Allow a minimum of 1 gal (4 L) to flow before obtaining samples.
4. Obtain samples of:
   - Asphalt binder from Hot Mix Asphalt (HMA) Plant from the line between the storage tank and the mixing plant or the storage tank while the plant is in operation, or from the delivery truck.
   - Cutback and Emulsified asphalt from distributor spray bar or application device; or from the delivery truck before it is pumped into the distributor. Sample emulsified asphalt at delivery or prior to dilution.

**CONTAINERS**

Sample containers must be new, and the inside may not be washed or rinsed. The outside may be wiped with a clean, dry cloth.

All samples shall be put in 1 qt (1 L) containers and properly identified on the outside of the container with contract number, date sampled, data sheet number, brand and grade of material, and sample number. Include lot and sublot numbers when appropriate.

Note: The filled sample container shall not be submerged in solvent, nor shall it be wiped with a solvent saturated cloth. If cleaning is necessary, use a clean dry cloth.

- Emulsified asphalt: Use wide-mouth plastic jars with screw caps. Protect the samples from freezing since water is a part of the emulsion.
- Asphalt binder & Cutbacks: Use metal cans.

Standard sample labels (WSDOT Form 350-016) shall be completely filled out and attached to each sample container.
Performance Exam Checklist

Sampling Bituminous Materials
WAQTC FOP for AASHTO T 40

Participant Name ___________________________ Exam Date ________________

Procedure Element

1. The tester has a copy of the current procedure on hand? Yes ☐ No ☐

2. Appropriate containers used?
   a. Wide-mouth plastic containers (emulsified). ☐ ☐
   b. Metal cans (all other bituminous liquids). ☐ ☐

3. Containers not washed or rinsed on inside? ☐ ☐

4. Minimum of 1 gallon allowed to flow before sample taken? ☐ ☐

5. Material obtained at correct location?
   a. Line between storage tank and mixing plant or the storage tank (HMA plants). ☐ ☐
   b. Spray bar or application device, if not diluted (distributors). ☐ ☐
   c. From delivery vehicle or prior to dilution, if diluted (distributors). ☐ ☐

Sample taken by: Contractor ☐ WSDOT ☐

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner ____________________________

Comments:

________________________________________________________________________
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WSDOT FOP for AASHTO TP 61

Determining the Percentage of Fracture in Coarse Aggregate

1. SCOPE

1.1. This test method covers the determination of the percentage, by mass, of a coarse aggregate sample that consists of fractured particles meeting specified requirements.

1.2. This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety problems 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.

1.3. The text of the standard reference notes provide explanatory material. These notes (excluding those in tables and figures) shall not be considered as requirements of the standard.

Method 1 will be used by WSDOT for determining the fracture of aggregate as required by the Standard Specifications.

2. REFERENCED DOCUMENTS

2.1. AASHTO Standards:

- M 92, Wire-Cloth Sieves for Testing Purposes
- M 231, Weighing Devices Used in the Testing of Materials
- T 2, Sampling of Aggregates
- T 11, Materials Finer Than 75-µm (No. 200) Sieve in Mineral Aggregates by Washing
- T 27, Sieve Analysis of Fine and Coarse Aggregates
- T 248, Reducing Samples of Aggregate to Testing Size
- T 255, Total Evaporable Moisture Content of Aggregate by Drying

3. SUMMARY OF TEST METHOD

3.1. A sample of aggregate is separated using the designated size of screen conforming to the specification controlling the determination of coarse and fine aggregate. The coarse aggregate particles are visually evaluated to determine their conformance to the defined fracture. The percentage of conforming particles, by mass, is determined for comparison to standard specifications.

4. APPARATUS

4.1. Balance—shall have sufficient capacity, be readable to 0.1 percent of the sample mass, or better, and conform to Meeting the requirements of M 231 for general-purpose balance required for the principle sample mass being tested.

4.2. Sieves—Meeting the requirements of M 92.

4.3. Splitter—Meeting the requirements of T 248.

This Test Procedure is based on AASHTO T 61-02
5. TERMINOLOGY

5.1. *fractured face*—an angular, rough, or broken surface of an aggregate particle created by crushing, or by other means. A face is considered a “fractured face” whenever one-half or more of the projected area, when viewed normal to that face, is fractured with sharp and well-defined edges: this excludes small nicks.

5.2. *fractured particle*—a particle of aggregate having at least the minimum number of fractured faces specified (usually one or two).

6. SAMPLING

Sample the aggregate in accordance with FOP for AASHTO T 2 and reduce the sample in accordance with FOP for AASHTO T 248, to the sample sizes shown in Table 1 of FOP for AASHTO T 27/11.

7. SAMPLE PREPARATION

7.1. Where the specifications list only a total fracture percentage, the sample shall be prepared in accordance with Method 1. When the specifications require that the fracture be counted and reported on each sieve, the sample shall be prepared in accordance with Method 2.

7.2. *Method 1*—Combined Fracture Determination

7.2.1. Dry the sample sufficiently to obtain a clean separation of fine and coarse material in the sieving operation. Sieve the sample in accordance with FOP for AASHTO T 27/11 over the No. 4 (4.75-mm) sieve, or the appropriate sieve listed in the agency specifications for this material.

*Note 1*—Where necessary, wash the sample over the sieve or sieves designated for the determination of fractured particles to remove any remaining fine material, and dry to a constant mass in accordance with FOP for AASHTO T 255.
7.2.2. Reduce the sample using a splitter in accordance with FOP for AASHTO T 248 to the appropriate size for test. This size of test sample should be slightly larger in mass than that shown in Table 1, to account for additional loss of fines after washing.

Table 1—Sample Size (Method 1, Combined Sieve Fracture)

<table>
<thead>
<tr>
<th>Nominal Maximum Particle Size</th>
<th>Minimum Sample Mass Retained No. 4 (4.75-mm) Sieve</th>
</tr>
</thead>
<tbody>
<tr>
<td>1½ in (37.5 mm)</td>
<td>6 lb (2500 g)</td>
</tr>
<tr>
<td>1 in (25 mm)</td>
<td>3.5 lb (1500 g)</td>
</tr>
<tr>
<td>¾ in (19.0 mm)</td>
<td>2.5 lb (1000 g)</td>
</tr>
<tr>
<td>⅜ in (16.0 mm)</td>
<td>2.0 lb (800 g)</td>
</tr>
<tr>
<td>⅜ in (12.5 mm)</td>
<td>1.5 lb (700 g)</td>
</tr>
<tr>
<td>⅜ in (9.5 mm)</td>
<td>0.9 lb (400 g)</td>
</tr>
<tr>
<td>No. 4 (4.75 mm)</td>
<td>0.4 lb (200 g)</td>
</tr>
</tbody>
</table>

* For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.

Note: For an aggregate specification having a generally unrestricted gradation (i.e. wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.

7.3. Method 2—Individual Sieve Fracture Determination WSDOT has deleted this section

8. PROCEDURE

8.1. Spread the dried cooled test sample on a clean flat surface large enough to permit careful inspection of each particle. To verify that a particle meets the fracture criteria, hold the aggregate particle so that the face is viewed directly. (See Section 5.1.)

8.2. To aid in making the fracture determination separate the sample into three categories: (1) fractured particles meeting the above criteria, (2) particles not meeting specification criteria, and (3) questionable or borderline particles.

8.3. Determine the mass of particles in the fractured category, the mass of questionable particles, and the mass of the unfractured particles.

8.4. If on any of the determinations, more than 15 percent of the total mass of the sample is placed in the questionable category, repeat the determination until no more than 15 percent is present in that category.
9. **CALCULATION REPORT**

9.1. *Report the following information:*

9.1.1. Calculate the mass percentage of fracture faces to the nearest 1 percent as follows:

\[ P = \left( \frac{F + Q/2}{F + Q + N} \right) \times 100 \]  

where:

\[ P \] = percent of fracture,

\[ F \] = mass of fractured particles,

\[ Q \] = mass of questionable or borderline particles, and

\[ N \] = mass of unfractured particles.

**REPORT**

Results shall be reported on standard forms approved for use by the agency. Report fracture to the nearest 1 percent.

Report the results using WSDOT Form 350-161 EF, 422-020X, or other report approved by the State Materials Engineer.

10. **PRECISION AND BIAS**

10.1. *Precision*—The research required to determine the precision of this standard has not been performed.

10.2. *Bias*—The research required to determine the bias of this standard has not been performed.
## Performance Exam Checklist

*Determining the Percentage of Fracture In Coarse Aggregate*

*WSDOT FOP for AASHTO TP 61*

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Sample reduced to correct size?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Sample dried and cooled, if necessary?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Sample properly sieved through specified sieve(s)?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Particles separated into fractured, unfractured, and questionable categories?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Dry mass of each category determined to nearest 0.1 g?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Calculation performed correctly?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐  Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

**Comments:**

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WSDOT FOP for AASHTO T 99¹

Moisture-Density Relations of Soils Using a 5.5-lb (2.5-kg) Rammer and a 12-in. (305-mm) Drop

1. SCOPE

1.1 These methods of test are intended for determining the relation between the moisture content and density of soils compacted in a mold of a given size with a 5.5-lb (2.5-kg) rammer dropped from a height of 12-in. (305-mm). Four alternate procedures are provided as follows:

   **Method A**
   A 4-in. (101.60-mm) mold: Soil material passing a No. 4 (4.75-mm) sieve
   Sections 3 and 4.

   **Method B**
   A 6-in. (152.40-mm) mold: Soil material passing a No. 4 (4.75-mm) sieve
   Sections 5 and 6.

   **Method C**
   A 4-in. (101.60-mm) mold: Soil material passing a 3/4-in. (19.0-mm) sieve
   Sections 7 and 8.

   **Method D**
   A 6-in. (152.40-mm) mold: Soil material passing a 3/4-in. (19.0-mm) sieve
   Sections 9 and 10.

   The preferred method of WSDOT is to use Method A.

   WSDOT recommends that the bulk specific gravity of coarse aggregate be determined.

   Native soils within the contract limits to be used for embankment construction and/or backfill material do not require the sampling by a qualified tester. For material that requires gradation testing such as but not limited to manufactured aggregates and Gravel Borrow, a qualified testers shall be required for sampling.

1.2 The method to be used should be indicated in the specifications for the material being tested. If no method is specified, the provisions of Method A shall govern.

1.3. This test method applies to soils mixtures that have 40% or less retained on the No. 4 (4.75 mm) sieve, when Method A or B is used and 30% or less retained on the 3/4-in. (19.0-mm) sieve, when Method C or D is used. The material retained on these sieves shall be defined as oversized particles (coarse particles).

1.4. If the test specimen contains oversize particles, and the test specimen is used for field density compaction control, corrections must be made according to T 224 to compare the total field density with the compacted specimen density. The person or agency specifying this method shall specify a minimum percentage below which correction for oversize need not be applied. If no minimum percentage is specified, correction shall be applied to samples with more than 5% by weight of oversize particles.

1.5. If the specified oversized maximum tolerances are exceeded, other methods of compaction control must be used.

¹This Test Method is based on AASHTO T 99-01
Note 1—One method for the design and control of the compaction of such soils is to use a test fill to determine the required degree of compaction and a method to obtain that compaction. Then use a method specification to control the compaction by specifying the type and size of compaction equipment, the lift thickness and the number of passes.

1.6. The following applies to all specified limits in this standard: For the purposes of determining conformance with these specifications, an observed value or a calculated value shall be rounded off “to the nearest unit” in the last right-hand place of figures used in expressing the limiting value, in accordance with R 11.

1.7. The values stated in SI units are to be regarded as the standard.

2. Referenced Documents

2.1. AASHTO Standards:

• M 92, Wire-Cloth Sieves for Testing Purposes
• M 231, Weighing Devices Used in the Testing of Materials
• R 11, Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values
• T 19/T 19M, Bulk Density(“Unit Weight”) and Voids in Aggregate
• T 224, Correction for Coarse Particles in the Soil Compaction Test
• T 255, Total Evaporable Moisture Content of Aggregate by Drying
• T 265, Laboratory Determination of Moisture Content of Soils

2.2. ASTM Standard:

• D 2168, Calibration of Laboratory Mechanical-Rammer Soil Compactors

3. APPARATUS

3.1 Molds — The molds shall be solid-wall, metal cylinders manufactured with dimensions and capacities shown in Sections 3.1.1 and 3.1.2 below. They shall have a detachable collar assembly approximately 2.375 in. (60 mm) in height, to permit preparation of compacted specimens of soil-water mixtures of the desired height and volume. The mold and collar assembly shall be so constructed that it can be fastened firmly to a detachable base plate made of the same material (Note 2). The base plate shall be plane to 0.005 in. as shown in Figures 1 and 2.

Note 2: Alternate types of molds with capacities as stipulated herein may be used, provided the test results are correlated with those of the solid-wall mold on several soil types and the same moisture-density results are obtained. Records of such correlation shall be maintained and readily available for inspection, when alternate types of molds are used.

3.1.1 A 4-in. (101.6-mm) mold having a capacity of 1/30 (0.0333) ± 0.0003 cu. ft. (0.000943 ± 0.000008 m³) with an internal diameter of 4.000 ± 0.016 in. (101.60 ± 0.41 mm) and a height of 4.584 ± 0.005 in. (116.43 ± 0.13 mm) (Figure 1).

3.1.2 A 6-in. (152.4-mm) mold having a capacity of 1/13.33 (0.07500) ± 0.00075 cu. ft. (0.002124 ± 0.000021 m³) with an internal diameter of 6.000 ± 0.026 in. (152.40 ± 0.66 mm) and a height of 4.584 ± 0.005 in. (116.43 ± 0.13 mm) (Figure 2).
Cylindrical Mold and Base Plate (101.6-mm mold)

Figure 1
Figure 2

Cylindrical Mold and Base Plate (152.4-mm mold)

(A) WING NUT (4)
(B) STUD (2)
(C) HANGER (4)
(D) WELD (Top and bottom of each hanger)
(E) COLLAR (1)
(F) MOLD (1)
(G) BASE PLATE (1)

NOTE:
ALL DIMENSIONS SHOWN IN MILLIMETERS UNLESS OTHERWISE NOTED.

LOCATION OF STUDS IN BASE PLATE

<table>
<thead>
<tr>
<th>Dimensional Equivalents</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>mm</strong></td>
</tr>
<tr>
<td>3.18 ± 0.64</td>
</tr>
<tr>
<td>3.81</td>
</tr>
<tr>
<td>6.35 ± 1.27</td>
</tr>
<tr>
<td>7.62</td>
</tr>
<tr>
<td>9.53 ± 0.64</td>
</tr>
<tr>
<td>12.70 ± 2.54</td>
</tr>
<tr>
<td>17.78 ± 1.27</td>
</tr>
<tr>
<td>20.32</td>
</tr>
<tr>
<td>38.10 ± 2.54</td>
</tr>
</tbody>
</table>

0.002123 ± 0.000021 m³ /13.33 ± 0.00075 ft³
3.1.3 Molds Out of Tolerance Due to Use — A mold that fails to meet manufacturing tolerances after continued service may remain in use provided those tolerances are not exceeded by more than 50 percent; and the volume of the mold, calibrated in accordance with Section 8 (Calibration of Measure) of T 19/T 19M, for Unit Mass of Aggregate, is used in the calculations.

3.2 Rammer

3.2.1 Manually Operated — Metal rammer with a mass of 5.5 ± 0.02 lb (2.495 ± 0.009 kg), and having a flat circular face of 2.000-in. (50.80-mm) diameter with a manufacturing tolerance of 0.01 in. (± 0.25 mm). The in-service diameter of the flat circular face shall be not less than 1.985 in. (50.42 mm). The rammer shall be equipped with a suitable guide-sleeve to control the height of drop to a free fall of 12.00 ± 0.06 in. (305 ± 2 mm) above the elevation of the soil. The guide-sleeve shall have at least 4 vent holes, no smaller than 3/8-in. (9.5-mm) diameter spaced approximately 90 degrees (1.57 rad) apart and approximately 3/4 in. (19 mm) from each end; and shall provide sufficient clearance so the free fall of the rammer shaft and head is unrestricted.

3.2.2 Mechanically Operated — A metal rammer which is equipped with a device to control the height of drop to a free fall of 12.00 ± 0.06 in. (305 ± 2 mm) above the elevation of the soil and uniformly distributes such drops to the soil surface (Note 3). The rammer shall have a mass of 5.5 ± 0.02 lb (2.495 ± 0.009 kg), and have a flat circular face of 2.000-in. (50.80 mm) diameter with a manufactured tolerance of 0.01 in. (± 0.25 mm). The in-service diameter of the flat circular face shall be not less than 1.985 in. (50.42 mm). The mechanical rammer shall be calibrated by ASTM D 2168.

Note 3: It may be impractical to adjust the mechanical apparatus so the free fall is 12 in. (305 mm) each time the rammer is dropped, as with the manually operated rammer. To make the adjustment of free fall, the portion of loose soil to receive the initial blow should be slightly compressed with the rammer to establish the point of impact from which the 12 in. (305 mm) drop is determined. Subsequent blows on the layer of soil being compacted may all be applied by dropping the rammer from a height of 12 in. (305 mm) above the initial-setting elevation; or, when the mechanical apparatus is designed with a height adjustment for each blow, all subsequent blows should have a rammer free fall of 12 in. (305 mm) measured from the elevation of the soil as compacted by the previous blow. A more detailed calibration procedure for laboratory mechanical-rammer soil compactors can be found in ASTM D 2168.

3.2.3 Rammer Face — The circular face rammer shall be used but a sector face may be used as an alternative provided the report shall indicate type of face used other than the 2-in. (50.8-mm) circular face and it shall have an area equal to that of the circular face rammer.

3.3 Sample Extruder (for Solid-Walled Molds Only) — A jack, lever, frame, or other device adopted for the purpose of extruding compacted specimens from the mold.

3.4 Balances and Scales — A balance or scale conforming to the requirements of AASHTO M 231, Class G 20. Also, a balance conforming to the requirements of AASHTO M 231, Class G 2.

Note 4: The capacity of the metric balance or scale should be approximately 11.5 kg when used to weigh the 6-in. (152.40-mm) mold and compacted, moist soil; however, when the 4-in. (101.60-mm) mold is used, a balance or scale of lesser capacity than the 11.5 kg may be used, if the sensitivity and readability is 5 g.
3.5 Drying Oven — A thermostatically controlled drying oven capable of maintaining a temperature of 230 ± 9°F (110 ± 5°C) for drying moisture samples.

3.6 Straightedge — A hardened-steel straightedge at least 10 in. (250 mm) in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.01 in. per 10 in. (0.250 mm per 250 mm) (0.1 percent) of length within the portion used for trimming the soil (Note 5).

Note 5: The beveled edge may be used for final trimming if the edge is true within a tolerance of 0.01 in. per 10 in. (0.250 mm per 250 mm) (0.1 percent) of length; however, with continued use, the cutting edge may become excessively worn and not suitable for trimming the soil to the level of the mold. The straightedge should not be so flexible that trimming the soil with the cutting edge will cause a concave soil surface.

3.7 Sieves — 2-in. (50-mm), 3/4-in. (19.0-mm), and No. 4 (4.75-mm) sieves conforming to the requirements of M 92.

3.8 Mixing Tools — Miscellaneous tools such as mixing pan, spoon, trowel, spatula, etc., or a suitable mechanical device for thoroughly mixing the sample of soil with increments of water.

3.9 Containers — Suitable containers made of material resistant to corrosion and not subject to change in mass or disintegration on repeated heating and cooling. Containers shall have close-fitting lids to prevent loss of moisture from samples before initial mass determination and to prevent absorption of moisture from the atmosphere following drying and before final mass determination. One container is needed for each moisture content determination.

### METHOD A

4. SAMPLE

4.1 If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus which is maintained at a temperature not exceeding 140°F (60°C). Then thoroughly break up the aggregations in such a manner as to avoid reducing the natural size of individual particles.

4.2 Sieve an adequate quantity of the representative pulverized soil over the No. 4 (4.75-mm) sieve. Discard the coarse material, if any, retained on the No. 4 (4.75-mm) sieve.

4.3 Select a representative sample, with a mass of approximately 7 lb (3 kg) or more, of the soil prepared as described in Sections 4.1 and 4.2.

Note 6—When developing a compaction curve for free draining soils, such as uniform sands and gravels, where seepage occurs at the bottom of the mold and base plate, taking a representative moisture content sample from the mixing bowl may be preferred in order to determine the amount of moisture available for compaction.

5. PROCEDURE

5.1 Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately four percentage points below optimum moisture content.

5.2 Form a specimen by compacting the prepared soil in the 4-in. (101.60-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5 in. (125 mm). Prior to compaction, place the loose soil into the mold and spread into a layer of uniform thickness. Lightly tamp the soil prior to compaction until it is not in a loose or fluffy state, using either the manual compaction rammer or similar device having a face diameter of approximately 2 in. (50 mm). Following compaction of each of the first two layers, any soil
adjacent to the mold walls that has not been compacted or extends above the compacted surface shall be trimmed using a knife or other suitable device, and be evenly distributed on top of the layer. Compact each layer by 25 uniformly distributed blows from the rammer dropping free from a height of 12 in. (305 mm) above the elevation of the soil when a sleeve-type rammer is used, or from 12 in. (305 mm) above the approximate elevation of compacted soil when a stationary mounted type of rammer is used. During compaction, the mold shall rest firmly on a dense, uniform, rigid, and stable foundation or base. This base shall remain stationary during the compaction process (Note 7).

Note 7: Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: A block of concrete, with a mass not less than 200 lb (90 kg), supported by a relatively stable foundation; a sound concrete floor; and for field application, such surfaces as are found in concrete box culverts, bridges, and pavements.

5.2.1 Following compaction, remove the extension collar, carefully trim the compacted soil even with the top of the mold by means of the straightedge, and determine the mass of the mold and moist soil in kilograms to the nearest 5 grams, or determine the mass in pounds to the nearest 0.01 pounds. For molds conforming to tolerances given in Section 3.1.1 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 1060, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in Section 3.1.1 and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 30, and record the result as the wet density, \( W_1 \), in pounds per cubic foot, of compacted soil. For used molds out of tolerance by not more than 50 percent (Section 3.1.3), use the factor for the mold as determined in accordance with Calibration of Measure in AASHTO T 19/T 19M.

5.3 Remove the material from the mold and slice vertically through the center. Take a representative sample of the material from one of the cut faces, weigh immediately and dry in accordance with T 255 or T 265, to determine the moisture content, and record the results.

5.4 Thoroughly break up the remaining portion of the molded specimen until it will pass a No. 4 (4.75-mm) sieve as judged by eye, and add to the remaining portion of the sample being tested. Add water in sufficient amount to increase the moisture content of the soil one to two percentage points (water content increments should not exceed 2.5 percent except when heavy clay soils or organic soils exhibiting flat elongated curves are encountered, the water content increments may be increased to a maximum of 4 percent), and repeat the above procedure for each increment of water added. Continue this series of determinations until there is either a decrease or no change in the wet unit mass, \( W_1 \), per cubic foot (cubic meter) of the compacted soil (Note 8).

Note 8: This procedure has been found satisfactory in most cases. However, in instances where the soil material is fragile in character and will reduce significantly in grain size due to repeated compaction, and in cases where the soil is a heavy-textured clayey material into which it is difficult to incorporate water, a separate and new sample shall be used in each compaction test. In these cases, separate samples shall be thoroughly mixed with amounts of water sufficient to cause the moisture contents of the samples to vary by approximately two percentage points. The moisture points selected shall bracket the optimum moisture content, thus providing samples which, when compacted, will increase in mass to the maximum density and then decrease in mass. The samples of soil-water mixtures shall be placed in covered containers and allowed to stand for not less than 12 hours before making the moisture-density test.
5.4.1 In instances where the soil material is fragile in character and will be reduced significantly in grain size by repeated compaction, a separate and new sample shall be used in each compaction test.

METHOD B

6. SAMPLE

6.1 Select the representative sample in accordance with Section 3.3, except that it shall have a mass of approximately 16 lb (7 kg).

7. PROCEDURE

7.1 Follow the same procedure as described for Method A in Section 4, except for the following: Form a specimen by compacting the prepared soil in the 6-in. (152.4-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5 in. (125 mm), each layer being compacted by 56 uniformly distributed blows from the rammer. For molds conforming to tolerances given in Section 2.1.2, and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 471, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in Section 2.1.2, and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 13.3, and record the result as the wet density, \( W_1 \), in pounds per cubic foot, of compacted soil. For used molds out of tolerance by not more than 50 percent (Section 3.1.3), use the factor for the mold as determined in accordance with Calibration of Measure in 19/T 19M.

METHOD C

8. SAMPLE

8.1 If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus which is maintained at a temperature not exceeding 140°F (60°C). Then thoroughly break up the aggregations in such a manner as to avoid reducing the natural size of individual particles.

8.2 Sieve an adequate quantity of the representative pulverized soil over the 19.0-mm sieve. Discard the coarse material, if any, retained on the \( \frac{3}{4} \) in. (19.0-mm) sieve. (Note 9).

Note 9: If it is advisable to maintain the same percentage of coarse material (passing a 2 in. (50-mm) sieve and retained on a No. 4 (4.75-mm) sieve) in the moisture-density sample as in the original field sample, the material retained on the \( \frac{3}{4} \) in. (19.0-mm) sieve shall be replaced as follows: Sieve an adequate quantity of the representative pulverized soil over the 2 in. - \( \frac{3}{4} \) in. (50- and 19.0-mm) sieves. Determine the mass of the material passing the 2 in. (50-mm) sieve and retained on the \( \frac{3}{4} \) in. (19.0-mm) sieve and replace it with an equal mass of material passing the \( \frac{3}{4} \) in. (19.0-mm) sieve and retained on the No. 4 (4.75-mm) sieve. Take the material for replacement from the remaining portion of the sample.

8.3 Select a representative sample, having a mass of approximately 11 lb (5 kg) or more, of the soil prepared as described in Sections 8.1 and 8.2.
9. **PROCEDURE**

9.1 Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately 4 percentage points below optimum moisture content.

9.2 Form a specimen by compacting the prepared soil in the 4-in. (101.60-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5 in. (125 mm). Prior to compaction, place the loose soil into the mold and spread into a layer of uniform thickness. Lightly tamp the soil prior to compaction until it is not in a loose or fluffy state, using either the manual compaction rammer or similar device having a face diameter of approximately 2 in. (50 mm). Following compaction of each of the first two layers, any soil adjacent to the mold walls that has not been compacted or extends above the compacted surface shall be trimmed using a knife or other suitable device, and be evenly distributed on top of the layer. Compact each layer by 25 uniformly distributed blows from the rammer dropping free from a height of 12 in. (305 mm) above the elevation of the soil when a sleeve-type rammer is used, or from 12 in. (305 mm) above the approximate elevation of each finely compacted layer when a stationary mounted type rammer is used. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation (Note 7).

9.2.1 Following compaction, remove the extension collar, carefully trim the compacted soil even with the top of the mold by means of the straightedge. Holes developed in the surface by removal of coarse material shall be patched with smaller sized material. Determine the mass of the mold and moist soil in kilograms to the nearest 5 grams, or determine the mass in pounds to the nearest 0.01 pounds. For molds conforming to tolerances given in Section 3.1.1 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 1060, and record the result as the wet density, $W_1$, in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in Section 3.1.1 and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 30, and record the result as the wet density, $W_1$, in pounds per cubic foot, of compacted soil. For used molds out of tolerance by not more than 50 percent (3.1.3), use the factor for the mold as determined in accordance with Section 8 (Calibration of Measure), AASHTO T 19/T 19M.

9.3 Remove the material from the mold and slice vertically through the center. Take a representative sample of the material fro one of the cut faces, determine the mass immediately and dry in accordance with T 255 or T 265, to determine the moisture content, and record the results.

9.4 Thoroughly break up the remainder of the material until it will pass a 3/4 in. (19.0-mm) sieve and 90 percent of the soil aggregations will pass a No. 4 (4.75-mm) sieve as judged by eye, and add to the remaining portion of the sample being tested. Add water in sufficient amounts to increase the moisture content of the soil sample by one or two percentage points, and repeat the above procedure for each increment of water added. Continue this series of determinations until there is either a decrease or no change in the wet mass, $W_1$, per cubic foot (cubic meter) of compacted soil (Note 8).

**METHOD D**

10. **SAMPLE**

10.1 Select the representative sample in accordance with Section 8.3 except that it shall have a mass of approximately 25 lb (11 kg).
11. PROCEDURE

11.1 Follow the same procedure as described for Method C in Section 9, except for the following:
Form a specimen by compacting the prepared soil in the 6-in. (152.4-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5 in. (125 mm), each layer being compacted by 56 uniformly distributed blows from the rammer. For molds conforming to tolerances given in Section 3.1.2, and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 471, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in Section 3.1.2, and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 13.33, and record the result as the wet density, \( W_1 \), in pounds per cubic foot, of the compacted soil. For used molds out of tolerance by not more than 50 percent (Section 3.1.3), use the factor for the mold as determined in accordance with Section 9 (Calibration of Measure), T 19/T 19M.

CALCULATIONS AND REPORT

12. CALCULATIONS

12.1 Calculate the moisture content and the dry unit mass of the soil as compacted for each trial, as follows:

\[
\begin{align*}
\text{w} & = \frac{A-B}{B-C} \times 100 \\
\text{and} \quad \text{W} & = \frac{W_1}{w+100} \times 100
\end{align*}
\]

where:
\( w \) = percentage of moisture in the specimen, based on oven dry mass of soil;
\( A \) = mass of container and wet soil;
\( B \) = mass of container and dry soil;
\( C \) = mass of container;
\( W \) = dry mass, in kilograms per cubic meter of compacted soil, or pounds per cubic foot of compacted soil; and
\( W_1 \) = wet mass, in kilograms per cubic meter of compacted soil, or pounds per cubic foot of compacted soil.

13. MOISTURE-DENSITY RELATIONSHIP

13.1 The calculations in Section 12.1 shall be made to determine the moisture content and corresponding oven-dry unit mass (density) in kilograms per cubic meter or pounds per cubic foot of the compacted samples. The oven-dry densities (unit mass) of the soil shall be plotted as ordinates and the corresponding moisture content as abscissas.
13.2 Optimum Moisture Content — When the densities and corresponding moisture contents for the soil have been determined and plotted as indicated in Section 13.1, it will be found that by connecting the plotted points with a smooth line, a curve is produced. The moisture content corresponding to the peak of the curve shall be termed the “optimum moisture content” of the soil under the above compaction.

13.3 Maximum Density — The oven-dry density in pounds per cubic foot (kilograms per cubic meter) of the soil at optimum moisture content shall be termed “maximum density” under the above compaction.

14. REPORT

14.1 The report shall include the following:

14.1.1 The method used (Method A, B, C, or D).

14.1.2 The optimum moisture content, as a percentage, to the nearest whole number.

14.1.3 The maximum density in pounds per cubic foot to the nearest whole number (kilograms per cubic meter to the nearest 10 kg/m³).

14.1.4 In Methods C and D indicate if the material retained on the 3/4 in. (19.0-mm) sieve was removed or replaced.

14.1.5 Type of face if other than 2 in. (50.8 mm) circular.

15. PRECISION STATEMENT

15.1 Repeatability — (Single operator) – Two results obtained by the same operation on the same sample in the same laboratory using the same apparatus, and on different days should be considered suspect if they differ by more than 10 percent of their mean for optimum moisture content and 2.2 lb/ft³ (35 kg/m³) for maximum density.

15.2 Reproducibility — (Multi-Laboratory) – Two results obtained by different operators in different laboratories should be considered suspect if they differ by more than 15 percent of their mean for optimum moisture and 4.5 lb/ft³ (72 kg/m³) for maximum density.
Tester Qualification Practical Exam Checklist

Moisture-Density Relations of Soils Using a 5.5-lb (2.5-kg) Rammer and a 12-in. (305-mm) Drop
FOP for AASHTO T 99

Participant Name ____________________________ Exam Date ____________

Procedure Element
1. The tester has a copy of the current procedure on hand? □ Yes □ No
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? □ Yes □ No

Sample Preparation
1. If damp, sample dried in air or drying apparatus, not exceeding 140°F (60°C)? □ Yes □ No
2. Sample pulverized and adequate amount sieved over the No. 4 (4.75 mm) sieve? □ Yes □ No
3. Material retained on the sieve discarded? □ Yes □ No
4. Sample passing the sieve has appropriate mass? □ Yes □ No

Procedure
1. Sample mixed with water to approximately 4 percent below expected optimum moisture content? □ Yes □ No
2. Layer of soil placed in mold with collar attached? □ Yes □ No
3. Mold placed on rigid and stable foundation? □ Yes □ No
4. Lightly tamp soil in mold? □ Yes □ No
5. Soil compacted with 25 blows? □ Yes □ No
6. Scrape sides of mold and evenly distributed on top of the layer? □ Yes □ No
7. Soil placed and compacted in three equal layers? □ Yes □ No
8. No more than ½ inch of soil above the top of the bottom portion of the mold? □ Yes □ No
9. Collar removed and soil trimmed to top of mold with straightedge? □ Yes □ No
10. Mass of mold and contents determined to appropriate precision? □ Yes □ No
11. Wet mass of specimen multiplied by appropriate factor to obtain wet density (.03333 lbs/ft³)? □ Yes □ No
12. Soil removed from mold using sample extruder? □ Yes □ No
13. Soil sliced vertically through center? □ Yes □ No
14. Moisture sample removed from one cut face and moist mass determined immediately? □ Yes □ No
**Procedure**

15. Moisture sample mass of at least 100 g?  
   - Yes ☐  No ☐

16. Sample dried and water content determined according to AASHTO T 255 or T 265?  
   - Yes ☐  No ☐

17. Remainder of material from mold broken up to about passing sieve size and added to remainder of original test sample?  
   - Yes ☐  No ☐

18. Water added to increase moisture content in approximately 2 percent increments?  
   - Yes ☐  No ☐

19. Steps 2 through 15 repeated for each increment of water added?  
   - Yes ☐  No ☐

20. If soil is plastic (clay types):
   a. Sample mixed with water varying moisture content by approximately 2 percent, bracketing the optimum moisture content?  
      - Yes ☐  No ☐
   b. Samples placed in covered containers and allowed to stand for at least 12 hours?  
      - Yes ☐  No ☐

21. Process continued until wet density either decreases or stabilizes?  
   - Yes ☐  No ☐

22. Water content and dry density calculated for each sample?  
   - Yes ☐  No ☐

23. Dry density plotted on vertical axis, moisture content plotted on horizontal axis, and points connected with a smooth curve?  
   - Yes ☐  No ☐

24. Water content at peak of curve recorded as optimum water content and recorded to nearest 1 percent?  
   - Yes ☐  No ☐

25. Dry density at optimum water content reported as maximum density, to nearest 1 lb/ft³ (10 kg/m³)?  
   - Yes ☐  No ☐

26. All calculations performed correctly?  
   - Yes ☐  No ☐

First attempt:  Pass ☐  Fail ☐  Second attempt:  Pass ☐  Fail ☐

Signature of Examiner __________________________________________

**Comments:**

_________________________________________________________________

_________________________________________________________________

_________________________________________________________________

_________________________________________________________________

_________________________________________________________________
WSDOT FOP for AASHTO T 106¹

Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)

1. SCOPE

1.1 This test method covers determination of the compressive strength of hydraulic cement mortars, using 2-in. or (50-mm) cube specimens.

*Note 1:* Test Method C 349 provides an alternative procedure for this determination (not to be used for acceptance tests).

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

1.3 Values in SI units shall be obtained by measurement in SI units or by appropriate conversion, using the Rules for Conversion and Rounding given in Standard IEEE/ASTM SI 10, of measurements made in other units.

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 *AASHTO Standards:*

M 152 Flow Table for Use in Tests of Hydraulic Cement  
M 201 Moist Cabinets, Moist Rooms, and Water Storage Tanks Used in the Testing of Hydraulic Cements and Concretes  
R11 Recommended Practice for Indicating Which Places of Figures Are to be Considered Significant in Specified Limiting Values  
T 162 Mechanical Mixing of Hydraulic Cement Pastes and Mortars of Plastic Consistency

2.2 *ASTM Standards:*

C 349 Test Method for Compressive Strength of Hydraulic Cement Mortars (Using Portions of Prisms Broken in Flexure)  
C 670 Practice for Preparing Precision and Bias Statements for Test Methods for Construction Materials  
C 778 Specification for Standard Sand 2  
C 1005 Specification for Weights and Weighing Devices for Use in Physical Testing of Hydraulic Cements  

¹This Test Method is based on AASHTO T 106-02
3. SUMMARY OF TEST METHOD

3.1 The mortar used consists of one part cement and 2.75 parts of sand proportioned by mass. Portland or air-entraining portland cements are mixed at specified water/cement ratios. Water content for other cements is sufficient to obtain a flow of 110 ± 5 in 25 drops of the flow table. Two-inch or (50-mm) test cubes are compacted by tamping in two layers. The cubes are cured 24 hours in the molds and stripped and immersed in lime water until tested.

4. SIGNIFICANCE AND USE

4.1 This test method provides a means of determining the compressive strength of hydraulic cement and other mortars and results may be used to determine compliance with specifications. Further, this test method is referenced by numerous other specifications and test methods. Caution must be exercised in using the results of this test method to predict the strength of concretes.

5. APPARATUS

5.1 Standard Masses and Balances, shall conform to the requirements of ASTM C 1005. The balance device shall be evaluated for precision and bias at a total load of 2000 g.

5.2 Glass Graduates, of suitable capacities (preferably large enough to measure the mixing water in a single operation) to deliver the indicated volume at 20°C. The permissible variation shall be ±2 mL. These graduates shall be subdivided to at least 5 mL, except that the graduation lines may be omitted for the lowest 10 mL for a 250-mL graduate and for the lowest 25 mL of a 500-mL graduate. The main graduation lines shall be circles and shall be numbered. The least graduations shall extend at least one seventh of the way around, and intermediate graduations shall extend at least one fifth of the way around.

5.3 Specimen Molds, for the 2-in. or (50-mm) cube specimens shall be tight fitting. The molds shall have not more than three cube compartments and shall be separable into not more than two parts. The parts of the molds when assembled shall be positively held together. The molds shall be made of hard metal not attacked by the cement mortar. For new molds the Rockwell hardness number of the metal shall be not less than 55 HRB. The sides of the molds shall be sufficiently rigid to prevent spreading or warping. The interior faces of the molds shall be plane surfaces and shall conform to the tolerances of Table 1.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Permissible Variations of Specimen Molds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
<td>2 in. Cube Molds</td>
</tr>
<tr>
<td>Planeness of Sides</td>
<td>New</td>
</tr>
<tr>
<td>Distance Between Opposite Sides</td>
<td>2 in. ± 0.005 in.</td>
</tr>
<tr>
<td>Height of Each Compartment</td>
<td>2 in. ± 0.001 in. to -0.005 in.</td>
</tr>
<tr>
<td>Angle Between Adjacent Faces</td>
<td>90 ± 0.5°</td>
</tr>
</tbody>
</table>

*Measured at points slightly removed from the intersection. Measured separately for each compartment between all the interior faces and the adjacent face and between interior faces and top and bottom planes of the mold.*
5.4 *Mixer, Bowl and Paddle*, an electrically driven mechanical mixer of the type equipped with paddle and mixing bowl, as specified in T 162.

5.5 *Flow Table and Flow Mold*, conforming to the requirements of M 152.

5.6 *Tamper*, a non-absorptive, nonabrasive, non-brittle material such as a rubber compound having a Shore A durometer hardness of 80 ± 10 or seasoned oak wood rendered non-absorptive by immersion for 15 min in paraffin at approximately 392°F or (200°C), shall have a cross section of about \(\frac{1}{2}\) by 1 in. or (13 by 25 mm) and a convenient length of about 5 to 6 in. or (120 to 150 mm). The tamping face shall be flat and at right angles to the length of the tamper.

5.7 *Trowel*, having a steel blade 4 to 6 in. (100 to 150 mm) in length, with straight edges.

5.8 *Moist Cabinet or Room*, conforming to the requirements of Specification M 201.

5.9 *Testing Machine*, either the hydraulic or the screw type, with sufficient opening between the upper bearing surface and the lower bearing surface of the machine to permit the use of verifying apparatus. The load applied to the test specimen shall be indicated with an accuracy of ± 1.0 %. If the load applied by the compression machine is registered on a dial, the dial shall be provided with a graduated scale that can be read to at least the nearest 0.1 % of the full scale load (Note 2). The dial shall be readable within 1 % of the indicated load at any given load level within the loading range. In no case shall the loading range of a dial be considered to include loads below the value that is 100 times the smallest change of load that can be read on the scale. The scale shall be provided with a graduation line equal to zero and so numbered. The dial pointer shall be of sufficient length to reach the graduation marks; the width of the end of the pointer shall not exceed the clear distance between the smallest graduations. Each dial shall be equipped with a zero adjustment that is easily accessible from the outside of the dial case, and with a suitable device that at all times until reset, will indicate to within 1 percent accuracy the maximum load applied to the specimen.

5.9.1 If the testing machine load is indicated in digital form, the numerical display must be large enough to be easily read. The numerical increment must be equal to or less than 0.10 percent of the full scale load of a given loading range. In no case shall the verified loading range include loads less than the minimum numerical increment multiplied by 100. The accuracy of the indicated load must be within 1.0 percent for any value displayed within the verified loading range. Provision must be made for adjusting to indicate true zero at zero load. There shall be provided a maximum load indicator that at all times until reset will indicate within 1 percent system accuracy the maximum load applied to the specimen.

*Note 2*: As close as can be read is considered \(\frac{1}{50}\) in. or (0.5 mm) along the arc described by the end of the pointer. Also, one half of the scale interval is about as close as can reasonably be read when the spacing on the load indicating mechanism is between \(\frac{1}{25}\) in. or (1 mm) and \(\frac{1}{16}\) in. or (1.6 mm). When the spacing is between \(\frac{1}{16}\) in. or (1.6 mm) and \(\frac{7}{6}\) in. or (3.2 mm), one third of the scale interval can be read with reasonable certainty. When the spacing is \(\frac{7}{8}\) in. or (3.2 mm) or more, one fourth of the scale interval can be read with reasonable certainty.
5.9.2 The upper bearing shall be a spherically seated, hardened metal block firmly attached at the center of the upper head of the machine. The center of the sphere shall lie at the center of the surface of the block in contact with the specimen. The block shall be closely held in its spherical seat, but shall be free to tilt in any direction. The diagonal or diameter (Note 3) of the bearing surface shall be only slightly greater than the diagonal of the face of the 2-in. or (50-mm) cube in order to facilitate accurate centering of the specimen. A hardened metal bearing block shall be used beneath the specimen to minimize wear of the lower platen of the machine. The bearing block surfaces intended for contact with the specimen shall have a Rockwell hardness number not less than 60 HRC. These surfaces shall not depart from plane surfaces by more than 0.0005 in. or (0.013 mm) when the blocks are new and shall be maintained within a permissible variation of 0.001 in. or (0.025 mm).

Note 3: A diameter of 3\(\frac{1}{8}\) in. or (79 mm) provided that the lower bearing block has a diameter slightly greater than the diagonal of the face of the 2-in. or (50-mm) cube but not more than 2.9 in. or (74 mm), and is centered with respect to the upper bearing block and held in position by suitable means.

6. MATERIALS

6.1 Graded Standard Sand:

6.1.1 The sand (Note 4) used for making test specimens shall be natural silica sand conforming to the requirements for graded standard sand in ASTM C 778.

Note 4: Segregation of Graded Sand — The graded standard sand should be handled in such a manner as to prevent segregation, since variations in the grading of the sand cause variations in the consistency of the mortar. In emptying bins or sacks, care should be exercised to prevent the formation of mounds of sand or craters in the sand, down the slopes of which the coarser particles will roll. Bins should be of sufficient size to permit these precautions. Devices for drawing the sand from bins by gravity should not be used.

7. TEMPERATURE AND HUMIDITY

7.1 Temperature — The temperature of the air in the vicinity of the mixing slab, the dry materials, molds, base plates, and mixing bowl, shall be maintained between 68 and 81.5°F or (20 and 27.5°C). The temperature of the mixing water, moist closet or moist room, and water in the storage tank shall be set at 73.5 °F or (23 °C) and shall not vary from this temperature by more than ± 3°F or (± 1.7°C).

7.2 Humidity — The relative humidity of the laboratory shall be not less than 50 percent. The moist closet or moist room shall conform to the requirements of M 201.

8. TEST SPECIMENS

8.1 Make two or three specimens from a batch of mortar for each period of test or test age.
9. PREPARATION OF SPECIMEN MOLDS

9.1 Apply a thin coating of release agent to the interior faces of the mold and non-absorptive base plates. Apply oils and greases using an impregnated cloth or other suitable means. Wipe the mold faces and the base plate with a cloth as necessary to remove any excess release agent and to achieve a thin, even coating on the interior surfaces. When using an aerosol lubricant, spray the release agent directly onto the mold faces and base plate from a distance of 6 to 8 in. or (150 to 200 mm) to achieve complete coverage. After spraying, wipe the surface with a cloth as necessary to remove any excess aerosol lubricant. The residue coating should be just sufficient to allow a distinct fingerprint to remain following light finger pressure (Note 5).

9.2 Seal the surfaces where the halves of the mold join by applying a coating of light cup grease such as petrolatum. The amount should be sufficient to extrude slightly when the two halves are tightened together. Remove any excess grease with a cloth.

9.3 After placing the mold on its base plate (and attaching, if clamp-type) carefully remove with a dry cloth any excess oil or grease from the surface of the mold and the base plate to which watertight sealant is to be applied. As a sealant, use paraffin, microcrystalline wax, or a mixture of three parts paraffin to five parts rosin by mass. Liquefy the sealant by heating between 230 and 248°F or (110 and 120°C). Effect a watertight seal by applying the liquefied sealant at the outside contact lines between the mold and its base plate.

Note 5: Because aerosol lubricants evaporate, molds should be checked for a sufficient coating of lubricant immediately prior to use. If an extended period of time has elapsed since treatment, retreatment may be necessary.

Note 6: Watertight Molds — The mixture of paraffin and rosin specified for sealing the joints between molds and base plates may be found difficult to remove when molds are being cleaned. Use of straight paraffin is permissible if a watertight joint is secured, but due to the low strength of paraffin it should be used only when the mold is not held to the base plate by the paraffin alone. A watertight joint may be secured with paraffin alone by slightly warming the mold and base plate before brushing the joint. Molds so treated should be allowed to return to the specified temperature before use.

10. PROCEDURE

Note: For Field fabrication of grout cubes, follow WSDOT Test Method 813.

10.1 Composition of Mortars

10.1.1 The proportions of materials for the standard mortar shall be one part of cement to 2.75 parts of graded standard sand by mass. Use a water-cement ratio of 0.485 for all portland cements and 0.460 for all air-entraining portland cements. The amount of mixing water for other than portland and air-entraining portland cements shall be such as to produce a flow of 110 ± 5 as determined in accordance with Section 10.3 and shall be expressed as mass percent of cement.
10.1.2 The quantities of materials to be mixed at one time in the batch of mortar for making six and nine test specimens shall be as follows in Table 2:

Table 2 —Mixing Proportions for 2-in. Cubes

<table>
<thead>
<tr>
<th>No. of Specimens</th>
<th>6</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cement, g</td>
<td>500</td>
<td>740</td>
</tr>
<tr>
<td>Sand, g</td>
<td>1375</td>
<td>2035</td>
</tr>
<tr>
<td>Water, mL:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Portland (0.485)</td>
<td>242</td>
<td>359</td>
</tr>
<tr>
<td>Air-entraining portland (0.460)</td>
<td>230</td>
<td>340</td>
</tr>
<tr>
<td>Other (to flow of 110 ± 5)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

10.2 Preparation of Mortar:

10.2.1 Mechanically mix in accordance with the procedure given in T 162.

10.3 Determination of Flow:

10.3.1 Carefully wipe the flow-table top clean and dry, and place the flow mold at the center. Place a layer of mortar about 1 in. or (25 mm) in thickness in the mold and tamp 20 times with the tamper. The tamping pressure shall be just sufficient to ensure uniform filling of the mold. Then fill the mold with mortar and tamp as specified for the first layer. Cut off the mortar to a plane surface, flush with the top of the mold, by drawing the straight edge of a trowel (held nearly perpendicular to the mold) with a sawing motion across the top of the mold. Wipe the table top clean and dry, being especially careful to remove any water from around the edge of the flow mold. Lift the mold away from the mortar 1 min after completing the mixing operation. Immediately, drop the table through a height of ½ in. or (13 mm) 25 times in 15 seconds. Using the calipers, determine the flow by measuring the diameters of the mortar along the lines scribed in the table top, adding the four readings. The total of the four readings from the calipers equals the percent increase of the original diameter of the mortar.

10.3.2 For portland and air-entraining portland cements, merely record the flow.

10.3.3 In the case of cements other than portland or air-entraining portland cements, make trial mortars with varying percentages of water until the specified flow is obtained. Make each trial with fresh mortar.

10.4 Molding Test Specimens:

10.4.1 Immediately following completion of the flow test, return the mortar from the flow table to the mixing bowl. Quickly scrape the bowl sides and transfer into the batch the mortar that may have collected on the side of the bowl and then remix the entire batch 15 seconds at medium speed. Upon completion of mixing, the mixing paddle shall be shaken to remove excess mortar into the mixing bowl.
10.4.2 When a duplicate batch is to be made immediately for additional specimens, the flow test may be omitted and the mortar allowed to stand in the mixing bowl 90 seconds without covering. During the last 15 seconds of this interval, quickly scrape the bowl sides and transfer into the batch the mortar that may have collected on the side of the bowl. Then remix for 15 seconds at medium speed.

10.4.3 Start molding the specimens within a total elapsed time of not more than 2 min and 30 seconds after completion of the original mixing of the mortar batch. Place a layer of mortar about 1 in. or (25 mm) (approximately one half of the depth of the mold) in all of the cube compartments. Tamp the mortar in each cube compartment 32 times in about 10 seconds in 4 rounds, each round to be at right angles to the other and consisting of eight adjoining strokes over the surface of the specimen, as illustrated in Figure 1. The tamping pressure shall be just sufficient to ensure uniform filling of the molds. The 4 rounds of tamping (32 strokes) of the mortar shall be completed in one cube before going to the next. When the tamping of the first layer in all of the cube compartments is completed, fill the compartments with the remaining mortar and then tamp as specified for the first layer. During tamping of the second layer bring in the mortar forced out onto the tops of the molds after each round of tamping by means of the gloved fingers and the tamper upon completion of each round and before starting the next round of tamping. On completion of the tamping, the tops of all cubes should extend slightly above the tops of the molds. Bring in the mortar that has been forced out onto the tops of the molds with a trowel and smooth off the cubes by drawing the flat side of the trowel (with the leading edge slightly raised) once across the top of each cube at right angles to the length of the mold. Then, for the purpose of leveling the mortar and making the mortar that protrudes above the top of the mold of more uniform thickness, draw the flat side of the trowel (with the leading edge slightly raised) lightly once along the length of the mold. Cut off the mortar to a plane surface flush with the top of the mold by drawing the straight edge of the trowel (held nearly perpendicular to the mold) with a sawing motion over the length of the mold.
10.5 Storage of Test Specimens — Immediately upon completion of molding, place the test specimens in the moist closet or moist room. Keep all test specimens, immediately after molding, in the molds on the base plates in the moist closet or moist room from 20 to 24 hrs with their upper surfaces exposed to the moist air but protected from dripping water. If the specimens are removed from the molds before 24 hrs, keep them on the shelves of the moist closet or moist room until they are 24 hrs old, and then immerse the specimens, except those for the 24-hr test, in saturated lime water in storage tanks constructed of non-corroding materials. Keep the storage water clean by changing as required.

10.6 Determination of Compressive Strength:

10.6.1 Test the specimens immediately after their removal from the moist closet in the case of 24 hrs specimens, and from storage water in the case of all other specimens. All test specimens for a given test age shall be broken within the permissible tolerance prescribed as follows in Table 3:

<table>
<thead>
<tr>
<th>Test Age</th>
<th>Permissible Tolerance</th>
</tr>
</thead>
<tbody>
<tr>
<td>24 hrs</td>
<td>± ½ hr</td>
</tr>
<tr>
<td>3 days</td>
<td>± 1 hr</td>
</tr>
<tr>
<td>7 days</td>
<td>± 3 hr</td>
</tr>
<tr>
<td>28 days</td>
<td>± 12 hr</td>
</tr>
</tbody>
</table>

If more than one specimen at a time is removed from the moist closet for the 24-hr tests, keep these specimens covered with a damp cloth until time of testing. If more than one specimen at a time is removed from the storage water for testing, keep these specimens in water at a temperature of 73.4±3°F or (23 ± 2°C) and of sufficient depth to completely immerse each specimen until time of testing.

10.6.2 Wipe each specimen to a surface-dry condition, and remove any loose sand grains or incrustations from the faces that will be in contact with the bearing blocks of the testing machine. Check these faces by applying a straightedge (Note 7). If there is appreciable curvature, grind the face or faces to plane surfaces or discard the specimen. A periodic check of the cross-sectional area of the specimens should be made.

*Note 7: Specimen Faces* — Results much lower than the true strength will be obtained by loading faces of the cube specimen that are not truly plane surfaces. Therefore, it is essential that specimen molds be kept scrupulously clean, as otherwise, large irregularities in the surfaces will occur. Instruments for cleaning molds should always be softer than the metal in the molds to prevent wear. In case grinding specimen faces is necessary, it can be accomplished best by rubbing the specimen on a sheet of fine emery paper or cloth glued to a plane surface, using only a moderate pressure. Such grinding is tedious for more than a few thousandths of an inch (hundredths of a millimeter); where more than this is found necessary, it is recommended that the specimen be discarded.
10.6.3 Apply the load to specimen faces that were in contact with the true plane surfaces of the mold. Carefully place the specimen in the testing machine below the center of the upper bearing block. Prior to the testing of each cube, it shall be ascertained that the spherically seated block is free to tilt. Use no cushioning or bedding materials. Bring the spherically seated block into uniform contact with the surface of the specimen. Apply the load rate at a relative rate of movement between the upper and lower platens corresponding to a loading on the specimen with the range of 200 to 400 lbs/s (900 to 1800 N/S). Obtain this designated rate of movement of the platen during the first half of the anticipated maximum load and make no adjustment in the rate of movement of the platen in the latter half of the loading especially while the cube is yielding before failure.

*Note 8:* It is advisable to apply only a very light coating of a good quality, light mineral oil to the spherical seat of the upper platen.

11. **CALCULATION**

11.1 Record the total maximum load indicated by the testing machine, and calculate the compressive strength as follows:

\[ fm = \frac{P}{A} \]  

where:

- \( fm \) = compressive strength in psi or (MPa),
- \( P \) = total maximum load in lbf or (N), and
- \( A \) = area of loaded surface in \( \text{in}^2 \) or (mm\(^2\)).

Either 2-in. or (50-mm) cube specimens may be used for the determination of compressive strength, whether inch-pound or SI units are used. However, consistent units for load and area must be used to calculate strength in the units selected. If the cross-sectional area of a specimen varies more than 1.5 percent from the nominal, use the actual area for the calculation of the compressive strength. The compressive strength of all acceptable test specimens (see Section 12) made from the same sample and tested at the same period shall be averaged and reported to the nearest 10 psi (0.1 MPa).

12. **REPORT**

12.1 Report the flow to the nearest 1 percent and the water used to the nearest 0.1 percent. Average compressive strength of all specimens from the same sample shall be reported to the nearest 10 psi (0.1 MPa).

13. **FAULTY SPECIMENS AND RETESTS**

13.1 In determining the compressive strength, do not consider specimens that are manifestly faulty.

13.2 The maximum permissible range between specimens from the same mortar batch, at the same test age is 8.7 percent of the average when three cubes represent a test age and 7.6 percent when two cubes represent a test age (Note 9).

*Note 9:* The probability of exceeding these ranges is 1 in 100 when the within-batch coefficient of variation is 2.1 percent. The 2.1 percent is an average for laboratories participating in the portland cement and masonry cement reference sample programs of the Cement and Concrete Reference Laboratory.
13.3 If the range of three specimens exceeds the maximum in 13.2, discard the result which differs most from the average and check the range of the remaining two specimens. Make a retest of the sample if less than two specimens remain after discarding faulty specimens or discarding tests that fail to comply with the maximum permissible range of two specimens.

**Note 10:** Reliable strength results depend upon careful observance of all of the specified requirements and procedures. Erratic results at a given test period indicate that some of the requirements and procedures have not been carefully observed; for example, those covering the testing of the specimens as prescribed in 10.6.2 and 10.6.3. Improper centering of specimens resulting in oblique fractures or lateral movement of one of the heads of the testing machine during loading will cause lower strength results.

14. **PRECISION AND BIAS**

See AASHTO T 106 for Precision

15. **KEYWORDS**

15.1 Compressive strength; hydraulic cement mortar; hydraulic cement strength; mortar strength; strength
Performance Exam Checklist

Compressive Strength of Hydraulic Cement Mortar for AASHTO T 106

Participant Name _______________________________ Exam Date __________________

Procedure Element

1. The tester has a copy of the current procedure on hand? Yes No
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? Yes No
3. Cubes are broken within permissible tolerance for time? Yes No
4. Cubes tested immediately after removal from moist closet? Yes No
5. Specimens covered with damp cloth while out of moist room closet? Yes No
6. Cubes wiped clean of sand, and wiped to surface dry condition prior to testing? Yes No
7. Faces to contact the bearing blocks are those that were in contact with the mold? Yes No
8. Faces that will contact the bearing blocks checked with a straightedge? Yes No
9. Cross-sectional area determined in respect to those faces contacting the bearing blocks? Yes No
10. Prior to testing each cube, the spherically seated block was checked for freedom to tilt? Yes No
11. Load rate of 200 to 400 lbf/s (900-1800 N/s) obtained during the first half of the anticipated load? Yes No
12. No adjustment in rate was made during the second half of the loading? Yes No
13. Compressive strength of cubes averaged and reported to the nearest 10 psi (0.1 MPa)? Yes No

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Date: _______________ Tech. ID Code: _______________ Insp. ID Code _______________

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________
AASHTO T 112 (Modified)
Clay Lumps and Friable Particles in Aggregate

AASHTO T 112 has been adopted by WSDOT.

The following definition for “Constant Mass” has been added to this procedure.

“Constant Mass - Test samples dried at a temperature of 110 ± 5C (230 ± 9F) to such a condition that it will not lose more than 0.1 per cent moisture after 2 hours of drying. Such a condition of dryness can be verified by determining the mass of the sample before and after successive 2 hour drying periods. In lieu of such determination, samples may be considered to have reached constant mass when they have been dried at a temperature of 110 ± 5C (230 ± 9 F) for an equal or longer period than that previously found adequate for producing the desired constant mass condition under equal or heavier loading condition of the oven.”
WSDOT Test Method T 113

Method of Test for Determination of Degradation Value

1. SCOPE
   a. This method covers the procedure for determining the susceptibility of an aggregate to degrade into plastic fines when abraded in the presence of water.

2. APPARATUS
   a. Balance — 5000 g capacity, sensitive to 0.1 g
   b. Degradation Shaker — Tyler Portable Sieve Shaker CL-305 modified to provide 300 ± 5 oscillations per minute with a 1 3/4 in. (44.5 mm) throw on the cam or a shaker with equivalent movement
   c. Washing Canister — Shall be either Plastic or Steel meeting the following:
      Plastic Canister — 7 1/2 in. ± 1/4 in (190.5 mm ± 6.3 mm) diameter x 6 ± 1/2 in. (152.4 mm ± 12.5 mm) high. Sidewalls of the plastic canister should meet the bottom at 90 degrees with little or no fillet
      Steel Canister: Meeting the requirements of AASHTO T 210 (ASTM D3744)
   d. Sand equivalent graduated cylinder and rubber stopper
   e. Sand equivalent stock solution
   f. Sieves — 1/2 in. (12.5 mm), 3/8 in. (9.5 mm), 1/4 in. (6.3 mm), U.S. No. 10 (2.00 mm) and U.S. No. 200 (0.075 mm) sieves conforming to the requirement of AASHTO M-92
   g. Graduates — 500 ml tall form, 100 ml
   h. Interval timer
   i. Funnel — Large enough to securely hold the nest of sieves and a mouth that fits into the 500 ml graduate
   j. Sieve Shaker – Shaker that meets the requirements of AASHTO T-27
   k. Oven — Sufficient size, capable of maintaining a uniform temperature of 230 ± 9F (110 ± 5C)
   l. Sprayer – Water sprayer, device to produce a low volume stream of water. i.e. 500 ml wash bottle
   m. Suitable Containers – Pans for washing and drying

3. SAMPLE PREPARATION
   a. If testing pit run material: dry at 230 ± 9F (110 ± 5C) to allow for clean separation from the fine material. Separate the material over the 1/2 in. (12.5 mm) sieve and discard that finer than the 1/2 in. (12.5 mm) and proceed to step 3d.
   b. If testing crushed and stockpiled material: dry at 230 ± 9F (110 ± 5C) to allow for clean separation from the fine material and proceed to step 3e.
   c. If testing quarry material: if necessary, separate the material over the 1/2 in. (12.5 mm) sieve and discard that finer than the 1/2 in. (12.5 mm).
d. Crush the material to be tested to pass the \( \frac{1}{2} \) in. sieve (12.5 mm).

e. Split out an adequate amount of crushed material (approximately 5000 grams).

f. Sieve the approx. 5000 g split over a \( \frac{1}{2} \) in. (12.5 mm), 3/8 in. (9.5 mm), \( \frac{1}{4} \) in. (6.3 mm), and U.S. No. 10 (2.00 mm) screens in a sieve shaker. Steps should be taken to avoid overloading the sieves. Use shaking time determined to meet the requirement of AASHTO T 27 Section 8.2 for the shaker being used.

g. By splitting or quartering, obtain from the sieved material approximately 550 g of \( \frac{1}{2} - 3/8 \) (12.5-9.5 mm), 550 g of 3/8- \( \frac{1}{4} \) (9.5-6.3 mm), and 1100 g of \( \frac{1}{4} - #10 \) (6.3-2.00 mm).

h. Combine the \( \frac{1}{2} - 3/8 \) (12.5-9.5 mm) with the 3/8-\( \frac{1}{4} \) (9.5-6.3 mm).

i. Wash the \( \frac{1}{2} - \frac{1}{4} \) (12.5-6.3 mm) and \( \frac{1}{4} - #10 \) (6.3-2.00 mm) portions separately by placing in a container and adding sufficient water to cover it. Agitate vigorously to ensure complete separation of the material finer than No. 200 (0.075 mm) from coarser particles and bring the fine material into suspension above the coarser material.

**Note 1:** The use of a mechanical aggregate washer is NOT permitted in the washing procedure.

Immediately pour the wash water containing the suspended and dissolved solids over a U.S. No. 10 (2.00 mm) sieve, being careful not to pour out the coarser particles. Add a second charge of water to the portion remaining in the container, agitate, and repeat the operation until the wash water is reasonably clear. Return all material retained on the sieve to the container. Repeat the process for the second portion.

j. Place washed portions into suitable containers and dry to a constant weight at 230 ± 9°F (110 ± 5°C).

k. Allow to cool to room temperature.

l. From the washed and dried material, prepare two - 1000 g test samples as follows:

1. Quarter or split the \( \frac{1}{2} - \frac{1}{4} \) (12.5-6.3 mm) to achieve two 500 ± 1 g portions; hand selection of up to 50 g to attain the 500 ± 1 grams is acceptable.

2. Split the \( \frac{1}{4} - #10 \) (6.3-2.00 mm) to achieve two 500 ± 1 g portions; hand selection of up to 50 g to attain the 500 ± 1 grams is acceptable.

3. Combine each of the \( \frac{1}{2} - \frac{1}{4} \) (12.5-6.3 mm) portions with one of the \( \frac{1}{4} - #10 \) (6.3-2.00 mm) portions to create two - 1000 ± 2 g test samples consisting of \( \frac{1}{2} - #10 \) (12.5-2.00 mm) material.

4. **PROCEDURE**

a. Place one test sample in the washing canister, add 200 ± 5 ml of water, cover tightly and place in degradation shaker.

b. Immediately agitate the material for 20 minutes.

c. At the end of the shaking time, empty the washing canister into nested U.S. No. 10 (2.00 mm) and U.S. No. 200 (0.075 mm) sieves fitted into the funnel placed over a 500 ml graduate to catch all wash water.
Note 2: IMPORTANT! It is critical to the test result that material finer than the U.S. No. 200 (0.075 m) is washed off the larger particles into the 500 ml graduate. This process has to be completed using approximately 300 ml of water such that the total amount water used in the test is only 500 ml. (200 ml with shaking, plus the 20-50 ml used for rinsing the canister and lid, plus that remaining to wash the fines off the particles) The process should be slow and meticulous, utilizing a high pressure, low volume spray of water. Use of a 500 ml squeeze type wash bottle has been found to work well for this process. The washing process should take 5 – 10 minutes.

d. Rinse material finer than U.S. No. 200 (0.075 mm) off the lid into the washing canister and then from the washing canister into the nested sieves using minimal amount of water. (20-50 ml)

e. Shake the nested sieves to spread the sample evenly. (Note 3)

f. Wash the sample using only 20-50 ml. of water. (Note 2)

g. Shake the nested sieves to release any water and 200- that may be sitting on the U.S. No. 200 (0.075 mm) sieve. (Note 3)

h. Raise the funnel and tilt slightly, insure that the mouth of the funnel remains over the 500 ml graduate and catches all of the wash water, to allow the sieves to drain easier. Observe the liquid for clarity.

i. Lower the funnel back into the 500 ml graduate.

j. Repeat steps 4e. through 4i. until the liquid in the graduate reaches the 500 ml mark. Do not allow drainage above the 500 ml mark.

Note 3: Shaking should be vigorous enough to move the aggregate but with care such that no spillage of wash water or loss of aggregate occurs.

k. Measure 7 + 1 ml of sand equivalent stock solution and pour into a sand equivalent cylinder.

l. Bring all solids in the 500 ml graduate into suspension by capping the top with the palm of the hand and turning it completely upside down and back as rapidly as possible, allowing the air bubble to traverse from end to end. Repeat this cycle 10 times, shaking the graduate on the first inversion to release sediment on the bottom.

m. After the tenth cycle, immediately pour the agitated liquid into the sand equivalent cylinder to the 15 ± 0.1 inch. (381 ± 2.5 mm) mark before any settling occurs. (Note 4)

Note 4: The pour should be immediate and continuous without pause. Allowing the agitated liquid to flow back into the 500 ml graduate and then resuming the pour will allow settling and yield inconsistent results.

n. Insert rubber stopper into the sand equivalent cylinder and mix the contents by turning the cylinder completely upside down and back as rapidly as possible, allowing the bubble to traverse from end to end. Repeat this cycle 20 times.

o. Gently place the sand equivalent cylinder on the table, remove stopper, and immediately start timer. Allow to stand undisturbed for 20 minutes. After 20 minutes read and record the height of the sediment column to the nearest 0.1 in. (2.5 mm).

p. Repeat steps 4a. thru 4o. for the second test sample.
5. CALCULATIONS

a. Calculate the degradation factors for the two test samples using the following formula:
\[
D_1 = \frac{15 - H_1}{15 + 1.75H_1} \times 100
\]
\[
D_2 = \frac{15 - H_2}{15 + 1.75H_2} \times 100
\]

Note: Table 1 may be used to determine the values of \(D_1\) and \(D_2\) by finding the corresponding \(H\) value.

b. Average the two degradation factors if they meet the requirements of Section 6, Repeatability:
\[
D = \frac{(D_1 + D_2)}{2}
\]

Where:
- \(D\) = Degradation Factor
- \(D_1\) = Degradation Factor for the first test sample
- \(D_2\) = Degradation Factor for the second test sample
- \(H_1\) = Height of Sediment in first sand equivalent cylinder
- \(H_2\) = Height of Sediment in second sand equivalent cylinder

c. Report the Degradation Factor \((D)\) to the nearest whole number.

d. Degradation Factors range from 0 to 100, with higher values representing the best materials.

6. REPEATABILITY

a. The two test samples, \(D_1\) & \(D_2\) must agree within 6 points.

b. Repeat the entire test if variation between the test samples exceeds 6 points, see following calculation:
\[
\text{Absolute Value}(D_1 - D_2) > 6
\]
### Table 1
Degradation Value “D”

\[
D = \frac{(15-H)}{(15+1.7H)} \times 100
\]

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Method of Test for Determination of Degradation Value  
(WSDOT TM 113)  
Performance Exam Checklist

**Procedure Element**

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<th>Equipment</th>
<th>Yes</th>
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<td>1. Balance - 5000g capacity, sensitive to 0.1g- Calibrated</td>
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<tr>
<td>2. Degradation Shaker – 1 3/4” throw, 300 ± 5 oscillations per minute – Verified</td>
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<td>☐</td>
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<tr>
<td>3. Canister – plastic, 7 1/2 in. diameter x 6 in. high, walls meet floor at 90 deg with min fillet, or steel meeting AASHTO T210, or ASTM D 3744</td>
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<tr>
<td>4. Sand Equivalent Cylinder &amp; Rubber Stopper</td>
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<tr>
<td>5. Sand Equivalent Stock Solution</td>
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<tr>
<td>7. Graduates – 500 ml tall form &amp; 100 ml</td>
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<td>8. Interval Timer – Verified</td>
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<tr>
<td>9. Funnel – Large enough to hold the sieves with a mouth that fits in the 500 ml graduate</td>
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<td>10. Sieve Shaker(s) – Verified</td>
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<td>11. Oven – verified at 230 ± 9 deg. F.- Calibrated</td>
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<tr>
<td>12. Sprayer – produces a low volume stream of water</td>
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<td>☐</td>
</tr>
<tr>
<td>13. Containers – suitable for drying and washing</td>
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</table>

**Procedure:**

1. a.  Pit Run – Dried and separated over the ½ in., ½- discarded | ☐ | ☐ |
   b. Processed material – Dried | ☐ | ☐ |
   c. Quarry material – prepared for crushing | ☐ | ☐ |
2. Material crushed to pass the ½” | ☐ | ☐ |
3. Split out approx. 5000g | ☐ | ☐ |
4. Separate the material over the ½, 3/8, ¼, and No. 10 | ☐ | ☐ |
5. Split or quarter approx. 550g ½-3/8, 550g 3/8-1/4, & 1100g ¼-No. 10 | ☐ | ☐ |
6. Combine the ½-3/8 with the 3/8-1/4 | ☐ | ☐ |
7. Hand wash the ½-1/4 and ¼-No. 10 separately | ☐ | ☐ |
8. Dry the portions in suitable containers at 230 ± 9 to a constant weight | ☐ | ☐ |
9. Split of quarter the two sizes into two 500 ± 1g portions, hand selection ok to 50g | ☐ | ☐ |
10. Combine to create two 1000 ± 2g, ½ - No. 10 test samples | ☐ | ☐ |
11. Place one sample into a canister, cover with 200 ± 5 ml water, cover & shake for 20 min. | ☐ | ☐ |
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<th>Yes</th>
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<tr>
<td>12. Empty canister into the nested No. 10 &amp; No. 200 fitted in the funnel</td>
<td></td>
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<tr>
<td>over the 500 ml grad.</td>
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<tr>
<td>13. Rinse the lid into the canister and then the canister into the nested sieves</td>
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<td></td>
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<tr>
<td>14. Shake the sieves to spread the sample</td>
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</tr>
<tr>
<td>15. Wash using only 20-50 ml.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16. Shake the sieves to release trapped water and then lift observing liquid for clarity</td>
<td></td>
<td></td>
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<tr>
<td>17. Repeat 14-16 until water reaches the 500 ml mark – water not to exceed 500 ml</td>
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<tr>
<td>18. No loss of fines or liquid during the washing process</td>
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<tr>
<td>19. Place 7 ± 1 ml of SE Stock Solution in a SE Graduated Cylinder</td>
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<tr>
<td>20. Turn capped 500 ml upside down &amp; back allowing bubble to traverse 10 cycles</td>
<td></td>
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<tr>
<td>21. Immediately pour into a SE Cylinder to the 15 ± .1 mark – no settling allowed</td>
<td></td>
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<tr>
<td>22. Rubber stopper inserted and SE Cylinder turned upside down &amp; back 20 cycles</td>
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<td>23. Place gently, remove stopper, start timer, allow to sit undisturbed for 20 min.</td>
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<tr>
<td>24. Record height of column to nearest 0.1 in.</td>
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<tr>
<td>25. Repeat for second sample</td>
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<tr>
<td>26. Calculations performed correctly</td>
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<tr>
<td>27. Second sample must be within 6 points</td>
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First attempt: Pass [ ] Fail [ ] Second attempt: Pass [ ] Fail [ ]

Signature of Examiner __________________________________________

Comments:

_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
1. SCOPE
   1.1 This test method covers determination of slump of concrete, both in the laboratory and in the field.

   1.2 The values stated in either inch-pound units or SI units are to be regarded separately as standard. Within the text, the SI units are shown in brackets. The values stated in each system are not exact equivalents; therefore, each system shall be used independently of the other. Combining values from the two systems may result in nonconformance with the standard.

   1.3 The text of the standard reference notes and footnotes provide explanatory material. These notes and footnotes (excluding those in tables and figures) shall not be considered as requirements of the standard.

   1.4 This standard does not purport to address all of the safety problems 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 AASHTO Standards:
      T 141 Sampling Freshly Mixed Concrete

   2.2 ASTM Standards:
      C 172 Practice for Sampling Freshly Mixed Concrete

3. SUMMARY OF TEST METHOD
   3.1 A sample of freshly mixed concrete is placed and compacted by rodding in a mold shaped as the frustum of a cone. The mold is raised, and the concrete allowed to subside. The distance between the original and displaced position of the center of the top surface of the concrete is measured and reported as the slump of the concrete.

4. SIGNIFICANCE AND USE
   4.1 This test method is intended to provide the user with a procedure to determine slump of plastic hydraulic-cement concretes.

   **Note 1:** This test method was originally developed to provide a technique to monitor the consistency of unhardened concrete. Under laboratory conditions, with strict control of all concrete materials, the slump is generally found to increase proportionally with the water content of a given concrete mixture, and thus to be inversely related to concrete strength. Under field conditions, however, such a strength relationship is not clearly and consistently shown. Care should therefore be taken in relating slump results obtained under field conditions to strength.

---

1This FOP is based on AASHTO T 119-99.
4.2 This test method is considered applicable to plastic concrete having coarse aggregate up to 1\(\frac{1}{2}\) in. (37.5 mm) in size. If the coarse aggregate is larger than 1\(\frac{1}{2}\) in. (37.5 mm) in size contact the State Materials Laboratory, the test method is applicable when it is performed on the fraction of concrete passing a 1\(\frac{1}{2}\)-in. (37.5-mm) sieve, with the larger aggregate being removed per FOP for WAOTC TM 2. Contact the Materials Laboratory for directions. in accordance with the section titled “Additional Procedure for Large Maximum Size Aggregate Concrete” in Practice T-141-.

4.3 This test method is not considered applicable to non-plastic and non-cohesive concrete.

Note 2— Concretes having slumps less than 0.5 in. (15 mm.) may not be adequately plastic and concretes having slumps greater than about 9 in. (230 mm) may not be adequately cohesive for this test to have significance. Caution should be exercised in interpreting such results.

5. APPARATUS

5.1 Mold — The test specimen shall be formed in a mold made of metal not readily attacked by the cement paste. The metal shall not be thinner than 0.060 in. (1.5 mm) and if formed by the spinning process, there shall be no point on the mold at which the thickness is less than 0.045 in. (1.15 mm). The mold shall be in the form of the lateral surface of the frustum of a cone with the base 8 in. (200 mm) in diameter, the top 4 in. (100 mm) in diameter, and the height 12 in. (300 mm). Individual diameters and heights shall be within ±\(\frac{1}{8}\) in. (3.2 mm) of the prescribed dimensions. The base and the top shall be open and parallel to each other and at right angles to the axis of the cone. The mold shall be provided with foot pieces and handles similar to those shown in Figure 1. The mold shall be constructed without a seam. The interior of the mold shall be relatively smooth and free from projections. The mold shall be free from projections. A mold which clamps to a nonabsorbent base plate is acceptable instead of the one illustrated provided the clamping arrangement is such that it can be fully released without movement of the mold and the base is large enough to contain all of the slumped concrete in an acceptable test.

5.1.1 Mold with alternative materials

5.1.1.1 Molds other than metal are permitted if the following requirements are met: The mold shall meet the shape, height, and internal dimensional requirements of Section 5.1. The mold shall be sufficiently rigid to maintain the specified dimensions and tolerances during use, resistant to impact forces, and shall be nonabsorbent. The mold shall be demonstrated to provide test results comparable to those obtained when using a metal mold meeting the requirements of 5.1. Comparability shall be demonstrated on behalf of the manufacturer by an independent testing laboratory. Test for comparability shall consist of not less than 10 pairs of comparisons performed at each of 3 different slumps ranging from 2 in. (50 mm) to 6 in. (150 mm). No individual test results shall vary by more than 0.50 in. (15 mm) from that obtained using the metal mold. The average test results of each slump range obtained using the mold constructed of alternative material shall not vary by more than 0.30 in. (10 mm) from the average of test results obtained using the metal mold. Manufacturer comparability test data shall be available to users and laboratory inspection authorities (see Note 3). If any changes in material or method of manufacture are made, tests for comparability shall be repeated.
**Note 3**—Because the slump of concrete decreases with time and higher temperatures, it will be advantageous for the comparability tests to be performed by alternating the use of metal cones and alternative material cones, to utilize several technicians, and to minimize the time between test procedures.

5.1.1.2 If the condition of any individual mold is suspected of being out of tolerance from the as manufactured condition, a single comparative test shall be performed. If the test results differ by more than 0.50 in. (15 mm) from that obtained using the metal mold, the mold shall be removed from service.

5.2 Tamping Rod — The tamping rod shall be a round, straight steel rod \(\frac{5}{8}\) in. (16 mm) in diameter and approximately 24 in. (600 mm) in length, having the tamping end or both ends rounded to a hemispherical tip, the diameter of which is \(\frac{5}{8}\) in. (16 mm).

5.3 Torpedo level

5.4 Base — Flat, nonabsorbent, rigid surface.

---

**Figure 1**

**Mold for Slump Test**

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6. SAMPLE

6.1 The sample of concrete from which test specimens are made shall be representative of the entire batch. It shall be obtained in accordance with FOP for WAQTC TM 2. With concrete using 1½ in. (37.5 mm), or larger aggregate, the aggregate larger than 1½ in. (37.5 mm) must be removed per FOP for WAQTC TM 2. Contact the Materials Laboratory for directions.

7. PROCEDURE

7.1 Dampen the mold and place it on a firm, flat, nonabsorbent, level surface. It shall be held firmly in place during filling by the operator standing on the two foot pieces. From the sample of concrete obtained in accordance with Section 6, immediately fill the mold in three layers, each approximately one third the volume of the mold.

**Note 3:** One third of the volume of the slump mold fills it to a depth of $2\frac{5}{8}$ in. (67 mm); two thirds of the volume fills it to a depth of $6\frac{1}{8}$ in. (155 mm).

7.2 Rod each layer with 25 strokes of the tamping rod. Uniformly distribute the strokes over the cross section of each layer. For the bottom layer this will necessitate inclining the rod slightly and making approximately half of the strokes near the perimeter, and then progressing with vertical strokes spirally toward the center. Rod the bottom layer throughout its depth. Rod the second layer and the top layer each throughout its depth, so that the strokes just penetrate into the underlying layer.

7.3 In filling and rodding the top layer, heap the concrete above the mold before rodding is started. If the rodding operation results in subsidence of the concrete below the top edge of the mold, add additional concrete to keep an excess of concrete above the top of the mold at all times. After the top layer has been rodded, strike off the surface of the concrete by means of a screeing and rolling motion of the tamping rod. Continue to hold the mold down firmly and remove concrete from the area surrounding the base of the mold to preclude interface with the movement of slumping concrete.

Remove the mold immediately from the concrete by raising it carefully in a vertical direction. Raise the mold a distance of approximately 12 in. (300 mm) in $5 \pm 2$ seconds by a steady upward lift with no lateral or torsional motion. Complete the entire test from the start of the filling through removal of the mold without interruption and complete it within an elapsed time of $2\frac{1}{2}$ min.

7.4 Immediately measure the slump by determining the vertical difference between the top of the mold and the displaced original center of the top surface of the specimen. If a decided falling away or shearing off of concrete from one side or portion of the mass occurs (Note 4), disregard the test and make a new test on another portion of the sample.

**Note 4:** If two consecutive tests on a sample of concrete show a falling away or shearing off of a portion of the concrete from the mass of the specimen, the concrete probably lacks necessary plasticity and cohesiveness for the slump test to be applicable. Report material cannot be slumped due to shearing or falling away.
8. REPORT

8.1 Report the slump in terms of inches (millimeters) to the nearest 1/4 in. (5 mm) of subsidence of the specimen during the test as follows:

- Slump = 12 inches of height after subsidence
- Slump = 300 mm of height after subsidence

Report results on concrete delivery ticket (i.e., Certificate of Compliance). The signature of the tester who performed the field acceptance test is required on concrete delivery tickets containing test results.

9. PRECISION AND BIAS

9.1 Precision:

See AASHTO T 119 for Precision and bias
## Performance Exam Checklist

### Slump of Hydraulic Cement Concrete

**FOP for AASHTO T 119**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
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**Procedure Element**

<table>
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<tr>
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<th>No</th>
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</table>

1. The tester has a copy of the current procedure on hand?  
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?  
3. Cone and floor or base plate dampened?  
4. Cone held firmly against the base by standing on the two foot pieces? Cone not allowed to move in any way during filling?  
5. Representative samples scooped into the cone?  
6. Cone filled in three approximately equal layers by volume?  
7. Each layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes?  
8. Middle and top layers rodded to just penetrate into the underlying layer?  
9. When rodding the top layer, excess concrete kept above the mold at all times?  
10. Concrete struck off level with top of cone using tamping rod?  
11. Excess concrete removed from around the base?  
12. Cone lifted upward approximately 12 in. (300 mm) in one smooth motion, without twisting the cone, in 5 ± 2 seconds?  
13. Slump measured to the nearest 1/4 in. (5 mm) from the top of the cone to the displaced original center of the top surface of the specimen?  
14. Test performed from start to finish within 2 1/2 minutes?

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner ________________________________________

Comments:

________________________________________________________________
________________________________________________________________
________________________________________________________________

________________________________________________________________
WSDOT Test Method T 123

Method of Test for Bark Mulch

1. SCOPE
   a. This method covers a procedure for determining the sieve analysis and material finer than \(\frac{1}{4}\) in. using a loose volume bucket.

2. EQUIPMENT
   a. A mechanical sieve shaker.
   b. Sieves — A 1 1/2 in. and No. 4 sieves conforming to the requirements of AASHTO M-92. Breaker sieves may be used.
   c. Volume Bucket — A container calibrated in 1 gal. increments from 1 to 5 gal. A 5-gal. bucket may be used when calibrated as follows:
      On a level surface calibrate the container by gradually filling it with water in 1 gal. increments. Mark the inner wall of the container after the addition of each gallon.

3. PROCEDURE
   a. Air dry (140°F max.) the sample for 15 hours, ± 4 hours.
   b. Reduce the sample to testing size per the FOP for AASHTO T 248.
   c. Place the sample in the volume bucket and record the volume as the total volume.
   d. Shake the sample over the 1 1/2 in. and No. 4 sieves. Using breaker sieves inserted between the two specified sieves so the No. 4 sieve will not be overloaded. Use caution to avoid over sieving as the wood material breaks down.
   e. The material retained on the 1 1/2 in. sieve is measured in the volume bucket and recorded.
   f. The material on the breaker sieves is added to the material retained on the No. 4 sieve and the volume measured in the volume bucket and recorded.
   g. The percent passing is calculated as follows:

\[
100 - \frac{(\text{Volume on sieve} \times 100)}{\text{Total Volume}} = \% \text{ passing}
\]
Method of Test for Bark Mulch  
WSDOT T 123  
Performance Exam Checklist

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure,</td>
<td>☐</td>
<td>☐</td>
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<tr>
<td>and if required, has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Bark mulch sample dried for 15 ± 4 hrs @ 140°F?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Five (5) gallon bucket calibrated in 1 gal. increments?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Sample quartered or split and placed in calibrated bucket?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Volume of sample in bucket recorded as total volume?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Sample screened in the shaker through 1½ in. screen, breaker screens and No. 4</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>screen?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Do not over shake to prevent degrading of sample?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. Remove 1½ in. screen and damp material in calibrated bucket and record volume</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>as volume on 1½ in. screen?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. Place all breaker screen material down to No. 4 screen in bucket and record</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>volume as volume on No. 4 screen?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>11. All calculations performed correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>12. Report results?</td>
<td>☐</td>
<td>☐</td>
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</tbody>
</table>

First attempt:  Pass ☐  Fail ☐  Second attempt:  Pass ☐  Fail ☐

Signature of Examiner __________________________________________

Comments:

________________________________________________________________
________________________________________________________________
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WSDOT Test Method T 124

Method for Testing Top Soils

1. SCOPE
   a. This method describes a Modified AASHTO T 88 procedure for the quantitative determination of the distribution of particle sizes in Top Soils.
   b. This method also describes the preparation procedure for the testing of the materials for PH and Organics.

2. EQUIPMENT
   a. Oven — An oven of appropriate size, capable of maintaining a uniform temperature of approximately 140°F (60°C).
   b. Balance — The balance or scale shall be sensitive to 0.1g.
   c. Sieves — A series of sieves conforming to the requirements of the Specifications for Sieves for Testing Purposes.
   d. Stirring Apparatus — A mechanically operated stirring apparatus and dispersion cup the same as described in AASHTO T 88.
   e. Hydrometer — As described in AASHTO T 88.
   f. Sedimentation Cylinders — As described in AASHTO T 88.
   g. Thermometer — As described in AASHTO T 88.
   h. Plunger — A plunger capable of reaching the bottom of the sedimentation cylinder.
   i. Dispersing Agent — 9 g of sodium hexametaphosphate made up to volume, with distilled water in a 1 L volumetric flask.

3. PROCEDURE
   Soil sample as received shall be air dried (140°F (60°C) for 15 hours, ± 4 hours). Run material on coarse screens 1 in. (25 mm) and No. 4 (4.75 mm) for 10 minutes. Record percentage of material retained on the 1 in. (25 mm) and No. 4 (4.75 mm) sieves. Split out approximately 3 lbs. (1500 g.) of No. 4 (4.75 mm) minus material for fine grading.
   Split No. 4 (4.75 mm) minus material to approximately 500 g. Shake by hand on the #10 (2.00 mm) screen and mull if needed for complete separation. Record percent passing. Hand shake approximately 150 g of #10 (2.00 mm) minus material for PH testing and approximately 50 g of #100 (0.150 mm) minus material for organic content testing. These two samples are to be tested by the Chemical Lab.
   Dispersion: Weigh out 40 g (± 0.5 g) of #10 (2.00 mm) minus, air dried soil and transfer to a 250 ml beaker. Add 190 ml of sodium hexametaphosphate solution. Let the soil-water mixture stand for at least 18 hours, but do not permit it to dry out. Using distilled water, wash soil into the dispersion cup, and add water to within 2 in. of the rim of the cup. Disperse the sample with the stirring apparatus for a period of 10 minutes.
Hydrometer Measurements: Wash the contents of the dispersion cup into a 1000 ml graduated cylinder. Place the hydrometer in the cylinder, and then add the distilled water to the 920 ml mark. Use a plunger with vigorous up and down motion to obtain a uniform suspension.

In the mixing process do not bring the perforated base of the plunger too close to the top of the water column or force it down through the column too vigorously or splashing will result and air may be introduced into the suspension. Incorporated air bubbles may cause sizeable positive errors in hydrometer readings. When the suspension is uniform, remove the plunger rapidly but carefully and start counting time as the base of the plunger clears the water surface. Insert the hydrometer quickly, but carefully, and limit bobbing with the finger tip. Record the hydrometer reading (top of the meniscus) after exactly 40 seconds. Move the hydrometer up and down in the suspension to displace soil particles which have settled on it and then remove it from the cylinder. Using the plunger, resuspend the soil, and obtain a second reading. If there is agreement within 0.5 of a scale unit, proceed to the final soil suspension; otherwise repeat the reading.

When the 40-second reading is made, sand will have settled out sufficiently so that only suspended silt and clay particles remain to influence the hydrometer level.

After two satisfactory 40-second readings have been obtained, resuspend the soil for the two hour reading which starts immediately after the plunger is removed.

About 1 minute prior to the two hour reading insert the hydrometer. Since the hydrometer was calibrated at a temperature of 68°F (20°C), data obtained at other temperatures must be corrected. Less error is introduced if the temperature is above than if it is below 68°F (20°C). Avoid extreme temperatures. Correct the observed hydrometer reading by adding a 0.2 unit for each 1° above 68°F (20°C), or subtracting a 0.2 unit for each 1° below 68°F (20°C). At each reading temperature will be taken.
Particle Size Analysis of Top Soils  
(WSDOT TM 124)  
Performance Exam Checklist

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Sample air dried @ 140 F (60 C) maximum?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Test sample obtained by AASHTO T-248?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Representative sample used to fill 1/2 cubic foot bucket?</td>
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</table>

Coarse Sieve Analysis

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Coarse material separated on 1 inch and 1/4 inch sieve in shaker for 10 minutes?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Coarse material separated on 1 inch and 1/4 inch sieve in shaker for 10 minutes?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. All material retained on the 1/4 inch sieve is placed in 1/10 cubic foot bucket?</td>
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<tr>
<td>4. If material fills 1/10 cubic foot bucket, sample is out of specification?</td>
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<tr>
<td>5. If not, percent filled, of 1/10 cubic foot bucket is recorded?</td>
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<tr>
<td>6. Approximately 3 pounds of 1/4 inch minus material is split out as a sample for fine grading?</td>
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<td></td>
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<tr>
<td>7. Split out 500 g sample for actual grading?</td>
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<td></td>
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<tr>
<td>8. Mull material if necessary?</td>
<td></td>
<td></td>
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<tr>
<td>9. Sieve analysis performed on material retained on the No. 10 sieve and percent passing recorded?</td>
<td></td>
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</table>

Samples Prepared for Testing by Others

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
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</thead>
<tbody>
<tr>
<td>1. Approximately 150 g of No. 10 minus material for PH testing?</td>
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<tr>
<td>2. Approximately 50 g of No. 100 minus material for Organic Content Testing?</td>
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<td></td>
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<tr>
<td>3. All material mulled if necessary?</td>
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</tbody>
</table>

Hydrometer Analysis

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Distilled or demineralized water used throughout the test?</td>
<td></td>
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<tr>
<td>2. Air dried sample weighs 40 ± 0.5 grams?</td>
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<tr>
<td>3. Sample placed in 250 ml beaker and 190 ml of Dispersing Agent added?</td>
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<tr>
<td>4. Sample soaked at least 18 hours in dispersing agent? (Do not allow to dry out)</td>
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<tr>
<td>5. Sample washed into dispersion cup with distilled water to within 2 inches of the rim of the cup?</td>
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<tr>
<td>6. Dispersed for 10 minutes?</td>
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<tr>
<td>7. Mixture transferred to 1000 ml glass graduate?</td>
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<tr>
<td>8. Hydrometer placed in the glass graduate and distilled water added to the 920 ml mark?</td>
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<tr>
<td>9. Hydrometer is removed and a plunger is used to mix to uniform suspension?</td>
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<tr>
<td>10. Plunger is removed and timer started?</td>
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</table>
Procedure Element

11. Hydrometer immediately placed in the graduate and reading @ 40 second is recorded? □ □
12. Hydrometer gently moved up and down then removed from graduate? □ □
13. Steps 9 and 10 are repeated for a second, 40 second reading? □ □
14. If the two readings do not agree within 0.5 ml, then steps 9 and 10 are repeated until they do? □ □
15. Plunger used to remix the suspension and timer started for 2 hour reading? □ □
16. Hydrometer placed into graduate one minute prior to 2 hour reading? □ □
17. Temperature is taken at each reading and corrected to 68 F? □ □

Calculations
1. Calculations performed in accordance with test method? □ □

First attempt: Pass □ Fail □
Second attempt: Pass □ Fail □

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________
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AIR CONTENT OF FRESHLY MIXED CONCRETE BY THE PRESSURE METHOD

Significance

Concrete is not a solid, but rather a solid with void spaces. The voids may contain gas such as air, or liquid, such as water. All concrete contains air voids, and the amount can be increased by the addition of an air entraining agent to the mix. When such an agent is used, the size of the voids drastically decreases and the number of voids greatly increases, providing a much greater dispersal of voids.

Air entrainment is necessary in concrete that will be saturated and exposed to cycles of freezing and thawing, and to deicing chemicals. The microscopic entrained air voids provide a site for relief of internal pressure that develops as water freezes and thaws inside the concrete. Without the proper entrained-air content, normal concrete that is saturated and is exposed to cycles of freezing and thawing can fail prematurely by scaling, spalling, or cracking.

Care must be taken, however, not to have too much entrained air. As the air content increases, there will be a corresponding reduction in the strength and other desirable properties of the concrete. Typically, this strength reduction will be on the order of 3 to 5 percent for each 1 percent of air content. A concrete mix design proportioned for 5 percent air, for example, will be approximately 15 to 25 percent lower in strength if the air content were to double.

Scope

This procedure covers determination of the air content in freshly mixed portland cement concrete containing dense aggregates in accordance with AASHTO T 152 (Type B meter). It is not for use with lightweight or highly porous aggregates. This procedure includes calibration of the "Type B" air meter gauge, and two methods for calibrating the gauge are presented. Concrete containing aggregate that would be retained on the 1 1/2 in (37.5 mm) sieve must be wet sieved. Sieve a sufficient amount of the sample over the 1 1/2 in (37.5 mm) sieve in accordance with per the FOP for WAQTC TM2.

Apparatus

- Air meter: Type B, as described in AASHTO T 152
- Balance or scale: Accurate to 0.3 percent of the test load at any point within the range of use (for Method 1 calibration only)
- Verified external or internal calibration vessel of known volume (usually 5% +/- of the volume of the meter base).
- Tamping rod: 5/8 in. (16 mm) diameter and approximately 24 in. (600 mm) long, having a hemispherical tip. (Hemispherical means half a sphere; the tip is rounded like half of a ball.)
- Vibrator: 7000 vibrations per minute, 0.75 to 1.50 in. (19 to 38 mm) in diameter, at least 3 in. (75 mm) longer than the section being vibrated for use with low slump concrete
- Scoop
- Container for water: rubber syringe (may also be a squeeze bottle)
- Strike-off bar: Approximately 12 in. x 3/4 in. x 1/8 in. (300 mm x 22 mm x 3 mm).

This Test Method is based on AASHTO T 152-05.
• Strike-off Plate: A flat rectangular metal plate at least 1/4 in. (6 mm) thick or a glass or acrylic plate at least 1/2 in. (12 mm) thick, with a length and width at least 2 in. (50 mm) greater than the diameter of the measure with which it is to be used. The edges of the plate shall be straight and smooth within tolerance of 1/16 in. (1.5 mm).

Note 1: Use either the strike-off bar or strike-off plate; both are not required.

• Mallet: With a rubber or rawhide head having a mass of 1.25 ±0.5 lb (0.57 ±0.23 kg)

Calibration of Air Meter Gauge

Note 2: There are two methods for calibrating the air meter, mass or volume.

1. Screw the short piece of straight tubing into the threaded petcock hole on the underside of the cover. Determine the mass of the dry, empty air meter base and cover assembly (Mass Method only).
2. Fill the base nearly full with water.
3. Clamp the cover on the base with the tube extending down into the water. Mark the petcock with the tube attached for future reference.
4. Add water through the petcock having the pipe extension below until all air is forced out the other petcock. Rock the meter slightly until all air is expelled through the petcock.
5. Wipe off the air meter base and cover assembly, and determine the mass of the filled unit (Mass Method only).
6. Pump up the air pressure to a little beyond the predetermined initial pressure indicated on the gauge. Wait a few seconds for the compressed air to cool, and then stabilize the gauge hand at the proper initial pressure by pumping up or relieving pressure, as needed.
7. Close both petcocks and immediately open the main air valve exhausting air into the base. Wait a few seconds until the meter needle stabilizes. The gauge should now read 0 percent. If two or more tests show a consistent variation from 0 percent in the result, change the initial pressure line to compensate for the variation, and use the newly established initial pressure line for subsequent tests.
8. Determine which petcock has the straight tube attached to it. Attach the curved tube to external portion of the same petcock.
9. Pump air into the air chamber. Open the petcock with the curved tube attached to it. Open the main air valve for short periods of time until 5 percent of water by mass or volume has been removed from the air meter. Remember to open both petcocks to release the pressure in the base and drain the water in the curved tube back into the base. To determine the mass of the water to be removed, subtract the mass found in Step 1 from the mass found in Step 5. Multiply this value by 0.05. This is the mass of the water that must be removed. To remove 5 percent by volume, remove water until the external calibrating vessel is level full.

Note 3: Many air meters are supplied with a calibration vessel(s) of known volume that are used for this purpose. Calibration vessels should be brass, not plastic, and must be protected from damage that would change their volume, crushing or denting.

If an external or internal calibration vessel is used, confirm what percentage volume it represents for the air meter being used. Vessels commonly represent 5 percent volume, but they are for specific size meters. This should be confirmed by mass.
10. Remove the curved tube. Pump up the air pressure to a little beyond the predetermined initial pressure indicated on the gauge. Wait a few seconds for the compressed air to cool, and then stabilize the gauge hand at the proper initial pressure by pumping up or relieving pressure, as needed.

11. Close both petcocks and immediately open the main air valve exhausting air into the base. Wait a few seconds until the meter needle is stabilized. The gauge should now read 5.0 ± 0.1 – 5.0 ± 0.2 percent. If the gauge is outside that range, the meter needs adjustment. (Consult the Region Materials Lab) The adjustment could involve adjusting the starting point so that the gauge reads 5.0 ± 0.1 – 5.0 ± 0.2 percent when this calibration is run, or could involve moving the gauge needle to read 5.0 percent. Any adjustment should comply with the manufacturer’s recommendations.

*Note 4:* Calibration shall be performed per agency standards, prior to field use, and weekly during construction use, at the frequency required by the agency. Record the date of the calibration, the calibration results, and the name of the technician performing the calibration in the log book kept with each air meter.

12. When the gauge hand reads correctly at 5.0 percent, additional water may be withdrawn in the same manner to check the results at other values such as 10 percent or 15 percent.

*Note 5:* Remove the extension tubing from threaded petcock hole in the underside of the cover before starting the test procedure.

An internal calibration vessel of known volume, usually 5% of the volume of the bucket, may be employed as a quick method to verify the calibration of the air meter during construction use. To employ this vessel proceed as follows.

13. Fill the base nearly full with water and place the internal calibration vessel into the base. Place the cover back on the base and gently add water through the petcock until all the air has been expelled. Do not disturb the meter to such an extent as to knock the calibration vessel from an upright position. Do not install either of the threaded tubes into the petcock when using the calibration vessels.

14. Pump up the air pressure to a little beyond the predetermined initial pressure indicated in the calibration record log book. Wait a few seconds for the compressed air to cool and then stabilize the gauge hand at the proper initial pressure by pumping up or relieving pressure, as needed.

15. Close both petcocks and immediately open the main air valve exhausting air into the base. Wait a few seconds and gently tap the back of the gauge until the meter needle stabilizes. The gauge should now read 5.0 +/- 0.2 percent. If the gauge is outside of that range follow step 1 through step 12 of the calibration procedure to re-calibrate the air meter. If further adjustment is required consult the Region Materials Lab.

16. If necessary, additional vessels may be placed into the base to verify the calibration of the air meter at 10% volume and 15% volume or the sum of the volumes indicated on the individual calibration vessels.

17. Record the date that the calibration of the air meter was verified in the calibration log book.

18. Gently release the air pressure in the base by opening one of the petcocks then remove and drain any water from within the calibration vessel and store it in a safe location. The air meter is now ready for use.
12. When the gauge hand reads correctly at 5.0 percent, additional water may be withdrawn in the same manner to check the results at other values such as 10 percent or 15 percent.

13. If an internal calibration vessel is used follow steps 1 thru 8 to set initial reading.

14. Release pressure from the base and remove cover. Place the internal calibration vessel into the base. This will displace 5 percent of the water in the base. (see AASHTO-T152 for more information on internal calibration vessels)

15. Place the cover back on the base and add water through the petcock until all the air has been expelled.

16. Pump up the air pressure chamber to the initial pressure. Wait a few seconds for the compressed air to cool, and then stabilize the gauge hand at the proper initial pressure by pumping up or relieving pressure, as needed.

17. Close both petcocks and immediately open the main air valve exhausting air into the base. Wait a few seconds until the meter needle stabilizes. The gauge should now read 5 percent.

Procedure Selection

There are two methods of consolidating the concrete – rodving and vibration. If the slump is greater than 3 in. (75 mm), consolidation is by rodving. When the slump is 1 to 3 in. (25 to 75 mm), internal vibration or rodving can be used to consolidate the sample, but the method used must be that required by the agency in order to obtain consistent, comparable results. For slumps less than 1 in. (25 mm), consolidate the sample by internal vibration.

PROCEDURE – RODDING

1. Obtain the sample in accordance with the FOP for WAQTC TM 2. If any aggregate 37.5 mm (1½ in.) or larger than 1½ in. (37.5 mm) is present, the larger aggregate must be removed. Sieve a sufficient amount of the sample over the 1½ in. (37.5 mm), sieve in accordance with the Wet Sieving portion of the FOP for WAQTC TM 2. Contact the Materials Laboratory for directions.

   Note 7: Testing shall begin within five minutes of obtaining the sample.

2. Dampen the inside of the air meter base and place on a firm, level surface.

3. Fill the base approximately 1/3 full with concrete.

4. Consolidate the layer with 25 strokes of the tamping rod, using the rounded end. Distribute the strokes evenly over the entire cross section of the concrete. Rod throughout its depth without hitting the bottom too hard.

5. Tap the sides of the base smartly 10 to 15 times with the mallet to close voids and release trapped air.

6. Add the second layer, filling the base about 2/3 full.

7. Consolidate this layer with 25 strokes of the tamping rod, penetrating about 1 in. (25 mm) into the bottom layer.

8. Tap the sides of the base 10 to 15 times with the mallet.

9. Add the final layer, slightly overfilling the base.

10. Consolidate this layer with 25 strokes of the tamping rod, penetrating about 1 in. (25 mm) into the second layer.
11. Tap the sides of the base smartly 10 to 15 times with the mallet.  

*Note 8:* The base should be slightly over full, about 1/8 in. (3 mm) above the rim. If there is a great excess of concrete, remove a portion with the trowel or scoop. If the base is under full, add a small quantity. This adjustment may be done only after consolidating the final layer and before striking off the surface of the concrete.

12. Strike off the surface of the concrete and finish it smoothly with a sawing action of the strike-off bar or plate, using great care to leave the base just full. The surface should be smooth and free of voids, as much as possible.

13. Clean the top flange of the base to ensure a proper seal.

14. Moisten the inside of the cover and check to see that both petcocks are open and the main air valve is closed.

15. Clamp the cover on the base.

16. Inject water into one petcock until water emerges from the second petcock. (Note: Water is injected into only one petcock during the entire procedure)

17. Jar or rock the air meter gently until no air bubbles appear to be coming out of the second petcock. The petcock expelling water should be higher than the petcock where water is being injected. Return the air meter to a level position and verify that water is present in both petcocks.

18. Close the air bleeder valve and pump air into the air chamber until the needle goes past the initial pressure line. Allow a few seconds for the compressed air to cool.

19. Tap the gauge gently with one hand while slowly opening the air bleeder valve until the needle rests on the initial pressure line. Close the air bleeder valve.

20. Close both petcocks.

21. Open the main air chamber valve.

22. Tap the sides of the base smartly with the mallet.

23. With the main air chamber valve open, lightly tap the gauge to settle the needle, and then read the air content to the nearest 0.1 percent, while the air chamber value is open.

24. Release or close the main air chamber valve.

25. Open both petcocks to release pressure, remove the concrete, and thoroughly clean the cover and base with clean water.

26. Open the main air valve to relieve the pressure in the air chamber.

**Procedure - Internal Vibration**

1. Obtain the sample in accordance with the FOP for WAQTC TM 2. If any aggregate 37.5mm (1 1/2 in.) or larger than 1 1/2 in (37.5 mm) is present, the larger aggregate must be removed. Sieve a sufficient amount of the sample over the 1 1/2 in (37.5 mm). sieve in accordance with the Wet Sieving portion of the FOP for WAQTC TM 2. Contact the Materials Laboratory for directions.

2. Dampen the inside of the air meter bowl and place on a firm level surface.

3. Fill the base approximately half full.
4. Insert the vibrator at three different points. Do not let the vibrator touch the bottom or sides of the base.

   **Note 9:** Remove the vibrator slowly, so that no air pockets are left in the material.

   **Note 10:** Continue vibration only long enough to achieve proper consolidation of the concrete. Over vibration may cause segregation and loss of appreciable quantities of intentionally entrained air.

5. Fill the base a bit over full.

6. Insert the vibrator as in Step 3. Do not let the vibrator touch the sides of the base, and penetrate the first layer approximately 1 in. (25 mm).

7. Return to Step 12 of the rodding procedure and continue.

**Report**

Results shall be reported on standard forms approved for use by the agency. Record the percent of air to the nearest 0.1 percent.

Report results on concrete delivery ticket, (i.e. Certificate of Compliance).

The signature of the tester who performed the field acceptance test is required on concrete delivery tickets containing test results.

**Note 11:** Some agencies require an aggregate correction factor in order to determine total percent air.

**Aggregate Correction Factor**

When available use the aggregate correction factor from the mix design in order to determine total percent entrained air.

Total percent entrained = gauge reading – aggregate correction factor from the mix design.

See AASHTO T 152.
Performance Exam Checklist

AIR CONTENT OF FRESHLY MIXED CONCRETE BY THE PRESSURE METHOD

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
</table>

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<thead>
<tr>
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</tr>
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<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Container filled in three equal layers, slightly overfilling the last layer?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Each layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Bottom layer rodded throughout its depth, without forcibly striking the bottom of the container?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Middle and top layers rodded, each throughout their depths and penetrating 1 in. (25 mm) into the underlying layer?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Sides of the container tapped 10 to 15 times with the mallet after rodding each layer?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Concrete struck off level with top of container using the bar and rim cleaned off?</td>
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<td></td>
</tr>
</tbody>
</table>

Using a Type B Meter

<table>
<thead>
<tr>
<th>Procedure Element</th>
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<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>9. Both petcocks open?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Air valve closed between air chamber and the bowl?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Inside of cover cleaned and moistened before clamping to base?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12. Water injected through petcock until it flows out the other petcock?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. Water injection into the petcock continued while jarring and tapping the meter to insure all air is expelled?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14. Air pumped up to initial pressure line?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15. A few seconds allowed for the compressed air to stabilize?</td>
<td></td>
<td></td>
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<tr>
<td>16. Gauge adjusted to the initial pressure?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17. Both petcocks closed?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18. Air valve opened between chamber and bowl?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>19. Sides of bowl tapped with the mallet?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Procedure Element</td>
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<td>No</td>
</tr>
<tr>
<td>-------------------</td>
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<td>----</td>
</tr>
<tr>
<td>20. With air valve open, Air percentage read after lightly tapping the gauge to stabilize the hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>21. Air valve closed and then petcocks opened to release pressure before removing the cover?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>22. Air content recorded to 0.1 percent?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>23. All calculations performed correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner ________________________________

This checklist is derived, in part, from copyrighted material printed in ACI CP-1, published by the American Concrete Institute.

Comments:

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WSDOT FOP for AASHTO T 166¹

Bulk Specific Gravity of Compacted Hot Mix Asphalt Using Saturated Surface-Dry Specimens

1. SCOPE

1.1 This method of test covers the determination of bulk specific gravity of specimens of compacted hot mix asphalt.

1.2 Definition:

1.3 Bulk specific gravity (of solids)—the ratio of the weight mass in air of a unit volume of a permeable material (including both permeable and impermeable voids normal to the material) at a stated temperature to the weight in air of equal density of an equal volume of gas-free distilled water at a stated temperature. The form of the expression shall be:

\[ \frac{\text{Bulk specific gravity}}{\text{x/y °C}} \]

where:

\[ x = \text{temperature of the material, and} \]
\[ y = \text{temperature of the water} \]

1.4 This method should not be used with samples that contain open or interconnecting voids and/or absorb more than 2 percent of water by volume, as determined in Sections 6.2 or 9.2 herein.

1.5 The bulk specific gravity of the compacted hot mix asphalt may be used in calculating the unit mass of the mixture.

1.6 The values stated in English SI units are to be regarded as the standard.

   Note: Method A shall be used for laboratory compacted specimens, and field specimens compacted using gyratory compactor.

   Method C shall be used for asphalt pavement cores.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:

   • M 231, Weighing Devices Used in the Testing of Materials
   • T 275, Bulk Specific Gravity of Compacted Bituminous Mixtures Using Paraffin-Coated Specimens

3. TEST SPECIMENS

3.1 Test specimens may be either laboratory-molded HMA asphalt mixtures or from HMA pavements. The mixtures may be surface, wearing, leveling or base course materials, or hot mix base.

3.2 Size of Specimens — It is recommended that: (1) the diameter of cylindrically molded or cored specimens, or the length of the sides of sawed specimens, be at least equal to four times the maximum size of the aggregate; and (2) the thickness of specimens be at least one-and-one-half times the maximum size of the aggregate.

¹This Test Method is based on AASHTO T 166-05.
3.3 Specimens shall be taken from pavements with core drill, diamond or carborundum saw, or by other suitable means.

3.4 Care shall be taken to avoid distortion, bending, or cracking of specimens during and after the removal from pavement or mold. Specimens shall be stored in a safe, cool place.

3.5 Specimens shall be free from foreign materials such as seal coat, tack coat, foundation material, soil, paper, or foil.

3.6 If desired, specimens may be separated from other pavement layers by sawing or other suitable means. Care should be exercised to ensure sawing does not damage the specimens.

METHOD A

4. APPARATUS

4.1 Weighing Device — The weighing device shall have sufficient capacity, be readable to 0.1 percent of the sample specimen mass, or better, and conform to the requirements of AASHTO M 231. The weighing device shall be equipped with suitable suspension apparatus and holder to permit weighing the specimen while suspended from the center of scale pan of the weighing device.

4.2 Suspension Apparatus — The wire suspending the container shall be the smallest practical size to minimize any possible effects of a variable immersed length. The suspension apparatus shall be constructed to enable the container to be immersed to a depth sufficient to cover it and the test specimen during weighing. Care should be exercised to ensure no trapped air bubbles exist under the specimen.

4.3 Water Bath — for immersing the specimen in water while suspended under the weighing device, equipped with an overflow outlet for maintaining a constant water level.

5. PROCEDURE

5.1 Dry the specimen to a constant mass (Note 1). Cool the specimen to room temperature at 77 ± 9°F (25 ± 5°C), and record the dry mass as A. Immerse each specimen in water at 77 ± 1.8°F (25 ± 1°C) for 4 ± 1 minute and record the immersed mass as C. Remove the specimen from the water, damp dry the specimen by blotting with a damp towel as quickly as possible, and determine the surface-dry mass as, B. Any water that seeps from the specimen during the weighing operation is considered part of the saturated specimen (Note 1). Each specimen shall be immersed and weighed individually.

Note 1: Constant mass shall be defined as the mass at which further drying at 125 ± 5°F (52 ± 3°C) does not alter the mass by more than 0.1 ± 0.05 percent. Sample Specimen saturated with water shall initially be dried overnight at 125 ± 5°F (52 ± 3°C) and then weighed at 2-hour drying intervals. Recently molded laboratory specimens which have not been exposed to moisture do not require drying.

Note 2: If desired, the sequence of testing operations may be changed to expedite the test results. For example, first the immersed mass (C) can be taken, then the surface-dry mass (B), and finally the dry mass (A).

Note 3: Terry cloth has been found to work well for an absorbent cloth. Damp is considered to be when no water can be wrung from towel.
6. **CALCULATION**

6.1 Calculate the bulk specific gravity of the specimens as follows (round and report the value to the nearest three decimal places):

\[
\text{Bulk Sp. Gr.} = \frac{A}{B-C}
\]

where:

- \(A\) = mass in grams of sample specimen in air,
- \(B\) = mass in grams of surface-dry specimen in air,
- \(C\) = mass in grams of sample specimen in water.

6.2 Calculate the percent water absorbed by the specimen (on volume basis) as follows:

\[
\text{Percent Water Absorbed by Volume} = \left(\frac{B-A}{B-C}\right) \times 100
\]

6.3 If the percent water absorbed by the specimen in Section 5.2 exceeds 2 percent, use T 275 (Bulk Specific Gravity of Compacted Bituminous Mixtures Using Paraffin-Coated Specimens) to determine the bulk specific gravity.

**METHOD B**

WSDOT does not use Method B and has removed this section from the procedure.

**METHOD C (RAPID TEST)**

10. **PROCEDURE**

10.1 This procedure can be used for testing specimens which are not required to be saved and which contain substantial amount of moisture. Specimens obtained by coring or sawing can be tested the same day by this method.

10.2 The testing procedure shall be the same as given in Sections 4 and 7 except for the sequence of operations. The dry mass \(A\) of the specimen is determined last as follows.

*Note 4:* A microwave oven can be used to speed up the process by initially heating the sample so that it can be broken into small pieces prior to placing it into the drying oven.

10.3 Place the specimen in a large flat bottom drying pan of known mass. Place the pan and specimen in a \(230 \pm 9^\circ F (110 \pm 5^\circ C)\) \(325 \pm 25^\circ F (164 \pm 14^\circ C)\) oven. Leave the specimen in the oven until it can be easily separated to the point where the particles of the fine aggregate-asphalt portion are not larger than 1/4 in. (6.4 mm). Place the separated specimen in the \(230^\circ F\) \((110^\circ C)\) \(325^\circ F\) \((164^\circ C)\) oven and dry to a constant mass. Constant mass shall be defined as the mass at which further drying at \(230 \pm 9^\circ F (110 \pm 5^\circ C)\) \(325 \pm 25^\circ F (164 \pm 14^\circ C)\) does not alter the mass by more than 0.1 to 0.5 percent when weighed at 2 hour intervals.

*Note:* If samples are placed in the oven overnight for a minimum of 6 hours at \(230^\circ F\), then the 2 hour weighting is not necessary.

10.4 Cool the pan and specimen to room temperature at \(77 \pm 9^\circ F (25 \pm 5^\circ C)\). Determine the mass of the pan and specimen, subtract the mass of the pan and record the dry mass of the pan and record the dry mass, A.
11. CALCULATIONS
   11.1 Calculate the bulk specific gravity in Sections 6.1 and 8.1.

12. REPORT
   12.1 The report shall include the following:
       12.1.1 The method used (A, B, or C).
       12.1.2 Bulk Specific Gravity reported to the nearest thousandth. (0.001)
       12.1.3 Absorption reported to the nearest hundredth. (0.01)

13. PRECISION
   13.1 Duplicate specific gravity results by the same operator should not be considered suspect unless they differ more than 0.02.
Performance Exam Checklist

Bulk Specific Gravity of Compacted HOT MIX ASPHALT Mixtures Using Saturated Surface-dry Specimens
AASHTO T 166

| Participant Name | ________________________________ | Exam Date | _____________ |

**Procedure Element**

<table>
<thead>
<tr>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

1. The tester has a copy of the current procedure on hand?

2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?

**Method A** (For use with laboratory compacted specimens.)

1. Compacted specimen cooled to room temperature, $77 \pm 9$ F, and record the dry mass?

2. Immerse each specimen in water at $77 \pm 2$ F for 3 to 5 minutes and record the immersed mass to the nearest 0.1 gram?

3. Remove sample from water, surface dry with damp towel and weigh the specimen in air at $77 \pm 9$ F to the nearest 0.1 gram?

4. Calculate the bulk specific gravity of the specimens by following the calculation in AASHTO T166 (Section 5.1)?

**Method C** (For use with pavement cores and chunks.)

1. Immerse specimen in water at $77 \pm 2$ F for 3 to 5 minutes and record the immersed weight to the nearest 0.1 gram?

2. Remove sample from water, surface dry by blotting with damp towel and immediately weigh specimen in air at $77 \pm 9$ F to the nearest 0.1 gram?

3. Place specimen in container (noting the empty container weight), then into an oven set at $230 \pm 9$ F until sample can be broken into small pieces?

4. Return container to oven until it has reached a constant weight?

5. Remove container and sample from oven and allow to cool to room temperature, $77 \pm 9$ F?

6. Weigh pan with sample and record to nearest 0.1 gram, deducting known weight of pan to arrive at oven-dried sample weight?

7. Calculate the bulk specific gravity of the specimen by following the calculation in AASHTO T166 (Section 5.1)?

*First attempt: Pass [ ]  Fail [ ]  Second attempt: Pass [ ]  Fail [ ]

Signature of Examiner __________________________________________
WSDOT FOP FOR WAQTC/AASHTO T168
SAMPLING OF HOT MIX ASPHALT PAVING MIXTURES
FOP FOR WAQTC T 168

SIGNIFICANCE

Testing bituminous paving mixtures in the field begins with obtaining and preparing the sample to be tested. Standardized procedures for obtaining a representative sample have been established. Producing strong, durable, reliable pavement in roadways requires careful sampling and accurate testing.

Technicians must be patient and follow these procedures. If one considers that the specifications require quality tests to be made on only a small portion of the total material placed, the need for a truly representative sample is apparent. For this reason, every precaution must be taken to obtain a sample that is truly representative of the entire batch and then to protect that sample from contamination and physical damage.

SCOPE

This procedure covers the sampling of bituminous paving mixtures from HMA plants, haul units, and roadways in accordance with AASHTO T 168. Sampling is as important as testing, and every precaution must be taken to obtain a truly representative sample. The sampling of aggregate used in bituminous paving mixtures shall be in accordance with the FOP for AASHTO T 2.

The Standard Specifications require that sample of Hot Mix Asphalt mixtures be taken from the hauling vehicle.

APPARATUS

- Flat-bottomed scoop 150 x 400 x 100 mm (6 x 16 x 4 in.) if sampling from a roadway
- Shovel
- Sample containers: such as cardboard boxes, metal cans, stainless steel bowls, or other agency-approved containers
- Template to match conveyor belt shape
- Scoops, trowels, or other equipment to obtain mix
- Sampling plate: heavy gauge metal plate 380 mm x 380 mm (15 in x 15 in) minimum 8-gauge thick with a wire attached to one corner long enough to reach from the center of the paver to the outside of the farthest auger extension. Holes ¼ in diameter should be provided in each corner.
- Cookie cutter sampling device: A 330mm (13 in.) square sampling template, constructed from 75mm x 50mm x 3mm (3 in. x 2 in. x 1/8 in.) formed steel angle with two 100mm x 150 mm x 9mm (4 in. x 6 in. x 3/8 in. handles. See diagram
GENERAL COMMENTS

1. Samples of mix upon which acceptance or rejection is based shall be selected at random, and may be obtained by, or under the observation of, the purchaser or authorized representative. Random selection will be accomplished by using WSDOT Test Method T716, Method of Random Sampling for Locations of Testing and Sampling Sites.

Note 2: Care shall be taken to prevent contamination of bituminous mixes by dust or other foreign matter, and to avoid segregation of aggregate and bituminous materials.

2. Some agencies require mechanical sampling devices for hot mix asphalt (HMA) and cold feed aggregate on some projects. These are normally permanently attached devices that allow a sample container to pass perpendicularly through the entire stream of material or divert the entire stream of material into the container. Operation may be hydraulic, pneumatic, or manual and allows the sample container to pass through the stream twice, once in each direction, without overfilling. Special caution is necessary with manually operated systems since a consistent speed is difficult to maintain and non-representative samples may result. Check agency requirements for the specifics of required sampling systems.

SAMPLE SIZE

Sample size depends on the test methods specified by the agency for acceptance. Check agency requirement for the size required.

WSDOT requires a minimum of four times the amount required for testing. This should be approximately 125 lbs.

SAMPLING

- General

1. The material shall be inspected to determine variations. The seller shall provide one of the following,
   a. A mechanical sampling device attached to the HMA plant.
   b. Platforms or devices to enable sampling from the hauling vehicle without entering the hauling vehicle for sampling HMA.

   equipment for safe and appropriate sampling including sampling devices on plants, when required.

2. Place dense graded mixture samples in cardboard boxes or stainless steel bowls or other agency approved containers. Place open graded mixture samples in stainless steel bowls. Do not put open graded mixture samples in boxes until they have cooled to the point that bituminous material will not migrate from the aggregate.

3. Sampling from the Roadway will require the contractor to repair the sampled location.

- Sampling from a Conveyor Belt

WSDOT has deleted this section.

- Attached Sampling Devices

WSDOT has revised this section see General 1b above.
• **Sampling from Truck Transports**
  1. Obtain samples in four approximately equal increments from haul units.
  2. Obtain each increment from approximately 12 in. (300 mm) below the surface, in each of the four quadrants of the load.
  3. Combine the increments to form a sample of the required size.

• **Sampling from a Roadway Prior to Compaction (Scoop Method)**
  WSDOT has deleted this section.

• **Sampling from Roadway Prior to Compaction (Plate Method)**
  WSDOT has deleted this section.

**IDENTIFICATION AND SHIPPING**

1. Identify sample containers as required by the agency.
2. Ship samples in containers that will prevent loss, contamination, or damage.
3. Refer to the sample identification requirements in FOP for WSDOT Test Method 712.
Performance Exam Checklist

**WSDOT FOP FOR WAQTC/AASHTO T 168**
**SAMPLING OF HOT MIX ASPHALT PAVING MIXTURES**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
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</table>

<table>
<thead>
<tr>
<th>Procedure Element</th>
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<th>No</th>
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</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
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<td>☐</td>
</tr>
<tr>
<td>2. Containers of correct type and ample size available?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Samples from truck transports taken from four quadrants at approximately depth 12 inches?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Sample size meets agency requirements?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Sample identified as required?</td>
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</tr>
</tbody>
</table>

First attempt:  Pass ☐  Fail ☐

Second attempt:  Pass ☐  Fail ☐

Signature of Examiner __________________________________________

Comments:

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WSDOT FOP For AASHTO T 176

Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test

1. SCOPE

1.1 This test is intended to serve as a rapid field test to show the relative proportions of fine dust or claylike material in soils or graded aggregates.

1.2 The following applies to all specified limits in this standard: For the purpose of determining conformance with these specifications, an observed value or a calculated value shall be rounded off “to the nearest unit” in the last right-hand place of figures used in expressing the limiting value, in accordance with R 11, Recommended Practice for Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values.

1.3 The values stated in English units are to be regarded as the standard.

1.4 Refer to R 16 for regulatory information for chemicals.

2. APPARATUS

2.1 A graduated plastic cylinder, rubber stopper, irrigator tube, weighted foot assembly, and siphon assembly, all conforming to their respective specifications and dimensions shown in Figure 1. Fit the siphon assembly to a 1 gal (4-L) bottle of working calcium chloride solution (see Section 2.8) placed on a shelf 36 ± 1 in. (915 ± 25 mm) above the work surface. In lieu of the specified 1 gal (4-L) bottle, a glass or plastic vat having a larger capacity may be used provided the liquid level of the working solution is maintained between 36 and 46 inches (915 and 1170 mm) above the work surface. (See Figure 2.)

Note 1: An older model of weighted foot assembly has a guide cap that fits over the upper end of the graduated cylinder and centers the rod in the cylinder, and the foot of the assembly has a conical upper surface and three centering screws to center it loosely in the cylinder. The older model does not have the same reading indicator affixed to the rod (Figure 1), but a slot in the centering screws of the weighted foot is used to indicate the sand reading. Apparatus with the sand reading indicator (Figure 1) is preferred for testing clayey materials.

Note 2: Glass or stainless steel may be substituted as a material type for the copper siphon and blow tubing.

¹This FOP is based on AASHTO T 176-02.
ASSEMBLY B

Note: all dimensions are shown in mm unless otherwise indicated.

FIGURE 1 Sand Equivalent Apparatus

WEIGHT
ASSEMBLY C

Note: all dimensions are shown in mm unless otherwise indicated.

FIGURE 1 Sand Equivalent Apparatus (continued)
# LIST OF MATERIAL

<table>
<thead>
<tr>
<th>Assembly</th>
<th>No.</th>
<th>Description</th>
<th>Stock size</th>
<th>Material</th>
<th>Heat Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>1</td>
<td>Siphon Tube</td>
<td>6.4 dia X 400</td>
<td>Copper Tube</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Siphon Hose</td>
<td>4.8 I.D. X 1220</td>
<td>Rubber Tube</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Blow Hose</td>
<td>4.8 I.D. X 50.8</td>
<td>Rubber Tube</td>
<td></td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>Blow Tube</td>
<td>6.4 dia X 50.8</td>
<td>Copper Tube</td>
<td></td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>Two-Hole Stopper</td>
<td>No. 6</td>
<td>Rubber</td>
<td></td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>Irrigating Tube</td>
<td>6.4 O.D. 0.89 Wall X 500 Stainless Steel Tube, Type 316</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>Clamp</td>
<td>Pinchcock, Day, BKH No. 21730 or Equiv.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>8</td>
<td>Tube</td>
<td>38.1 O.D. X 430</td>
<td>Trans. Acrylic Plastic</td>
<td></td>
</tr>
<tr>
<td></td>
<td>9</td>
<td>Base</td>
<td>12.7 X 102 X 102</td>
<td>Trans. Acrylic Plastic</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>10</td>
<td>Sand Reading Indicator</td>
<td>6.4 dia. X 14.9</td>
<td>Nylon 101 Type 66 Annealed</td>
<td></td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>Rod</td>
<td>6.4 dia. X 438.2</td>
<td>Brass</td>
<td></td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>Weight</td>
<td>0.8 dia. X 52.78</td>
<td>C.R. SH</td>
<td></td>
</tr>
<tr>
<td></td>
<td>13</td>
<td>Roll Pin</td>
<td>0.16 dia. X 12.7</td>
<td>Steel</td>
<td></td>
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<tr>
<td></td>
<td>14</td>
<td>Foot</td>
<td>0.16 Hex. X 13.7</td>
<td>Brass</td>
<td></td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>Solid Stopper</td>
<td>No. 7</td>
<td>Rubber</td>
<td></td>
</tr>
</tbody>
</table>

Notes:

1. "C" Mounted Foot Assembly to weigh 1000 ± 5 g.
2. Graduations of graduate to be 2.54 mm apart and every tenth mark to be numerically designated as shown. Every fifth line should be approximately 9.5 mm long. All other lines should be approximately 5.5 mm long. Depth to be 0.4 mm. Width to be 0.8 mm across the top.
3. Accuracy of scale to be ± 0.25 mm. Error at any point on scale to be ± 0.75 mm of true distance to zero.
4. Glass or stainless steel may be substituted as a material type for the copper siphon and blow tubing.

---

**FIGURE 1 Sand Equivalent Apparatus (continued)**

2.2 A tinned measure, having a capacity of 3 oz (85 ± 5 mL), approximately 2.25 in. (57 mm) in diameter.

2.3 A wide-mouth funnel approximately 4 in. (100 mm) in diameter at the mouth.

2.4 A clock or watch reading in minutes and seconds.

2.5 A mechanical shaker having a throw of 8.00 ± 0.04 in. (203.2 ± 1.0 mm) and operating at 175 ± 2 cycles per minute (2.92 ± 0.03 Hz) (Note 2). Prior to use, fasten the mechanical sand equivalent shaker securely to a firm and level mount.

*Note 2:* The mechanical shaker shall be used when performing referee sand equivalent determinations. Either the mechanical or manually operated shaker should be used in lieu of the hand method whenever possible.

2.6 A manually operated shaker capable of producing an oscillating motion at the rate of 100 complete cycles in 45 ± 5 seconds, with a hand-assisted half stroke length of 5.0 ± 0.2 in. (127 ± 5 mm). The shaker shall be fastened securely to a firm and level mount by bolts or clamps if a large number of determinations are to be made.
2.7 **Stock Solution** – Shall meet the requirements of AASHTO T 176.

2.8 Working calcium chloride solution: Prepare the working calcium chloride by diluting one measuring tin full 3 oz. (85 ± 5 mL), or from a graduated cylinder of the stock calcium chloride solution to 1 gal (3.8 L) with water [finished product will equal 1 gallon]. Use distilled or demineralized water for the normal preparation of the working solution. However, if it is determined that the local tap water is of such purity that it does not affect the test results, it is permissible to use in lieu of distilled or demineralized water except in the event of dispute. Record the date made on the gallon bottle. Working solutions more than 30 days old shall be discarded.

2.9 A straightedge or spatula, suitable for striking off the excess soil from the tin measure.

2.10 A thermostatically controlled drying oven capable of maintaining a temperature of 230 ± 9°F (110 ± 5°C), or other suitable sources of heat may be used, such as an electric or gas hot plate, electric heat lamp, or a ventilated microwave oven.

2.11 Quartering or splitting cloth, approximately 2 ft square, nonabsorbent material such as plastic or oil cloth.

2.12 Optional Handle for Irrigation Tube — A 25-mm diameter wooden dowel to aid in pushing the irrigation tube into firm materials. See Figure 1, Assembly B.

3. **CONTROL**

3.1 The temperature of the working solution should be maintained at 67-77 72 ± °F (22 ± 3°C) during the performance of this test. If field conditions preclude the maintenance of the temperature range, frequent reference samples should be submitted to a laboratory where proper temperature control is possible. It is also possible to establish temperature correction curves for each material being tested where proper temperature control is not possible. However, no general correction curve should be utilized for several materials even within a narrow range of sand equivalent values. Samples which meet the minimums and equivalent requirement at a working solution temperature below the recommended range need not be subject to reference testing.

4. **SAMPLE PREPARATION**

4.1 The sand equivalent test shall be performed on soils or graded aggregate materials passing the No. 4 (4.75-mm) sieve. All aggregations of fine-grained soil material shall be pulverized to pass the No. 4 (4.75-mm) sieve, and all fines shall be cleaned from the particles retained on the No. 4 (4.75-mm) sieve and included with the material passing the No. 4 (4.75-mm) sieve.

4.2 Split or quarter enough of the original sample to yield approximately 1,000 g, slightly more than eight 3 oz (85 mL) tin measures of material passing the No. 4 (4.75-mm) sieve. Use extreme care to obtain a truly representative portion of the original sample (Note 5).

Sieve the aggregate past the No. 4 sieve with a mechanical shaker per FOP for WAQTC/ AASHTO T27/11 at SSD or drier. Use caution to avoid overloading the No. 4 sieve, additional sieving may be necessary.

2Available from Aldrich Chemical Company, P.O. Box 2060, Milwaukee, WI 53201 or Fisher Scientific, 711 Forbes Ave., Pittsburg, PA 15219

3Kathon CG/ICP may be obtained from Rohm and Hass Chemical Company, Independence Mall West, Philadelphia, PA 19105
Note 5: Experiments show that as the amount of material being reduced by splitting or quartering is decreased, the accuracy of providing representative portions is reduced. It is imperative that the sample be split or quartered carefully. When it appears necessary, dampen the material before splitting or quartering, to avoid segregation or loss of fines.

4.3 Prepare the desired number of two test samples by one of the following methods:

4.3.1 Alternate Method No. 1 — Air Dry

4.3.1.1 Split or quarter enough material from the portion passing the No. 4 (4.75-mm) sieve to fill the 30 oz (85-mL) tin measure so it is slightly rounded above the brim. While filling the measure, tap the bottom edge of the tin on the worktable or other hard surface to cause consolidation of the material and allow the maximum amount to be placed in the tin. Strike off the tin measure level full with a spatula or straightedge. For the second determination, remix the sample, quarter and fill the tin again.

4.3.2 Alternate Method No. 2 — Pre-Wet

4.3.2.1 The sample must be in the proper moisture condition to achieve reliable results. This condition is determined by tightly squeezing a small portion of the thoroughly mixed sample in the palm of the hand. If the cast that is formed permits careful handling without breaking, the correct moisture range has been obtained. If the material is too dry, the cast will crumble and it will be necessary to add water and remix and retest until the material forms a cast. If the material shows any free water it is too wet to test and must be drained and air-dried, mixing it frequently to insure uniformity. This overly wet material will form a good cast when checked initially, so the drying process should continue until a squeeze check on the drying material gives a cast which is more fragile and delicate to handle than the original. If the moisture content of the original sample prepared in Section 4.2 is within the limits described above, the test sample may be obtained immediately. If the moisture content is altered to meet these limits, the altered sample should be placed in a pan, covered with a lid or with a damp cloth which does not touch the material, and allowed to stand for a minimum of 15 minutes.

4.3.2.2 After the minimum 15-minute tempering period, Place the sample on the splitting cloth and mix by alternately lifting each corner of the cloth and pulling it over the sample toward the diagonally opposite corner, causing the material to be rolled. When the material appears homogeneous, finish the mixing with the sample in a pile near the center of the cloth.

4.3.2.3 Fill the 3-oz (85-mL) tin measure by pushing it through the base of the pile while exerting pressure with the hand against the pile on the side opposite the measure. As the tin is moved though the pile, hold enough pressure with the hand to cause the material to fill the tin to overflowing. Press firmly with the palm of the hand, compacting the material and allowing the maximum amount to be placed in the tin. Strike off the tin measure level full with a spatula or straightedge. For the second determination, remix the sample and fill the tin again.
Dry the test sample to constant mass in accordance with FOP for AASHTO T 255, and cool to room temperature before testing. It is acceptable to place the test sample in a larger container to aid drying.

4.3.3 Reference Method (Mechanical Shaker) — Obtain the 3-oz (85-mL) tin measure of material by one of the alternate methods, Section 4.3.1 or 4.3.2, above; then dry the test sample to constant mass at 230 ± 9°F (110 ± 5°C), and cool to room temperature before testing.

5. PROCEDURE

5.1 Start the siphon by forcing air into the top of the solution bottle through the bent copper, glass, or stainless steel blow tube while the pinch clamp is open. The apparatus is now ready for use.

5.2 Siphon 4.0 ± 0.1 in. (101.6 ± 2.5 mm) of working calcium chloride solution into the plastic cylinder. Pour the prepared test sample from the measuring tin into the plastic cylinder using the funnel to avoid spillage. (See Figure 3). Tap the bottom of the cylinder sharply on the heel of the hand several times to release air bubbles and to promote thorough wetting of the sample.

5.3 Allow the wetted sample to stand undisturbed for 10 ± 1 minute. At the end of the 10-minute soaking period, stopper the cylinder, then loosen the material from the bottom by partially inverting the cylinder and shaking it simultaneously.

5.4 After loosening the material from the bottom of the cylinder, shake the cylinder and contents by any one of the following methods:

5.4.1 Mechanical Shaker Method (Reference Method) — Place the stoppered cylinder in the mechanical sand equivalent shaker, set the timer, and allow the machine to shake the cylinder and contents for 45 ± 1 second.

5.4.2 Manual Shaker Method — Secure the stoppered cylinder in the three spring clamps on the carriage of the hand-operated sand equivalent shaker and reset the stroke counter to zero. Stand directly in front of the shaker and force the pointer to the stroke limit marker painted on the backboard by applying an abrupt horizontal thrust to the upper portion of the right hand spring steel strap. Then remove the hand from the strap and allow the spring action of the straps to move the carriage and cylinder in the opposite direction without assistance or hindrance. Apply enough force to the right hand spring steel strap during the thrust portion of each stroke to move the pointer to the stroke limit marker by pushing against the strap with the ends of the fingers to maintain a smooth oscillating motion. The center of the stroke limit marker is positioned to provide the proper stroke length and its width provides the maximum allowable limits of variation. The proper shaking action is accomplished only when the tip of the point reverses direction within...
the marker limits. Proper shaking action can best be maintained by using only the forearm and wrist action to propel the shaker. Continue the shaking action for 100 strokes.

5.4.3 Hand Method — Hold the cylinder in a horizontal position as illustrated in Figure 5 and shake it vigorously in a horizontal linear motion from end to end. Shake the cylinder 90 cycles in approximately 30 seconds using a throw of 9 ± 1 in. (229 ± 25 mm). A cycle is defined as a complete back and forth motion. To properly shake the cylinder at this speed, it will be necessary for the operator to shake with the forearms only, relaxing the body and shoulders.

![Manually-operated shaker](image)

Manually-operated shaker

Figure 4

5.5 Following the shaking operation, set the cylinder upright on the work table and remove the stopper.

5.6 Irrigation Procedure — Insert the irrigator tube in the cylinder and rinse material from the cylinder walls as the irrigator is lowered. Force the irrigator through the material to the bottom of the cylinder by applying a gentle stabbing and twisting action while the working solution flows from the irrigator tip. This flushes the fine material into suspension above the coarser sand particles, (See Figure 5.) Continue to apply the stabbing and twisting action while flushing the fines upward until the cylinder is filled to the 15 in. (381 mm) mark. Then raise the irrigator slowly without shutting off the flow so that the liquid level is maintained at above 15 in. (381 mm) while the irrigator is being withdrawn. Regulate the flow just before the irrigator is entirely withdrawn and adjust the final level to 15 in. (381 mm). Final level as judged by the bottom of the meniscus shall be between the top two gradations on the tube but shall not be above the 15 in. (381 mm) level.

*Note 7:* For certain soils, particularly on crushed materials, the stabbing action may not be possible. For these materials, the irrigation technique is as follows: Continue to apply a twisting action as the irrigation tube is slowly withdrawn. As the tube is withdrawn, it is essential that as many fines as possible flushed upward until the cylinder is filled to the 15 in (381 mm) mark.

5.7 Allow the cylinder and contents to stand undisturbed for 20 minutes ± 15 seconds. Start the timing immediately after withdrawing the irrigator tube.
5.8 At the end of the 20 minute sedimentation period, read and record the level of the top of the clay suspension. This is referred to as the “clay reading.” If no clear line of demarcation has formed at the end of the specified 20 minute sedimentation period, allow the sample to stand undisturbed until a clear reading can be obtained, then immediately read and record the level of the top of the clay suspension and the total sedimentation time. If the total sedimentation time exceeds 30 minutes, it will be rejected rerun the test using three individual samples of the same material. Read and record the clay column height of that sample requiring the shortest sedimentation period only.

5.9 After the clay reading has been taken, the “sand reading” shall be obtained by one of the following methods:

5.9.1 When using the weighted foot assembly having the sand indicator on the rod of the assembly, place the assembly over the cylinder and gently lower the assembly toward the sand. Do not allow the indicator to hit the mouth of the cylinder as the assembly is being lowered. As the weighted foot comes to rest on the sand, tip the assembly toward the graduations on the cylinder until the indicator touches the inside of the cylinder. Subtract 10 in. (254 mm) from the level indicated by the extreme top edge of the indicator and record this value as the “sand reading.” (See Figure 6.)

5.9.2 If an older model weighted foot assembly having centering screws is used, keep one of the centering screws in contact with the cylinder wall near the graduations so that it can be seen at all times while the assembly is being lowered. When the weighted foot has come to rest on the sand, read the level of the centering screw and record this value as the “sand reading.”
5.10 If clay or sand readings fall between 0.1 in. (2.5 mm) graduations, record the level of the higher graduation as the reading. For example, a clay reading of 7.95 would be recorded as 8.0, and a sand reading of 3.22 would be recorded as 3.3.

**Note 8:** Samples obtained from aggregate proposed for use in hot asphalt paving mixtures shall be prepared by oven-drying if acceptance of the material is based on tests on material that has passed through a hot plant drier.

6. **CALCULATIONS**

6.1 Calculate the sand equivalent (SE) to the nearest 0.1 using the following formula:

\[
SE = \frac{\text{Sand Reading} \times 100}{\text{Clay Reading}}
\]

6.2 If the calculated sand equivalent is not a whole number, report it as the next higher whole number, as in the following example:

\[
SE = \frac{3.3}{8} \times 100 = 41.25
\]

which is reported as 42.

6.3 If it is desired to average a series of sand equivalent values, average. Average the whole number values determined as described above. If the average of these values is not a whole number, raise it to the next higher whole number, as in the following example:

Calculated SE values: 41.2, 43.8, 40.9

After raising each to the next higher whole number, they become: 42, 44, 41.

The average of these values is then determined:

\[
\frac{42 + 44 + 41}{3} = 41.5
\]

Which is reported as 42.

If the two results from the same SE sample vary by more than 8 points, the test shall be invalid and a new test completed.

6.3.1 Since the average value is not a whole number, it is raised to the next higher whole number and the reported averages and equivalent value is reported as 43 42.

Report the results using WSDOT Form 350-161 EF, 422-020X, or other report approved by the State Materials Engineer.

7. **PRECAUTIONS**

See AASHTO T 176 for Precision

8. **OPERATOR QUALIFICATIONS**

WSDOT has deleted this section see Section 9-5.5 of the Construction Manual.
Performance Exam Checklist

Plastic Fines in Graded Aggregates and Soils by the Use of the Sand Equivalent Test
FOP for AASHTO T 176

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Preparation</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Sample passed through No. 4 (4.75 mm) sieve?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Split or quarter proper amount of material?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Material in clods broken up and re-screened?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. No fines lost?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Temperature of working solution 72±5 F (22 ±3°C)?</td>
<td>☐</td>
<td>☐</td>
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<tr>
<td>8. Working calcium chloride solution 36 ± 1 in. (915 mm ± 25 mm) above the work surface?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. 4 ± 0.1 in (101.6 ± 2.5 mm) working calcium chloride solution siphoned into cylinder?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. Working solution dated?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td><strong>Sample Preparation</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. If necessary, sample sprayed with water to prevent loss of fines?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Material checked for moisture condition by tightly squeezing small portion in palm of hand and forming a cast?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Sample at proper water content?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>a. If too dry, (cast crumbles easily), water added and re-mixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>b. If too wet (shows free water), sample drained, air dried and mixed frequently?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Sample placed on splitting cloth and mixed by alternately lifting each corner of the cloth and pulling it over the sample toward diagonally opposite corner, causing material to be rolled?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Is material thoroughly mixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. When material appears to be homogeneous, mixing finished with sample in a pile near center of cloth?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Fill the 85 mL tin by pushing through base of pile with other hand on opposite side of pile?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>
Procedure Element | Yes | No
--- | --- | ---
8. Material fills tin to overflowing? | [ ] | [ ]
9. Material compacted into tin with palm of hand? | [ ] | [ ]
10. Tin struck off level full with spatula or straightedge? | [ ] | [ ]
11. Test sample dried to a constant mass? | [ ] | [ ]
12. Sample cooled to room temperature | [ ] | [ ]

Procedure
1. Prepared sample funneled into cylinder with no loss of fines? | [ ] | [ ]
2. Bottom of cylinder tapped sharply on heel of hand several times to release air bubbles? | [ ] | [ ]
3. Wetted sample allowed to stand undisturbed for 10 min. ± 1 min.? | [ ] | [ ]
4. Cylinder stoppered and material loosened from bottom by shaking? | [ ] | [ ]
5. Properly performed shaking method?
   Mechanical Shaker Method | [ ] | [ ]
   Manual Shaker Method | [ ] | [ ]
6. Following shaking, cylinder set vertical on work surface and stopper removed? | [ ] | [ ]
7. Irrigator tube inserted in cylinder and material rinsed from cylinder walls as irrigator is lowered? | [ ] | [ ]
8. Irrigator tube forced through material to bottom of cylinder by gently stabbing and twisting action? | [ ] | [ ]
9. Stabbing and twisting motion applied until cylinder filled to 15 in. (381 mm) mark? | [ ] | [ ]
10. Liquid raised and maintained at 15 in. (381 mm) mark while irrigator is being withdrawn? | [ ] | [ ]
11. No clear solution at top of column? | [ ] | [ ]
12. Contents let stand 20 minutes ± 15 seconds? | [ ] | [ ]
13. Timing started immediately after withdrawal of irrigator? | [ ] | [ ]
14. No vibration or disturbance of the sample? | [ ] | [ ]
15. Readings taken at 20 minutes or up to 30 minutes, when a definite line appears? | [ ] | [ ]
16. Weighted foot assembly lowered into cylinder without hitting mouth of cylinder? | [ ] | [ ]
17. Calculations made to 0.1 and reported to the next higher whole number? | [ ] | [ ]
18. SE is based on the average results of two samples? | [ ] | [ ]
19. If the two SE values vary by more than 8 points additional tests run? | [ ] | [ ]
20. All calculations performed correctly? | [ ] | [ ]
First attempt:  Pass ☐ Fail ☐  
Second attempt:  Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

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WSDOT FOP for AASHTO T 180
Moisture–Density Relations of Soils Using a 10-lb (4.54-kg) Rammer and an 18-in. (457-mm) Drop

1. SCOPE

1.1. This method of test is intended for determining the relationship between the moisture content and density of soils when compacted in a given mold of a given size with a 4.54-kg (10-lb) rammer dropped from a height of 457 mm (18 in.). Four alternate procedures are provided as follows:

• Method A—A 101.60-mm (4-in.) mold: Soil material passing a 4.75-mm (No. 4) sieve Sections 4 and 5.
• Method B—A 152.40-mm (6-in.) mold: Soil material passing a 4.75-mm (No. 4) sieve Sections 6 and 7.
• Method C—A 101.60-mm (4-in.) mold: Soil material passing a 19.0-mm (3/4-in.) sieve Sections 8 and 9.
• Method D—A 152.40-mm (6-in.) mold: Soil material passing a 19.0-mm (3/4-in.) sieve Sections 10 and 11.

The preferred method of WSDOT is to use FOP for AASHTO T 180 Method D.

WSDOT recommends that the bulk specific gravity of coarse aggregate be determined. Native soils within the contract limits to be used for embankment construction and/or backfill material do not require the sampling by a qualified tester. For material that requires gradation testing such as but not limited to manufactured aggregates and Gravel Borrow, a qualified testers shall be required for sampling.

1.2. The method to be used should be indicated in the specifications for the material being tested. If no method is specified, the provisions of Method A shall govern.

1.3. This test method applies to soils mixtures that have 30 percent or less retained on the 4.75-mm (No. 4) sieve, when Method A or B is used and 30 percent or less retained on the 19.0-mm (3/4-in.) sieve, when Method C or D is used. The material retained on these sieves shall be defined as oversize particles (coarse particles).

1.4. If the test specimen contains oversize particles, and the test specimen used for field density compaction control, corrections must be made according to T 224 to compare the total field density with the compacted specimen. The person or agency specifying this method shall specify a minimum percentage of oversize particles below which correction for oversize need not be applied. If no minimum percentage is specified, correction shall be applied to samples with more than five percent by mass of oversize particles.

1.5. If more than 30 percent of the sample is retained on the 3/4 inch (19 mm) sieve, WSDOT test Method T 606 shall be used.

Note 1—One method for the design and control of the compaction of such soils is to use **a test fill to determine the required degree of compaction and a method to obtain that compaction. Then use a method specification to control the compaction by specifying the type and size of compaction equipment, the lift thickness and the number of passes.

1 This Test Method is based on AASHTO T 180-01
1.6. The following applies to all specified limits in this standard: For the purposes of determining conformance with these specifications, an observed value or a calculated value shall be rounded off “to the nearest unit” in the last right-hand place of figures used in expressing the limiting value, in accordance with R 11.

1.7. The values stated in SI units are to be regarded as the standard.

2. referenced documents

2.1. AASHTO Standards:
- M 92, Wire-Cloth Sieves for Testing Purposes
- M 231, Weighing Devices Used in the Testing of Materials
- R 11, Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values
- T 19/T 19M, Bulk Density (“Unit Weight”) and Voids in Aggregate
- T 224, Correction for Coarse Particles in the Soil Compaction Test
- T 265, Laboratory Determination of Moisture Content of Soils

2.2. ASTM Standards
- D 2168, Calibration of Laboratory Mechanical-Rammer Soil Compactors

3. APPARATUS

3.1. Molds—The molds shall be solid-wall, metal cylinders manufactured with dimensions and capacities shown in Sections 3.1.1, 3.1.2 and Figures 1 and 2 below. They shall have a detachable collar assembly approximately 60 mm (2 3/8 in.) in height, to permit preparation of compacted specimens of soil-water mixtures of the desired height and volume. The mold and collar assembly shall be so constructed that it can be fastened firmly to a detachable base plate made of the same material (Note 2). The base plate shall be plane to 0.13 mm (0.005 in.) as shown in Figures 1 and 2.

Note 2—Alternate types of molds with capacities as stipulated herein may be used, provided the test results are correlated with those of the solid-wall mold on several soil types and the same moisture-density results are obtained. Records of such correlation shall be maintained and readily available for inspection, when alternate types of molds are used.

3.1.1. A 101.6-mm (4-in.) mold having a capacity of $0.000943 \pm 0.000008 \text{ m}^3 (1/30 (0.0333) \pm 0.0003 \text{ cu ft})$ with an internal diameter of $101.60 \pm 0.41 \text{ mm (4.000 \pm 0.016 in.)}$ and a height of $116.43 \pm 0.13 \text{ mm (4.584 \pm 0.005 in.)}$ (Figure 1).

3.1.2. A 152.4-mm (6-in.) mold having a capacity of $0.002124 \pm 0.00021 \text{ m}^3 (1/13.33 (0.07500) \pm 0.00075 \text{ cu ft})$ with an internal diameter of $152.40 \pm 0.66 \text{ mm (6.000 \pm 0.026 in.)}$ and a height of $116.43 \pm 0.13 \text{ mm (4.584 \pm 0.005 in.)}$ (Figure 2).

3.1.3. Molds Out of Tolerance Due to Use—A mold that fails to meet manufacturing tolerances after continued service may remain in use provided those tolerances are not exceeded by more than 50 percent; and the volume of the mold, calibrated in accordance with T 19/T 19M, is used in the calculations.
3. Figure 1 is to be used for all compaction molds purchased after the publication of the 21st edition (HM-21).

4. Not to scale.

Notes:

1. All dimensions shown in millimeters unless otherwise noted.

2. Hanger on the mold portion only cannot extend above the midheight line.

3. Figure 1 is to be used for all compaction molds purchased after the publication of the 21st edition (HM-21).

4. Not to scale.

Figure 1—Cylindrical Mold and Base Plate (101.60-mm Mold)
3. Figure 2 is to be used for all compaction molds purchased after the publication of the 21st edition (HM-21).

4. Not to scale.

Figure 2— Cylindrical Mold and Base Plate (152.40-mm Mold)

Notes:
1. All dimensions shown in millimeters unless otherwise noted.
2. Hanger on the mold portion only cannot extend above the midheight line.
3. Figure 2 is to be used for all compaction molds purchased after the publication of the 21st edition (HM-21).
4. Not to scale.
### Dimensional Equivalents for Figure 1

<table>
<thead>
<tr>
<th>mm</th>
<th>in.</th>
<th>mm</th>
<th>in.</th>
<th>Mm</th>
<th>in.</th>
</tr>
</thead>
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<tr>
<td>3.18 ± 0.64</td>
<td>0.125 ± 0.025</td>
<td>17.78 ± 1.27</td>
<td>0.700 ± 0.050</td>
<td>107.95 ± 1.27</td>
<td>4.250 ± 0.050</td>
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<tr>
<td>3.81</td>
<td>0.150</td>
<td>20.32</td>
<td>0.800</td>
<td>114.30 ± 2.54</td>
<td>4.300 ± 0.100</td>
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<tr>
<td>6.35 ± 1.27</td>
<td>0.250 ± 0.050</td>
<td>38.10 ± 2.54</td>
<td>1.500 ± 0.100</td>
<td>116.43 ± 0.13</td>
<td>4.384 ± 0.005</td>
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<td>7.62</td>
<td>0.300</td>
<td>50.80 ± 0.64</td>
<td>2.000 ± 0.025</td>
<td>152.40 ± 2.54</td>
<td>6.000 ± 0.100</td>
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<tr>
<td>9.53 ± 0.64</td>
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<td>2.375 ± 0.050</td>
<td>165.10 ± 2.54</td>
<td>6.500 ± 0.100</td>
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<tr>
<td>12.70 ± 2.54</td>
<td>0.500 ± 0.100</td>
<td>101.60 ± 0.41</td>
<td>4.000 ± 0.016</td>
<td>172.72 ± 2.54</td>
<td>6.800 ± 0.100</td>
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<tr>
<td>0.000943 ± 0.00008 m³</td>
<td>1/30 ± 0.0003 ft³</td>
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</table>

### Dimensional Equivalents for Figure 2

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<th>mm</th>
<th>in.</th>
<th>Mm</th>
<th>in.</th>
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</thead>
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<tr>
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<td>0.125 ± 0.025</td>
<td>17.78 ± 1.27</td>
<td>0.700 ± 0.050</td>
<td>132.40 ± 0.66</td>
<td>6.000 ± 0.026</td>
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<td>3.81</td>
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<td>20.32</td>
<td>0.800</td>
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<td>38.10 ± 2.54</td>
<td>1.500 ± 0.100</td>
<td>165.10 ± 2.54</td>
<td>6.500 ± 0.100</td>
</tr>
<tr>
<td>7.62</td>
<td>0.300</td>
<td>50.80 ± 0.64</td>
<td>2.000 ± 0.025</td>
<td>172.72 ± 2.54</td>
<td>6.800 ± 0.100</td>
</tr>
<tr>
<td>9.53 ± 0.64</td>
<td>0.375 ± 0.025</td>
<td>60.33 ± 2.54</td>
<td>2.375 ± 0.050</td>
<td>203.23 ± 2.54</td>
<td>8.000 ± 0.100</td>
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<tr>
<td>12.70 ± 2.54</td>
<td>0.500 ± 0.100</td>
<td>116.43 ± 0.13</td>
<td>4.384 ± 0.005</td>
<td>215.90 ± 2.54</td>
<td>8.500 ± 0.100</td>
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<tr>
<td>0.000943 ± 0.00008 m³</td>
<td>1/30 ± 0.0003 ft³</td>
<td></td>
<td></td>
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</tbody>
</table>

3.2. Rammer:

3.2.1. Manually Operated—Metal rammer with a mass of 4.536 ± 0.009 kg (10.00 ± 0.02 lb) (Note 3), and having a flat circular face of 50.80 mm (2.000 in.) diameter with a manufacturing tolerance of ±0.25 mm (±0.01 in.). The in-service diameter of the flat circular face shall be not less than 50.42 mm (1.985 in.). The rammer shall be equipped with a suitable guide sleeve to control the height of drop to a free fall 457 ± 2 mm (18.00 ± 0.06 in.) above the elevation of the soil. The guide sleeve shall have at least four vent holes, no smaller than 9.5-mm (3/8-in.) diameter, spaced approximately 90 degrees (1.57 rad) apart and approximately 19 mm (3/4 in.) from each end; and shall provide sufficient clearance so the free fall of the rammer shaft and head is unrestricted.

3.2.2. Mechanically Operated—A metal rammer which is equipped with a device to control the height of drop to a free fall of 457 ± 2 mm (18.00 ± 0.06 in.) above the elevation of the soil, and uniformly distributes such drops to the soil surface (Note 3). The rammer shall have a mass of 4.536 ± 0.009 kg (10.00 ± 0.02 lb) (Note 2), and have a flat circular face of 50.80 mm (2.000 in.) diameter with a manufacturing tolerance of ±0.25 mm (±0.01 in.). The in-service diameter of the flat circular face shall be not less than 50.42 mm (1.985 in.). The mechanical rammer shall be calibrated by ASTM D 2168 to give the same moisture-density results as with a manually operated rammer.

Note 3—The mechanical rammer apparatus shall be calibrated with several soil types and the mass of the rammer adjusted, if necessary, to give the same moisture-density results as with the manually operated rammer.
It may be impractical to adjust the mechanical apparatus so the free fall is 457-mm (18 in.) each time the rammer is dropped, as with the manually operated rammer. To make the adjustment of free fall, the portion of loose soil to receive the initial blow should be slightly compressed with the rammer to establish the point of impact from which the 457-mm (18-in.) drop is determined; subsequent blows on the layer of soil being compacted may all be applied by dropping the rammer from a height of 457 mm (18 in.) above the initial-setting elevation, or when the mechanical apparatus is designed with a height adjustment for each blow, all subsequent blows should have a rammer free fall of 457 mm (18 in.) measured from the elevation of the soil as compacted by the previous blow. A more detailed calibration procedure for laboratory mechanical rammer soil compactors can be found in ASTM D 2168.

3.2.3. Rammer Face—The circular face rammer shall be used but a sector face rammer may be used as an alternative provided the report shall indicate type of face used other than the 50.8-mm (2-in.) circular face and it shall have an area equal to that of the circular face rammer.

3.3. Sample Extruder (For Solid-Walled Molds Only)—A jack, lever, frame, or other device adapted for the purpose of extruding compacted specimen from the mold.

3.4. Balances and Scales—A balance or scale conforming to the requirements of M 231, Class G 20. Also, a balance conforming to the requirements of M 231, Class G 2.

Note 4—The capacity of the metric balance or scale should be approximately 11.5 kg when used to determine the mass of the 152-mm (6-in.) mold and compacted, moist soil; however, when the 102-mm (4-in.) mold is used, a balance or scale of lesser capacity than 11.5 kg may be used, if the sensitivity and readability is 5 g.

3.5. Drying Oven—A thermostatically controlled drying oven capable of maintaining a temperature of 110 ± 5ºC (230 ± 9ºF) for drying moisture samples.

3.6. Straightedge—A hardened steel straightedge at least 250 mm (10 in.) in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.1 percent of the length within the portion used for trimming the soil (Note 5).

Note 5—The beveled edge may be used for final trimming if the edge is true within a tolerance of 0.25 mm per 250 mm (0.1 percent) of length; however, with continued use, the cutting edge may become excessively worn and not suitable for trimming the soil to the level of the mold. The straightedge should not be so flexible that trimming the soil surface with the cutting edge will cause a concave soil surface.

3.7. Sieves—50, 19.0, and 4.75 mm sieves conforming to the requirements of M 92.

3.8. Mixing Tools—Miscellaneous tools such as mixing pan, spoon, trowel, spatula, etc., or a suitable mechanical device for thoroughly mixing the sample of soil with increments of water.

3.9. Containers—Suitable containers made of material resistant to corrosion and not subject to change in mass or disintegration on repeated heating and cooling. Containers shall have close-fitting lids to prevent loss of moisture from samples before initial mass determination and to prevent absorption of moisture from the atmosphere following drying and before final mass determination. One container is needed for each moisture content determination.
METHOD A

4. SAMPLE

4.1. If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus which is maintained at a temperature not exceeding 60°C (140°F). Then thoroughly break up the aggregation in such a manner as to avoid reducing the natural size of individual particles.

4.2. Sieve an adequate quantity of the representative pulverized soil over the 4.75-mm sieve. Discard the coarse material, if any, retained on the 4.75-mm sieve.

4.3. Select a representative sample, with a mass of approximately 3 kg (7 lb) or more, of the soil prepared as described in Sections 4.1 and 4.2.

Note 6—When developing a compaction curve for free draining soils, such as uniform sands and gravels, where seepage occurs at the bottom of the mold and base plate, taking a representative moisture content sample from the mixing bowl may be preferred in order to determine the amount of moisture available for compaction.

5. PROCEDURE

5.1. Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately four percentage points below optimum moisture content.

5.2. Form a specimen by compacting the prepared soil in the 101.60-mm (4-in.) mold (with collar attached) in five approximately equal layers to give a total compacted depth of about 125 mm (5 in.). Prior to compaction, place the loose soil into the mold and spread into a layer of uniform thickness. Lightly tamp the soil prior to compaction until it is not in a loose or fluffy state, using either the manual compaction rammer or similar device having a face diameter of approximately 50 mm (2 in.). Following compaction of each of the first four layers, any soil adjacent to the mold walls that has not been compacted or extends above the compacted surface shall be trimmed using a knife or other suitable device, and be evenly distributed on top of the layer. Compact each layer by 25 uniformly distributed blows from the rammer dropping free from a height of 457 mm (18 in.) above the elevation of the soil when a sleeve-type rammer is used, or from 457 mm (18 in.) above the approximate elevation of the soil as compacted by the previous blow when a stationary mounted type of rammer is used (Note 7).

Note 7—Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: A block of concrete, with a mass of not less than 90 kg (200 lb), supported by a relatively stable foundation; a sound concrete floor; and for field application, such surfaces as found in concrete box culverts, bridges, and pavements.

5.2.1. Following compaction, remove the extension collar, carefully trim the compacted soil even with the top of the mold by means of the straightedge, and determine the mass of the mold and moist soil in kilograms to the nearest five grams, or determine the mass in pounds to the nearest 0.01 pounds. For molds conforming to the tolerances given in Section 3.1.1 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 1060, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in Section 3.1.1 and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 30, and record the result as the wet density, \( W_1 \), in pounds per cubic foot of completed soil. For used molds out of tolerance by not more than 50 percent
(Section 3.1.3), use the factor for the mold as determined in accordance with T 19/T 19M. During compaction, the mold shall rest firmly on a dense, uniform, rigid, and stable foundation or base. This base shall remain stationary during the compaction process (Note 7).

5.3. Remove the material from the mold and slice vertically through the center. Take a representative sample of the material from one of the cut faces, determine the mass immediately and dry in accordance with T 265, Laboratory Determination of Moisture Content of Soils, to determine the moisture content, and record the results.

5.4. Thoroughly break up the remaining portion of the molded specimen until it will pass a 4.75-mm sieve as judged by eye, and add to the remaining portion of the sample being tested. Add water in sufficient amount to increase the moisture content of the soil by approximately one to two percentage points (water content increments should not exceed 2.5 percent, except when heavy clay soils or organic soils exhibiting flat elongated curves are encountered, the water content increments may be increased to a maximum of four percent), and repeat the above procedure for each increment of water added. Continue this series of determinations until there is either a decrease or no change in the wet unit mass per cubic meter (cubic foot) of the compacted soil (Note 8).

Note 8—In instances where the soil material is fragile in character and will reduce significantly in grain size due to repeated compaction, and in cases where the soil is heavy-textured, clayey material into which it is difficult to incorporate water, a separate and new sample shall be used in each compaction test. In these cases, separate samples shall be thoroughly mixed with amounts of water sufficient to cause the moisture contents of the samples to vary by approximately two percentage points. The moisture contents selected shall bracket the optimum moisture content, thus providing samples which, when compacted, will increase in mass to the maximum density and then decrease in mass. The samples of soil-water mixtures shall be placed in covered containers and allowed to stand for not less than 12 hours before making the moisture–density test.

5.4.1. In instances where the soil material is fragile in character and will be reduced significantly in grain size by repeated compaction, a separate and new sample shall be used in each compaction test.

METHOD B

6. SAMPLE

6.1. Select the representative sample in accordance with Section 4.3 except that it shall have a mass of approximately 7 kg (16 lb).

7. PROCEDURE

7.1. Follow the same procedure as described for Method A in Section 5, except for the following: Form a specimen by compacting the prepared soil in the 152.40-mm (6-in.) mold (with collar attached) in five approximately equal layers to give a total compacted depth of about 125 mm (5 in.), each layer being compacted by 56 uniformly distributed blows from the rammer. For molds conforming to tolerances given in Section 3.1.2 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold by 471, and record the result as the wet density, $W_1$, in kilograms per cubic meter, of the compacted soil. For molds conforming to tolerances in Section 3.1.2, and masses recorded in
pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 13.33 and record the result as the wet density, $W_1$, in pounds per cubic foot, of the compacted soil. For used molds out of tolerance by not more than 50 percent (Section 3.1.3), use the factor for the mold as determined in accordance with T 19/T 19M.

**METHOD C**

8. **SAMPLE**

8.1. If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus which is maintained at a temperature not exceeding 60°C (140°F). Then, thoroughly break up the aggregations in such a manner as to avoid reducing the natural size of individual particles.

8.2. Sieve an adequate quantity of the representative pulverized soil over the 19.0-mm sieve. Discard the coarse material, if any, retained on the 19.0-mm sieve (Note 9).

Note 9—The use of replacement method previously specified, where the oversized particles are replaced with finer particles, to maintain the same percentage of coarse material, is not considered appropriate to compute the maximum density.

8.3. Select a representative sample, with a mass of approximately 5 kg (12 lb), or more, of the soil prepared as described in Sections 8.1 and 8.2.

9. **PROCEDURE**

9.1. Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately four percentage points below optimum moisture content.

9.2. Form a specimen by compacting the prepared soil in the 101.60-mm (4-in.) mold (with collar attached) in five approximately equal layers to give a total compacted depth of about 125 mm (5 in.). Prior to compaction, place the loose soil into the mold and spread into a layer of uniform thickness. Lightly tamp the soil prior to compaction until it is not in a loose or fluffy state, using either the manual compaction rammer or similar device having a face diameter of approximately 50 mm (2 in.). Following compaction of each of the first four layers, any soil adjacent to the mold walls that has not been compacted or extends above the compacted surface shall be trimmed using a knife or other suitable device, and be evenly distributed on top of the layer. Compact each layer by 25 uniformly distributed blows from a rammer dropping free from a height of 457 mm (18 in.) above the elevation of the soil when a sleeve-type rammer is used, or from 457 mm (18 in.) above the approximate elevation of each finely compacted layer when a stationary mounted type of rammer is used. During compaction, the mold shall rest firmly on a dense, uniform, rigid, and stable foundation or base. This base shall remain stationary during the compaction process. (Note 7).

9.2.1. Following compaction, remove the extension collar, carefully trim the compacted soil even with the top of the mold by means of the straightedge. Fill any holes in the surface with unused or trimmed soil from the specimen, press in with the fingers, and again scrape the straight edge across the top of the mold. And Determine the mass of the mold and moist soil in kilograms to the nearest 5 grams, or determine the mass in pounds to the nearest 0.01 pounds. For molds conforming to the tolerances given in Section 3.1.1 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 1060, and record the result as the wet density, $W_1$, in kilograms per cubic meter, of compacted soil. For molds conforming to the tolerances given in Section 3.1.1 and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the
mold, by 30, and record the result as the wet density, \( W_1 \), in pounds per cubic foot, of compacted soil. For used molds out of tolerance by not more than 50 percent (Section 3.1.3), use the factor for the mold as determined in accordance with T 19/T 19M.

9.3. Remove the material from the mold and slice vertically through the center. Take a representative sample of the material from one of the cut faces, determine the mass immediately and dry in accordance with T 265, and record the results.

9.4. Thoroughly break up the remainder of the material until it will pass a 19.0-mm sieve and 90 percent of the soil aggregations will pass a 4.75-mm sieve as judged by eye, and add to the remaining portion of the sample being tested. Add water in sufficient amounts to increase the moisture content of the soil sample by one or two percentage points, and repeat the above procedure for each increment of water added. Continue this series of determinations until there is either a decrease or no change in the wet unit mass, \( W_1 \), per cubic meter (cubic foot) of the compacted soil (Note 8).

METHOD D

10. SAMPLE

10.1. Select the representative sample in accordance with Section 8.3, except that it shall have a mass of approximately 11 kg (25 lb).

11. PROCEDURE

11.1. Follow the same procedure as described for Method C in Section 9, except for the following: Form a specimen by compacting the prepared soil in the 152.40-mm (6-in.) mold (with collar attached) in five approximately equal layers to give a total compacted depth of about 127 mm (5 in.), each layer being compacted by 56 uniformly distributed blows from the rammer. For molds conforming to tolerances given in Section 3.1.2 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 471, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances in Section 3.1.2, and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 13.33, and record the result as the wet density, \( W_1 \), in pounds per cubic foot of the compacted soil. For used molds out of tolerance by not more than 50 percent (Section 3.1.3), use the factor for the mold as determined in accordance with T 19/T 19M.
CALCULATIONS AND REPORT

12. CALCULATIONS

12.1. Calculate the moisture content and the dry mass of soil as compacted for each trial as follows:

\[ w = \frac{A - B}{B - C} \times 100 \]  \hspace{2cm} (1)

\[ W = \frac{W_1}{w + 100} \times 100 \]  \hspace{2cm} (2)

where:

\( w \) = percentage of moisture in the specimen,

\( A \) = mass of the container and wet soil,

\( B \) = mass of the container and dry soil,

\( C \) = mass of the container,

\( W \) = dry density in kilograms per cubic meter or pounds per cubic foot of compacted soil, and

\( W_1 \) = wet density in kilograms per cubic meter or pounds per cubic foot of compacted soil.

13. MOISTURE–DENSITY RELATIONSHIP

13.1. The calculations in Section 12.1 shall be made to determine the moisture content and corresponding oven-dry density (unit mass) in kilograms per cubic meter or pounds per cubic foot for each of the compacted samples. The oven-dry densities of the soil shall be plotted as ordinates and corresponding moisture contents as abscissae.

13.2. Optimum Moisture Content—When the densities and corresponding moisture contents for the soil have been determined and plotted as indicated in Section 13.1, it will be found that by connecting the plotted points with a smooth line, a curve is produced. The moisture content corresponding to the peak of the curve shall be termed the “optimum moisture content” of the soil under the above compaction.

13.3. Maximum Density—The oven-dry density in kilograms per cubic meter or pounds per cubic foot of the soil at optimum moisture content shall be termed “maximum density” under the above compaction.

14. REPORT

14.1. The report shall include the following:

14.1.1. The method used (Method A, B, C, or D);

14.1.2. The optimum moisture content, as a percentage, to the nearest whole number;

14.1.3. The maximum density in kilograms per cubic meter to the nearest 10 kg/m³ or in pounds per cubic foot, to the nearest whole number;

14.1.4. In Methods C and D, whether the 19.0-mm material was removed or replaced; and

14.1.5. Type of face if other than 50.8-mm (2-in.) circular.
### Tester Qualification Practical Exam Checklist

**Moisture-Density Relations of Soils Using a 10-lb (4.54-kg) Rammer and a 18-in. (457-mm) Drop FOP for AASHTO T 180**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
</table>

#### Procedure Element

1. The tester has a copy of the current procedure on hand?  
   - Yes  
   - No

2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?  
   - Yes  
   - No

#### Sample Preparation

1. If damp, sample dried in air or drying apparatus, not exceeding 140°F (60°C)?  
   - Yes  
   - No

2. Sample pulverized and adequate amount sieved over the ¾ inch (19 mm) sieve?  
   - Yes  
   - No

3. Material retained on the sieve discarded?  
   - Yes  
   - No

4. Sample passing the sieve has appropriate mass?  
   - Yes  
   - No

#### Procedure

1. Sample mixed with water to approximately 4 percent below expected optimum moisture content?  
   - Yes  
   - No

2. Layer of soil placed in mold with collar attached?  
   - Yes  
   - No

3. Mold placed on rigid and stable foundation?  
   - Yes  
   - No

4. Lightly tamp soil in mold?  
   - Yes  
   - No

5. Soil compacted with 56 blows?  
   - Yes  
   - No

6. Scrape sides of mold and evenly distributed on top of the layer?  
   - Yes  
   - No

7. Soil placed and compacted in five equal layers?  
   - Yes  
   - No

8. No more than ½ inch of soil above the top of the bottom portion of the mold?  
   - Yes  
   - No

9. Collar removed and soil trimmed to top of mold with the straightedge?  
   - Yes  
   - No

10. Mass of mold and contents determined to appropriate precision?  
    - Yes  
    - No

11. Wet mass of specimen multiplied by appropriate factor to obtain wet density (.075 lbs/ft³)?  
    - Yes  
    - No

12. Soil removed from mold using sample extruder?  
    - Yes  
    - No

13. Soil sliced vertically through center?  
    - Yes  
    - No

14. Moisture sample removed from one cut face and moist mass determined immediately?  
    - Yes  
    - No
**Procedure**

<table>
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<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>15. Sample dried and water content determined according to AASHTO T 255 or T 265?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>16. Remainder of material from mold broken up to about passing sieve size and added to remainder of original test sample?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>17. Water added to increase moisture content in approximately 2 percent increments?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>18. Steps 2 through 15 repeated for each increment of water added?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>19. If soil is plastic (clay types):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Sample mixed with water varying moisture content by approximately 2 percent, bracketing the optimum moisture content?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>b. Samples placed in covered containers and allowed to stand for at least 12 hours?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>20. Process continued until wet density either decreases or stabilizes?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>21. Water content and dry density calculated for each sample?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>22. Dry density plotted on vertical axis, moisture content plotted on horizontal axis, and points connected with a smooth curve?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>23. Water content at peak of curve recorded as optimum water content and recorded to nearest 1 percent?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>24. Dry density at optimum water content reported as maximum density, to nearest 1 lb/ft³ (10 kg/m³)?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>25. All calculations performed correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt:  Pass ☐  Fail ☐  Second attempt:  Pass ☐  Fail ☐

Signature of Examiner __________________________________________

**Comments:**

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________
Performance Exam Checklist

Air Content of Concrete (Volumetric Method) for AASHTO T 196

Participant Name ___________________________________________ Exam Date __________

Procedure Element Yes No
1. Bowl filled in three equal layers? □ □
2. Each layer rodded 25 times? □ □
3. Bowl tapped (sharply) 10 to 15 times after rodding each layer? □ □
4. Excess concrete removed with strike-off bar? □ □
5. Flange of bowl wiped clean? □ □
6. Using funnel, water added until it appears in neck? □ □
7. Funnel removed & water adjusted to zero mark using rubber syringe? □ □
8. Screw cap is attached and tightened? □ □
9. Unit inverted and agitated until concrete is free from base? □ □
10. Unit rolled and rocked with neck elevated to remove air from concrete (approx. 1 min.)? □ □
11. Apparatus placed upright, jarred lightly and allowed to stand until air rises to the top? □ □
12. Agitation and rolling repeated until level of water column remains constant within 0.25%? □ □
13. One measureful of alcohol added with the syringe? □ □
14. Level of liquid read and recorded to nearest 0.25%? □ □

First attempt: Pass □ Fail □ Second attempt: Pass □ Fail □

Date: _____________ Tech. ID Code: _____________ Insp. ID Code: _____________

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________

________________________________________________________________________
1. SCOPE

1.1 This test method covers the determination of the theoretical maximum specific gravity and density of uncompacted hot-mix asphalt paving mixtures at 77°F (25°C).

Note 1—The precision of the method is best when the procedure is run on samples that contain aggregates that are completely coated. In order to assure complete coating it is desirable to run the method on samples that are close to the optimum asphalt content.

1.2 The values stated in English units are to be regarded as the standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:

- M 132, Terms Relating to Density and Specific Gravity of Solids, Liquids, and Gases
- R 10, Definition of Terms for Specifications and Procedures
- T 168, Sampling Bituminous Paving Mixtures

2.2 ASTM Standards:

- D 431, Practice for Determining Asphalt Volume Correction to a Base Temperature
- E 1, Specification for ASTM Thermometers

2.3 Other Standards:

- T 168 WSDOT FOP for WAQTC/AASHTO for Sampling Bituminous Paving Mixtures
- T 712 WSDOT Standard Method of Reducing Bituminous Paving Mixtures
- SOP 729 InPlace Density of Bituminous Mixes Using the Nuclear Moisture-Density Gauge FOP for WAQTC TM 8
- SOP 730 Standard Operating Procedure for Correlation of Nuclear Gauge Determined Density with Asphalt Concrete Pavement Cores
- SOP 731 Method for Determining Volumetric Properties of Asphalt Concrete Pavement Class Superpave
- SOP 732 Standard Operating Procedure for Superpave Volumetric Design for Hot-Mix Asphalt (HMA)
- SOP 733 Determination of Pavement Density Differentials Using the Nuclear Density Gauge
- SOP 734 Sampling Hot Mix Asphalt After Compaction (Obtaining Cores)
- SOP 735 Standard Operating Procedure for Longitudinal Joint Density

This FOP is based on WAQTC T 209 and has been modified per WSDOT standards.
To View the redline modifications, contact WSDOT Quality Systems Manager (360) 709-5497.
3. TERMINOLOGY

3.1 The terms specific gravity and density used in this test method are in accordance with M 132.

3.2 Definitions:

3.2.1 Density, as determined by this test method—the mass of a cubic meter of the material at 77°F (25°C) in English SI units, or the mass of a cubic foot of the material at 77°F (25°C) in inch-pound units.

3.2.2 Residual pressure, as employed by this test method—the pressure in a vacuum vessel when vacuum is applied.

3.2.3 Specific gravity, as determined by this test method—the ratio of a given mass of material at 77°F (25°C) to the mass of an equal volume of water at the same temperature.

4. SUMMARY OF TEST METHOD

4.1 A weighed sample of oven-dry HMA paving mixture in the loose condition is placed in a tared vacuum vessel. Sufficient water at a temperature of 77°F ± 0.9°F (25°C ± 0.5°C) is added to completely submerge the sample. Vacuum is applied for 15 ± 2 min to gradually reduce the residual pressure in the vacuum vessel to 30 mm ± 2.5 mm Hg (4.0 kPa ± 0.3 kPa). At the end of the vacuum period, the vacuum is gradually released. The volume of the sample of paving mixture is obtained either by (Section 9.5.1) immersing the vacuum container with sample into a water bath and weighing or by (Section 9.5.2) filling the vacuum container level full of water and weighing in air. At the time of weighing the temperature is measured as well as the mass. From the mass and volume measurements, the specific gravity or density at 77°F (25°C) is calculated. If the temperature employed is different from 77°F (25°C), an appropriate correction is applied.

5. SIGNIFICANCE AND USE

5.1 The theoretical maximum specific gravities and densities of hot-mix asphalt paving mixtures are intrinsic properties whose values are influenced by the composition of the mixtures in terms of types and amounts of aggregates and asphalt binder materials.

5.1.1 They are used to calculate values for percent air voids in compacted hot-mix asphalt paving mixtures.

5.1.2 They provide target values for the compaction of HMA paving mixtures.

5.1.3 They are essential when calculating the amount of asphalt binder absorbed by the internal porosity of the individual aggregate particles in a hot-mix asphalt paving mixture.

6. APPARATUS

6.1 Vacuum Container:

6.1.1 Six different vacuum containers are described. Each must be capable of withstanding the full vacuum applied, and each must be equipped with the fittings and other accessories required by the test procedure being employed. The opening in the container leading to the vacuum pump shall be covered by a piece of fine wire mesh such as No. 200 (75-μm) to minimize the loss of fine material.

6.1.2 The vacuum container size should be between 2000 and 10,000-mL and depends on the minimum sample size requirements given in Section 7.2. Avoid using a small sample in a large container.
6.3. Vacuum Bowl—Either a metal or plastic bowl with a diameter of approximately 7.1 to 10.2 in. (180 to 260 mm) and a bowl height of at least 6.3 in. (160 mm) shall be equipped with a transparent cover fitted with a rubber gasket and a connection for the vacuum line.

6.4. Vacuum Flask for Weighing in Air Only—A thick-walled volumetric glass flask and a rubber stopper with a connection for the vacuum line.

6.5. Pycnometer for Weighing in Air Only—A glass, metal or plastic pycnometer.

6.2. Balance, with ample capacity, and with sufficient sensitivity to enable the specific gravity of samples of uncompacted HMA paving mixtures to be calculated to at least four significant figures: that is, to at least three decimal places. For the bowl method (Type A), the balance shall be equipped with a suitable apparatus and holder to permit weighing the sample while suspended below the balance. The apparatus must have the same sensitivity, capacity and accuracy as the top pan.

6.2.1. Wire suspending the holder should be the smallest practical size to minimize any possible effects of a variable immersed length.

6.3. Vacuum pump or water aspirator, capable of evacuating air from the vacuum container to a residual pressure of 30 mm Hg (4.0 kPa) or less.

6.3.1. When a vacuum pump is used, a suitable trap of one or more 1000 mL filter flasks, or equivalent, shall be installed between the vacuum vessel and vacuum source to reduce the amount of water vapor entering the vacuum pump.

6.4. Residual Pressure Manometer, or vacuum gauge used for annual calibration and traceable to NIST (mandatory) to be connected directly to the vacuum vessel and to be capable of measuring residual pressure down to 30 mm Hg (4.0 kPa), or less (preferably to zero). It is to be connected at the end of the vacuum line using an appropriate tube and either a “T” connector on the top of the vessel or by using a separate opening (from the vacuum line) in the top of the vessel to attach the hose. To avoid damage, the manometer itself is not to be situated on top of the vessel but adjacent to it.

Note 2—A residual pressure of 30 mm Hg (4.0 kPa) absolute pressure is approximately equivalent to 730 mm Hg (97 kPa) reading on vacuum gauge at sea level.

Note 3—Residual pressure in the vacuum vessel, measured in millimeters of mercury, is the difference in the height of mercury in the Torricellian vacuum leg of the manometer and the height of mercury in the other leg of the manometer that is attached to the vacuum vessel.

6.5. Manometer or Vacuum Gauge, suitable for measuring the vacuum being applied at the source of the vacuum. This device can be connected directly to the vacuum source or be in the vacuum line close to the source. This is required to check the reading given by the residual pressure manometer attached directly to the vacuum vessel.

Note 4—The Torricellian vacuum leg of the manometer occasionally acquires one or more bubbles of air that introduce error into the residual pressure reading. By the addition of a vacuum gauge this error can often be quickly detected by the differences between two vacuum measurements.

6.6. Temperature measuring device Thermometers, calibrated liquid-in-glass thermometers of suitable range with subdivisions and maximum scale error of 0.2°F (0.1°C) 0.5°C (0.9°F), or any other thermometric device of equal accuracy, precision and sensitivity shall be used. Thermometers shall conform to the requirements of ASTM E 1.
6.7 Water Bath:

6.7.1. For vacuum bowls Type A or B containers, a water bath that can be maintained at a constant temperature between 68 and 86 ° F (20 and 30°C) is required. *(See Appendix.)* *(optional)*

6.8 Bleeder Valve, attached to the vacuum train to facilitate adjustment of the vacuum being applied to the vacuum vessel.

6.9 Protective Gloves, used when handling glass equipment under vacuum.

6.10 Mallet: With a rubber or rawhide head.

Note 5—An example of a correct arrangement of the testing equipment is shown in Figure 1. In the figure, the purpose of the train of small filter flasks is to trap water vapor from the vacuum vessel that otherwise would enter the oil in the vacuum pump and decrease the pump’s ability to provide high vacuum.

Figure 1—An example of the correct arrangement of testing apparatus

7. SAMPLING

7.1. Obtain the sample in accordance with WSQTC FOP for AASHTO T 168 and WSDOT T 712.

7.2. The size of the sample shall conform to the following requirements. Samples larger than the capacity of the container may be tested a portion at a time.

<table>
<thead>
<tr>
<th>Nominal Max. Agg. * Size</th>
<th>Class of Mix</th>
<th>Minimum Mass of Specimen, lbs (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Superpave</td>
<td>Other</td>
</tr>
<tr>
<td>No. 4 (4.75)</td>
<td></td>
<td>1 (500)</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td>3/8 In.</td>
<td>Class G &amp; D</td>
</tr>
<tr>
<td>½ (12.5)</td>
<td>1/2 In.</td>
<td>Class A, B, &amp; ATB</td>
</tr>
<tr>
<td>1 (25.0)</td>
<td>1 In.</td>
<td></td>
</tr>
<tr>
<td>1 ½ (37.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 (50.0)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.

Note: For a aggregate specification having a generally unrestrictive gradation (i.e., wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.
8. CALIBRATION OF FLASKS, BOWLS, AND PYCNOMETERS

This section has been deleted by WSDOT and replaced with the following:

The volumetric flask or metal vacuum pycnometer will be calibrated periodically in conformance with established verification procedures or per AASHTO T209. Calibration shall be done at 77 degrees F.

9. PROCEDURE

9.1. Separate the particles of the sample of paving mixture by hand, taking care to avoid fracturing the aggregate, so that the particles of the HMA fine aggregate portion are not larger than ¼ in (6.3 mm). If a sample of paving mixture is not sufficiently soft to be separated manually, place it in a flat pan, and warm it in an oven until it can be separated as described.

9.2. Samples prepared in a laboratory shall be cured and dried in an oven at 135 ± 5°C for a minimum of 2 hours, or as appropriate to match the mix design procedure being used. Longer drying time may be necessary for the sample to achieve a constant mass (mass repeats within 0.1 percent). Paving mixtures which have not been prepared in a laboratory with oven-dried aggregates shall be dried to a constant mass at a temperature of 105 ± 5°C. This drying and curing shall be combined with any warming described in Section 9.1.

Note 7—The minimum 2-hour time in the oven is specified as cure time for laboratory-prepared specimens. The curing at the specified temperature is especially important when absorptive aggregates are used. This will ensure the computation of realistic values for the amount of asphalt absorbed by the aggregate and void properties of the mix. Plant-produced materials should not be cured since absorption takes place during production.

9.3. Cool the sample to room temperature, and place it in a tared and calibrated flask, bowl, or pycnometer. The sample is to be placed directly into a Type A, B, C, D, or E vacuum container. A container within a container is not to be used. Weigh and designate the net mass of the sample as A. Add sufficient water at a temperature of approximately 77°F (25°C) to cover the sample completely.

9.4. Remove air trapped in the sample by applying gradually increased vacuum until the residual pressure vacuum gauge manometer reads 30 mm or less ± 2.5 mm Hg (3.7 ± 0.3 kPa). Maintain this residual pressure for 15 ± 2 min. Agitate the container and contents during the vacuum period either continuously by a mechanical device, or manually by vigorous shaking at intervals of about 2 minutes. Glass vessels should be shaken on a resilient surface such as a rubber or plastic mat, and not on a hard surface, so as to avoid excessive impact while under vacuum. To aid in releasing the trapped air from the metal vacuum pycnometer, tap the sides of the metal vacuum pycnometer 3 to 5 times with the mallet at approximately two minutes intervals.

Note 8—The release of entrapped air may be facilitated by the addition of a suitable wetting agent such as a few drops of Aerosol OT in concentration of 0.001 percent or 0.2 grams in 7.75 gal (20 L) of water. This solution is then diluted by about 20:1 to make a wetting agent of which 0.17 to 0.34 fl oz (5 to 10 mL) may be added to the apparatus.

9.5. At the end of the vacuum period, release the vacuum by increasing the pressure at a rate not to exceed 1.2 PSI (8 kPa) per second and proceed with one of the following determinations:

9.5.1. Weighing in Water—Suspend the container and contents in the water bath and determine the mass after 10 ± 1 min immersion. Measure the water bath temperature, and if different from 25 ± 1°C (77 ± 1.8°F), correct the mass to 25°C using the calibration temperature adjustment developed in Section 8.1. Designate the mass of the sample in water at 25°C as C.
Note 9—Instead of using a chart like Figure 2 to establish the mass correction for the temperature of the vacuum vessel submerged by itself in the water bath, this correction can be easily established by rapidly and completely emptying the vacuum container immediately following the final weighing, and then without delay, weighing the vessel by itself when totally submerged in the water bath.

9.5.2. Weighing in Air—Fill the flask (Type C), or any one of the pycnometers (Type D, E, or F) with water and adjust the contents to a temperature of 77 ± 1.8°F (25 ± 1°C) in a constant temperature water bath. Determine the mass of the container (and contents), completely filled, in accordance with Section 8.2 9 to 11 minutes minute after completing Section 9.4. Designate this mass as E.- Accurate filling may be ensured by the use of a glass cover plate.

In lieu of a constant temperature water bath described in 9.5.2, determine the temperature of the water within the flask or metal vacuum pycnometer and determine the appropriate density correction factor “R” using Table 2.

Note 10—See Appendix for correcting the theoretical maximum specific gravity when measurements are made at temperatures other than 25°C:

10. CALCULATION

10.1. Calculate the theoretical maximum specific gravity of the sample at 77°F (25°C) as follows:

10.1.1. Weighing in Water:

\[
\text{Theoretical Maximum Specific Gravity} = \frac{A}{A - C} \quad (2)
\]

where:

A = mass of oven-dry sample in air, g; and
C = mass of water displaced by sample at 25°C (77°F), g.

10.1.2. Weighing in Air:

\[
\text{Theoretical Maximum Specific Gravity} = \frac{A}{A + D - E} \quad (3)
\]

where:

A = mass of oven-dry sample in air, g;
D = mass of container filled with water at 77°F (25°C), g; and
E = mass of container filled with sample and water at 77°F (25°C), g.

10.1.3. Large-Size Plastic Pycnometer (Type F) Determinations:

10.1.3.1. If the test temperature is within +1.7 or –2.8°C (+3 or –5°F) of 25°C (77°F), that is, between 22.2 and 26.7°C (72 and 80°F), Equation 2.2 may be used to calculate specific gravity within 0.001 points or less error due to thermal effects.
10.1.3.2. If the test temperature differs significantly from 77°F (25°C), correct for thermal effects as follows:

WSDOT has removed the AASHTO calculation and replaced it with the following three calculation:

1. Determination using temperature correction:

\[
\text{Rice Sp. Gr.} = \frac{A}{A + D - E} \times R
\]

where:

\( A \) = mass of oven-dry sample in air, g;
\( D \) = mass of container filled with water at 77°F (25°C), g; and
\( E \) = mass of container filled with sample and water at 77°F (25°C), g.

\( R \) = Factor from Table 2 to correct density of water from the test temperature to 77°F (25°C).

Note: The flask calibration is done at 77 ± 0.4°F (25 ± 0.2°C).

2. Determination using weighted average:

\[
\text{Weighted Average Maximum Specific Gravity} = \frac{(\text{Sp. G}_1 \times A_1) + (\text{Sp. G}_2 \times A_2)}{(A_1 + A_2)}
\]

where:

\( \text{Sp. G}_1 \) = Specific gravity of first test segment
\( \text{Sp. G}_2 \) = Specific gravity of second test segment
\( A_1 \) and \( A_2 \) = Mass of dry sample in air of respective test segments

3. Calculate the rice density (calculate to one decimal place):

\[
\text{Rice density} = \text{Rice sp. gr. x 62.245 lb/ft}^3 \ (997 \text{ kg/m}^3)
\]
Table 2: Temperature Correction Factor

<table>
<thead>
<tr>
<th>C°</th>
<th>F°</th>
<th>&quot;R&quot;</th>
<th>C°</th>
<th>F°</th>
<th>&quot;R&quot;</th>
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<td>23.2</td>
<td>73.8</td>
<td>1.00045</td>
<td>35.0</td>
<td>95.0</td>
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<td>23.3</td>
<td>73.9</td>
<td>1.00042</td>
<td>35.6</td>
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<td>23.4</td>
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<td>36.1</td>
<td>97.0</td>
<td>0.99659</td>
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<td>23.6</td>
<td>74.5</td>
<td>1.00035</td>
<td>36.7</td>
<td>98.1</td>
<td>0.99638</td>
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<td>23.8</td>
<td>74.8</td>
<td>1.00030</td>
<td>37.2</td>
<td>99.0</td>
<td>0.99621</td>
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<td>23.9</td>
<td>75.0</td>
<td>1.00028</td>
<td>37.8</td>
<td>100.0</td>
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<td>24.0</td>
<td>75.2</td>
<td>1.00025</td>
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<td>24.2</td>
<td>75.6</td>
<td>1.00020</td>
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<td>24.4</td>
<td>75.9</td>
<td>1.00015</td>
<td>39.4</td>
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<td>24.6</td>
<td>76.3</td>
<td>1.00010</td>
<td>40.0</td>
<td>104.0</td>
<td>0.99516</td>
</tr>
</tbody>
</table>

Note: Water Temperatures should be maintained within the limits shown in bold face type.
10.2. Theoretical maximum density at 77°F (25°C):

10.2.1. Calculate the corresponding theoretical maximum density at 77°F (25°C) as follows:

Theoretical maximum density at 77°F (25°C) = theoretical maximum specific gravity × 997.1 kg/m³ in SI units, or

Theoretical maximum density at 77°F (25°C) = theoretical maximum specific gravity × 62.245 lb/ft³ in inch-pound units.

where:

The specific gravity of water at 77°F (25°C) = 997.1 in SI units or = 62.245 in inch-pound units.

11. SUPPLEMENTAL PROCEDURE FOR MIXTURES CONTAINING POROUS AGGREGATE

WSDOT has removed this section.

12. REPORT

12.1. Report the following information:

12.1.1. Specific gravity and density of the mixture to the third decimal place as: sp gr 25/25°C or density at 77°F (25°C),

12.1.2. Type of mixture,

12.1.3. Size of sample,

12.1.4. Number of samples,

12.1.5. Type of container, and

12.1.6. Type of procedure.

13. PRECISION

See AASHTO T-209 for Precision.

APPENDIX

Nonmandatory Information

A1. THEORETICAL MAXIMUM SPECIFIC GRAVITY FOR A LOOSE-PAVING MIXTURE

WSDOT has removed this section.

1 Sargent Welch, 39745 Gauge-Vacuum, Mercury Prefilled (or equivalent).
## Performance Exam Checklist

### Theoretical Maximum Specific Gravity and Density of HOT MIX ASPHALT Paving Mixtures

**FOP for AASHTO T 209**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
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<tbody>
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</table>

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Particles of sample separated?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Care used not to fracture mineral fragments?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. After separation, fine HMA particles not larger than ¼ inch?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Sample at room temperature?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Mass of bowl or flask determined?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Mass of sample and bowl or flask determined?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Mass of sample determined?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Water at approximately 77°F (25°C) added to cover sample?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Entrapped air removed using partial vacuum for 15 ± 2 min?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12. Container and contents agitated continuously by mechanical device or manually by vigorous shaking at intervals of about 2 minutes?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. For metal pycnometer, strike 3 to 5 times with a mallet?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14. Release of entrapped air facilitated by addition of suitable wetting agent (optional)?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15. Flask determination:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Flask filled with water?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b. Flask then placed in constant temperature water bath (optional)?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c. Contents at 77 ± 1° F or Table 2 in FOP used?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>d. Mass of filled flask determined 9 to 11 minutes after removal of entrapped air completed?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16. All calculations performed correctly?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt:  Pass [ ]  Fail [ ]

Second attempt:  Pass [ ]  Fail [ ]

Signature of Examiner ________________________________
WSDOT FOP for AASHTO T 217

Determination of Moisture in Soils by Means of a Calcium Carbide Gas Pressure Moisture Tester

1. SCOPE

1.1 This method of test is intended to determine the moisture content of soils by means of a calcium carbide gas pressure moisture tester. The manufacturer’s instructions shall be followed for the proper use of the equipment.

1.2 The following applies to all specified limits in this standard: For the purposes of determining conformance with these specifications, an observed value or a calculated value shall be rounded off “to the nearest unit” in the last right-hand place of figures used in expressing the limiting value, in accordance with R 11, Recommended Practice for Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values.

Note 1: This method shall not be used on granular materials having particles large enough to affect the accuracy of the test in general any appreciable amount retained on a No. 4 (4.75-mm) sieve. The super 200 D tester is intended to be used to test aggregate.

1.3 The values stated in English units are to be regarded as the standard.

1.4 Refer to R 16 for regulatory information for chemicals.

2. REFERENCED DOCUMENT

2.1 AASHTO Standards:

R 11, Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values

T 265, Laboratory Determination of Moisture Content of Soils

3. APPARATUS

3.1 Calcium carbide pressure moisture test – a chamber with attached pressure gage for the water content of specimens having a mass of at least 20 g. (Figure 1).

Those “Speed Moisture Testers” which use a 20 g sample may be used to test aggregates and soil-aggregate mixtures where the maximum particle size is 3/4 in. (20 mm) or less.

3.2 Balance – shall conform to AASHTO M 231, Class G-2.

3.3 Two 1.25-in. (31.75-mm) steel balls

3.4 Cleaning brush and cloth.

3.5 Scoop for measuring calcium carbide reagent.

This FOP is based on AASHTO T 217-02
4. MATERIAL

4.1 Calcium carbide reagent.

**Note 2:** The calcium carbide must be finely pulverized and should be of a grade capable of producing acetylene gas in the amount of at least 2.25 ft³/lb (0.14 m³/kg) of carbide.

**Note 3:** The “shelf life” of the calcium carbide reagent is limited, so it should be used according to the manufacturer’s recommendations. When a can of calcium carbide is opened, it shall be dated. After 3 months of use, or if the can becomes contaminated, it shall be discarded.

5. PROCEDURE

5.1 When using the 20-g or 26-g tester, place three scoops (approximately 24 g) of calcium carbide in the body of the moisture tester (or per the manufacturers recommendations). When using the super 200 D tester to test aggregate, place six scoops (approximately 48 g) of calcium carbide in the body of the moisture tester.

**Note 4:** Care must be exercised to prevent the calcium carbide from coming into direct contact with water.

5.2 Weigh a sample of the exact mass specified by the manufacturer of the instrument in the balance provided, and place the sample in the cap of the tester. When using the 20-g or 26-g size tester, place two 1.25-in. (31.75-mm) steel balls in the body of the tester with the calcium carbide (or per the manufacturers recommendations).

**Note 5** Manufacturer’s instructions shall be followed for the use of steel balls, particularly when testing sand.

**Note 6** If the moisture content of the sample exceeds the limit of the pressure gage (12 percent moisture for aggregate tester to 20-percent moisture for soil tester), a one-half size sample must be used and the dial reading must be multiplied by 2. This proportional method is not directly applicable to the dry mass percent scale on the super 200 D tester.

5.3 With the pressure vessel in an approximately horizontal position, insert the cap in the pressure vessel and seal the unit by tightening the clamp, taking care that no carbide comes in contact with the soil until a complete seal is achieved.
5.4 Raise the moisture tester to a vertical position so that the soil in the cap will fall into the pressure vessel.

5.5 Shake the instrument vigorously so that all lumps will be broken up to permit the calcium carbide to react with all available free moisture. When steel balls are being used in the tester and when using the large tester to test aggregate, the instrument should be shaken with a rotating motion so the steel balls or aggregate will not damage the instrument or cause soil particles to become embedded in the orifice leading to the pressure diaphragm.

**Note 7:** Shaking should continue for at least 60 seconds with granular soils and for up to 180 seconds for other soils so as to permit complete reaction between the calcium carbide and the free moisture. Time should be permitted to allow dissipation of the heat generated by the chemical reaction.

5.6 When the needle stops moving, read the dial while holding the instrument in a horizontal position at eye level.

5.7 Record the sample mass and the dial reading.

5.8 With the cap of the instrument pointed away from the operator, and away from open flame or source of ignition, slowly release the gas pressure. Empty the pressure vessel and examine the material for lumps. If the sample is not completely pulverized, the test should be repeated using a new sample. Clean the cap thoroughly of all carbide and soil before running another test.

**Note 8:** When removing the cap, care should be taken to point the instrument away from the operator to avoid breathing the fumes, and away from any potential source of ignition for the acetylene gas.

5.9 The dial reading is the percent of moisture by wet mass and must be converted to dry mass. With the super 200 D tester the dial reading is the percent of moisture by dry mass, and no further calculation is required.

6. **CALCULATION**

6.1 The percentage of moisture by dry mass of the soil may be determined from a correction curve similar to Figure 2.

6.2 A correction curve similar to Figure 2 is normally supplied with each moisture tester. Each moisture tester, however, should be checked for the accuracy of its gage, and for the accuracy of its correction curve.

5.2.1 The accuracy of the moisture tester gage should be checked by using a calibration kit (available from the manufacturer), equipped with a standard gage. In case of discrepancy, the gage on the tester should be adjusted to conform with the standard gage.

5.2.2 The accuracy of the correction curve should be checked by comparing curve-corrected moisture contents to moisture contents of locally prepared soils determined using T 265. In case of discrepancy, develop a new correction curve based on moisture contents determined from T 265.
5.2.3 The range of the factory-supplied or laboratory-determined curves may be extended by additional testing.

Figure 2 — Correction Curve for Moisture Tester Reading

Note 9: It may be more convenient for field use of the apparatus to prepare a table of moisture tester readings versus oven-dry moisture content for the moisture tester.

6.3 Determine the percentage of moisture to the nearest whole percent.
Performance Exam Checklist

**Determination of Moisture in Soils by Means of Calcium Carbide Gas Pressure Moisture Tester**

**FOP for AASHTO T 217**

<table>
<thead>
<tr>
<th>Procedure Element</th>
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<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>and if required, has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Shelf life of calcium carbide reagent checked?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Correct amount of reagent placed in body of tester?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Number and size of steel balls correct?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Correct mass of moist soil placed in cap of tester?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Cap clamped to body with tester in horizontal position?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Shaking done for proper time (60 seconds for granular soils,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>180 seconds for other soils)?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Shaking done without steel balls hitting cap or bottom of tester?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Reading taken with tester in horizontal position at eye level?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Reading taken after gauge stops moving?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12. Gauge reading recorded?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. Tester positioned with cap away from user and away from open flame or</td>
<td></td>
<td></td>
</tr>
<tr>
<td>source of ignition before gas slowly released?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14. Moisture content on wet mass basis converted to dry mass basis?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner __________________________________________

Comments:

__________________________________________________________________

__________________________________________________________________

__________________________________________________________________
This FOP is based on AASHTO T 224-0

Correction for Coarse Particles in the Soil
Compaction Test

1. SCOPE

1.1. This method describes a procedure for adjusting the densities of soil and soil aggregate mixtures to compensate for differing percentages of coarse particles retained on either the No.4 (4.75-mm) or 3/4 in. (19.0-mm) sieve. This is necessary to adjust either the field wet density to a dry density of the material passing these sieves or the reverse, by adjusting the lab density to the field density when doing compaction control testing. Comparisons are made by comparing the field densities with the maximum dry density as determined by T 99 or T 180.

The standard for the WSDOT is to use:

- materials with less than 30 percent by weight retained on the U.S. No. 4 sieve shall be determined using FOP for AASHTO T 99 Method A.
- materials with 30 percent or more by weight retained on the U.S. No. 4 sieve and less than 30 percent retained on the 3/4 inch sieve shall be determined by WSDOT Test Method No. 606 or FOP for AASHTO T 180 Method D.

The Standard Specifications define non-granular material as 30% or less retained on the No. 4 (4.75-mm) sieve.

1.2. Two methods are available for correction, either, lab to field or field to lab density. The method specified in Section 4.1 adjusts the compacted lab density to the field density. The method specified in Section 4.2 adjust the field wet density to dry density of the fine fraction and compares its results with the compacted lab density.

WSDOT uses the calculation specified in section 4.1 for adjusting the compacted lab density to the field density. Appendix B contains a nomograph for this adjustment.

1.3. This test method applies to soil mixtures that have 30 percent or less retained on the No. 4 (4.75-mm) sieve, when Method A or B of T 99 or T 180 is used, or mixtures that have 30 percent or less retained on the 3/4 in. (19.0 mm) sieve, when Method C or D of T 99 or T 180 is used. The material retained on these sieves shall be defined as oversize particles (coarse particles).

1.4. This method applies to soils with any percentage of oversize particles as specified in Section 1.3. However, the correction may not be of practical significance for soils with only a small percentage of oversize particles. The person or agency specifying this method shall specify a minimum percentage of oversize particles below which the method need not be applied. If a minimum percentage is not specified, this method shall be applied to samples with more than 5 percent by weight of oversize particles.

1.5. The following applies to all specified limits in this standard: For the purposes of determining conformance with these specifications, an observed value or a calculated value shall be rounded off to the nearest 1 pcf (10 kg/m³), according to R 11.

1.6. The values stated in SI units are to be regarded as the standard.

1This FOP is based on AASHTO T 224-01
2. REFERENCE DOCUMENTS

2.1. AASHTO Standards:

- R 11, Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values
- T 85, Specific Gravity and Absorption of Coarse Aggregate
- T 99, Moisture-Density Relations of Soils Using a 5.5-lb, (2.5-kg) Rammer and a 12-in. (305-mm) Drop
- T 180, Moisture-Density Relations of Soils Using a 10-lb (4.54-kg) Rammer and a 18-in. (457-mm) Drop
- T 217, Determination of Moisture in Soils by Means of a Calcium Carbide Gas Pressure Moisture Tester
- T 255, Total Evaporable Moisture Content of Aggregate by Drying
- T 272, Family of Curves-One Point Method

Other Methods
- WSDOT SOP 615, Determination of the % Compaction for Embankments & Untreated Surfacing Materials using the Nuclear Moisture-Density Gauge

3. OUTLINE OF METHOD

3.1. When Method A or Method B of WSDOT FOP for AASHTO T 99 or WSDOT FOP for AASHTO T 180 is employed, the total field wet density is compared with the dry density of the soil particles passing the No. 4 (4.75-mm) sieve.

3.2. When Method C or Method D of WSDOT FOP for AASHTO T 99 or WSDOT FOP for AASHTO T 180 is employed, the total field wet density is compared with the dry density of the soil particles passing the 3/4 in. (19.0-mm) sieve.

3.3. Significant figures are as follows:

3.3.1. Adjusted wet density of the fine material passing the No.4 (4.75-mm) sieve, Methods A and B; or 3/4 in. (19.0-mm) sieve, Method C and D; \( D_f \) 0.1 pcf (1 kg/m\(^3\)).

3.3.2. Bulk specific gravity of the coarse material on the 4.75-mm (No. 4) sieve, Methods A and B; or 3/4 in. (19.0-mm) sieve, Methods C & D; \( G_m \) 0.01.

3.3.3. Percent by mass, of coarse and fine particles, of material retained and passing the No. 4 (4.75-mm) sieve, Methods A & B; or 3/4 in. (19.0-mm) sieve, Methods C and D; \( P_c \) and \( P_f \) 0.1 percent.

3.3.4. In-place (field) wet density of the total sample \( D \) 0.1 pcf (1 kg/m\(^3\)).

4. ADJUSTMENT EQUATION

4.1. Compacted Laboratory Dry Density Corrected to Field Dry Density

4.1.1. This Section corrects the laboratory density obtained by either WSDOT FOP for AASHTO T 99 or WSDOT FOP for AASHTO T 180 for the moisture content and density of the material retained on the No. 4 (4.75-mm) sieve, Methods A & B; or the material retained on the 3/4 in. (19.0-mm) sieve, Methods C and D. The maximum lab dry density, adjusted for oversized particles and total moisture content are compared
with the field dry density and field moisture content. This method is limited to field samples containing 30 percent or less for material retained on the No. 4 (4.75-mm) sieve, Methods A and B; or 30 percent or less of material retained on the 3/4 in. (19.0-mm) sieve, Methods C and D.

4.1.2. Determine the moisture content of the fine particles and oversize particles of the material used during compaction. The moisture contents can be determined by WSDOT FOP for AASHTO T 217 or T 255. The moisture content of the oversize material retained on the sieve can be assumed to be two (2) percent for most construction applications. If the moisture content of the oversized material is generally known, substitute that moisture content in the calculations. It is recommended if drying equipment is available, determine the actual moisture contents.

4.1.3. Calculate the dry mass of the coarse and fine particles as follows:

\[
MD = \frac{MM}{1 + MC}
\]

where:

- \(MD\) = mass of dry material (fine or oversize particles);
- \(MM\) = mass of moist material (fine or oversize particles);
- \(MC\) = moisture content of respective fine or oversized particles, expressed as a decimal.

4.1.4. Calculate the percentage of the fine particles and oversized particles by dry weight of the total sample as follows:

\[
P_f = \frac{100 \times M_{DF}}{M_{DF} + M_{DC}}
\]

and

\[
P_C = \frac{100 \times M_{DC}}{M_{DF} + M_{DC}}
\]

where:

- \(P_f\) = percent of fine particles, of sieve used, by weight;
- \(P_C\) = percent of oversize particles, of sieve used, by weight;
- \(M_{DF}\) = mass of dry particles, and
- \(M_{DC}\) = mass of oversize particles.

4.1.5. Calculate the corrected optimum moisture content and corrected dry density of the total sample as follows:

**Optimum Moisture Content:**

\[
MC_T = MC_f \times \frac{P_C}{100}
\]

where:

- \(MC_T\) = corrected optimum moisture content
- \(MC_f\) = moisture content from the maximum density curve
- \(P_C\) = percent passing the US No. 4 sieve for T99, or passing the ¾ sieve for T180
Density:

\[
D_d = \frac{100(D_f \cdot k)}{[(D_f) (P_C) + (k)(P_f)]}
\]

where:

- \( D_d \) = corrected total dry density (combined fine and oversized particles) pcf (kg/m\(^3\)),
- \( D_f \) = Laboratory Maximum Density (T99 or T180) dry density of the fine particles pcf (kg/m\(^3\)),
- \( P_C \) = percent of oversize particles, of sieve used, by weight,
- \( P_f \) = percent of fine particles, (US No. 4 - for T99, or ¾ - for T180) of sieve used, by weight,
- \( k \) = 62.4 * Bulk Specific Gravity (\( G_m \)) (oven dry basis) of coarse particles (pcf),

**Note 1** – If the bulk specific gravity has been determined, this value may be used in the calculations. Determine the Bulk Specific Gravity according to T 85, or determine the bulk specific gravity for the coarse aggregate according the WSDOT Test Method T-606. For most construction activities bulk the specific gravity can be assumed to be 2.60-2.67

4.2 Field Wet Density Corrected to Compacted Laboratory Density

WSDOT has Deleted this section

5. PRECISION

5.1. Since this correction for coarse particles involves no testing but instead utilizes the results of other tests and mathematically combines the results, determination of the precision and accuracy is not applicable.

APPENDIX A

A1. NOTES

A1.1. These methods, described for coarse particle correction, are applicable to one type of soil and soil aggregate material only. If the characteristic of the material changes, then a moisture density relationship (T 99 or T 180) test is performed to determine a new maximum density.

A1.2. T 272 describes the methods for determining different maximum densities of soil and soil-aggregate materials which reveal certain similarities and trends characteristic of the material type and source.

A1.2.1. Utilization of a Family of Curves-When using the One Point Method (T 272), Note 3 of T 272 does not apply. The percentage of oversized particles, when performing the density of soil and soil aggregate in-place, must be determined to adjust the T 99 or T 180 maximum density to compensate for this percentage.
APPENDIX B

WSDOT has added the following nomograph and prefers its use.

B1. SOLUTION

B.1.1. The solution to this equation by nomograph is shown in Figure 1, wherein the maximum density of material passing the No. 4 (4.75 mm) sieve (Section 3.1) determined in the laboratory test is plotted and a line is drawn from this point to a point on the line representing the bulk specific gravity of the coarse particles. When the percentage of coarse particles contained in a field density of soil and soil-aggregate in-place has been determined and plotted on the above line as the abscissa on the chart (Figure 1), the ordinate will give the maximum laboratory dry density corrected to the new percentage of coarse particles.

B.1.2. Figures 2 and 3 illustrate a solution when Method C or Method D of WSDOT FOP for AASHTO T 99 or T180 is utilized (Section 3.2).

B2. Report

Report the maximum density on DOT Form 350-074 and DOT Form 351-015.
EXAMPLE
METHOD A OR METHOD B

Maximum laboratory dry density of 4.75 mm minus material, $D_L = 1826 \text{ kg/m}^3 (114.0 \text{ lb/ft}^3)$. Plot at A.\n
Specific gravity of coarse particles (4.75 mm plus) = 2.50. Plot at B.\n
Percent of coarse particles (4.75 mm plus) found when performing the density of soil and soil-aggregate in-place = 29.0. Plot at C.\n
Draw line AB.\n
Locate intersection of line extended vertically from C to line AB (point E).\n
Draw line horizontally from E, intersecting the ordinate at F.\n
Point F = 1949 kg/m$^3$ (121.7 lb/ft$^3$), the corrected maximum dry density of total material, D.

---

Figure 1. Density Correction Chart for Coarse Particles

EXAMPLE
METHOD C OR METHOD D

Maximum laboratory dry density of 19.0 mm minus material, $D_L = 1220 \text{ lb/ft}^3$. Plot at A.\n
Percent of coarse particles (4.75 mm plus), including any coarse particles replaced in the T99 or T180 Moisture Density Test, = 30.0. Plot at B.\n
Draw a line horizontally from A and a line vertically from B, meeting at point C.\n
Specific gravity of coarse particles (4.75 mm plus) = 2.50. Plot at E. Draw a straight line, EC, and extend toward ordinate.

(ADJUSTING MAXIMUM DENSITY WITH LESS THAN 30 PERCENT COARSE PARTICLES)

Percent of coarse particles (4.75 mm plus) found when performing the density of soil and soil-aggregate in-place = 15.0. Plot at F and draw line vertically from F to intersection with EC extended (point G).\n
Draw line horizontally from G, intersecting ordinate at H. Point H = 1890 kg/m$^3$ (118.0 lb/ft$^3$), corrected maximum dry density of total material, D.

(ADJUSTING MAXIMUM DENSITY WITH MORE THAN 30 PERCENT COARSE PARTICLES)

Percent of coarse particles (4.75 mm plus) found when performing the density of soil and soil-aggregate in-place = 45.0. Plot at J and draw line vertically from J to intersection with EC (point K).\n
Draw line horizontally from K, intersecting the ordinate at L. Point L = 2018 kg/m$^3$ (126.0 lb/ft$^3$), the corrected maximum dry density of total material, D.

---

Figure 2. Density Correction Chart for Coarse Particles
Maximum laboratory dry density of 19.0 mm minus, $D_1 = 1938 \text{ kg/m}^3$ (121.0 lb/ft$^3$). Plot at A.
Specific gravity of coarse particles (19.0 mm plus) = 2.50. Plot at B.
Percent of coarse particles (19.0 mm plus) found when performing the density of soil and soil-aggregate in-place = 25.0. Plot at C.

Draw line AB.
Locate intersection of line extended vertically from C to line AB (point E).
Draw line horizontally from E, intersecting the ordinate at F.
Point F = 2015 kg/m$^3$ (125.8 lb/ft$^3$), the corrected maximum dry density of total material, D.

Figure 3. Density Correction Chart for Coarse Particles
If the specific gravity is unknown, then use 2.67.
Performance Exam Checklist

WSDOT FOP for AASHTO T 224

Correction for Coarse Particles in the Soil Compaction Test

Participant Name _______________________________ Exam Date ____________________

Procedure Element

1. The tester has a copy of the current procedure on hand? ☐ ☐
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? ☐ ☐

Gradation Analysis

1. WSDOT SOP 615 used to identify percent of oversize material? ☐ ☐
2. Sample Dried to a SSD condition (dried until no visible surface moisture present) and mass recorded? ☐ ☐
3. Sample allowed to cool sufficiently prior to sieving? ☐ ☐
4. Sample was shaken by hand through a No. 4 sieve for a sufficient period of time? ☐ ☐
5. Recorded mass of material retained on No. 4 sieve? ☐ ☐
6. Calculated and recorded percent of material retained and passing No 4 sieve? ☐ ☐

Correction for Coarse Particles

7. Maximum density of material passing No. 4 sieve, as determined by AASHTO T-99, correctly plotted onto nomograph? ☐ ☐
8. Line correctly drawn from maximum density plot to the correct specific gravity? ☐ ☐
9. Percent of material retained on the No. 4 screen correctly plotted onto nomograph? ☐ ☐
10. Corrected maximum density correctly identified from the nomograph? ☐ ☐
11. All calculations performed correctly? ☐ ☐

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________
WSDOT FOP for AASHTO T 231¹

Capping Cylindrical Concrete Specimens

1. SCOPE

1.1 This method covers apparatus, materials, and procedures for capping freshly molded concrete cylinders with neat cement and hardened cylinders and drilled concrete cores with high-strength gypsum plaster or sulfur mortar.

1.2 The values stated in English units are the preferred standard.

1.3 This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety problems 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. For specific precaution statements, see Sections 4.3 and 6.2.3.1.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:
   M 85 Portland Cement
   T 22 Compressive Strength of Cylindrical Concrete Specimens
   T 106 Compressive Strength of Hydraulic Cement Mortar (Using 2-in. (50-mm) Cube Specimens)
   M 240 Blended Hydraulic Cement

2.2 ASTM Standards:
   C 287 Specification for Chemical-Resistant Sulfur Mortar
   C 472 Physical Testing of Gypsum, Gypsum Plasters and Gypsum Concrete

2.3 ANSI Standards:
   B 46.1 Standard for Surface Texture

3. SIGNIFICANCE AND USE

3.1 This practice describes procedures for providing plane surfaces on the end surfaces of freshly molded concrete cylinders, hardened cylinders, or drilled concrete cores when the end surfaces do not conform with the planeness and perpendicularity requirements of applicable standards.

4. CAPPING EQUIPMENT

4.1 Capping Plates — Neat cement caps and high-strength gypsum-plaster caps shall be formed against a glass plate of at least \( \frac{1}{4} \) in. (6 mm) thick, a machined metal plate at least 0.45 in. (11 mm) thick (Note 1), on a polished plate of granite or diabase at least 3 in. (75 mm) thick. Sulfur mortar caps shall be formed against similar metal or stone plates. In all cases, plates shall be at least 1 in. (25 mm) greater in diameter than the test specimen and the working surfaces shall not depart from a plane by more than 0.002 in. (0.05 mm) in 6 in. (150 mm).

¹This Test Method is based on AASHTO T 231-05.
The surface roughness of newly finished metal plates shall not exceed that set forth in Table 4 of the American National Standard for Surface Texture (ANSI B46.1) or 125 μ in. (0.003 mm) for any type of surface and direction of lay. The surface when new shall be free of gouges, grooves, or indentations beyond those caused by the finishing operation. Metal plates that have been in use shall be free of gouges, grooves, or indentations greater than 0.010 in. (0.25 mm) deep or greater than 0.05 in.² (32 mm²) in surface area. If a recess is machined into the metal plate, the thickness of the plate beneath the recessed area shall be at least 1/2 in. (13 mm). In no case shall the recess in the plate be deeper than 1/2 in. (13 mm).

**Note 1:** In vertical capping devices, use of two-piece metal capping plates is advantageous as this facilitates refinishing of the capping surface should it become necessary to do so. In such devices, the lower section is a solid plate and the upper section has a circular hole which forms the recess. The two sections are customarily fastened together with machine screws. It is advantageous to have the upper surface of the lower plate case hardened. A Rockwell hardness of HRC 48 is suggested.

4.2 Alignment Devices — Suitable alignment devices such as guide bars or bull’s-eye levels shall be used in conjunction with capping plates to ensure that no single cap will depart from perpendicularity of cylindrical specimen by more than 0.5° (approximately equivalent to 1/8 in. in 12 in. (3.2 mm in 305 mm)). The same requirement is applicable to the relationship between the axis of the alignment device and the surface of a capping plate when guide bars are used. In addition, the location of each bar with respect to its plate must be such that no cap will be off-centered on a test specimen by more than 1/16 in. (1.6 mm).

4.3 Melting Pots for Sulfur Mortars — Pots used for melting sulfur mortars shall be equipped with automatic temperature controls and shall be made of metal or lined with a material that is nonreactive with molten sulfur.

**Caution:** Melting pots equipped with peripheral heating will ensure against accidents during reheating of cooled sulfur mixtures which have a crusted-over surface. When using melting pots not so equipped, a build up of pressure under the hardened surface crust on subsequent reheating may be avoided by use of a metal rod which contacts the bottom of the pot and projects above the surface of the fluid sulfur mix as it cools. The rod should be of sufficient size to conduct enough heat to the top of reheating to melt a ring around the rod first and thus avoid the development of pressure. A large metal ladle can be substituted for the rod.

Sulfur melting pots should be used under a hood to exhaust the fumes to outdoors. Heating over an open flame is dangerous because the flash point of sulfur is approximately 440°F (227°C) and the mixture can ignite due to overheating. Should the mixture start to burn, covering will snuff out the flame. The pot should be recharged with fresh material after the flame has been extinguished.
5. CAPPING MATERIALS

5.1 The strength of the capping material and the thickness of the caps shall conform to the requirements of Table 1.

Table 1—Compressive Strength and Maximum Thickness of Capping Materials

<table>
<thead>
<tr>
<th>Cylinder Compressive Strength M Pa (psi)</th>
<th>Minimum Strength of Capping Material</th>
<th>Maximum Average Thickness of Cap</th>
<th>Maximum Thickness Any Part of Cap</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5 to 50 M Pa (500 to 7000 psi)</td>
<td>35 M Pa (5000 psi) or cylinder strength, whichever is greater</td>
<td>6 mm (0.25 in.)</td>
<td>8 mm (0.31 in.)</td>
</tr>
<tr>
<td>greater than 50 M Pa (7000 psi)</td>
<td>Compressive strength not less than cylinder strength, except as provided in Section 5.1.1.</td>
<td>3 mm (0.125 in.)</td>
<td>5 mm (0.20 in.)</td>
</tr>
</tbody>
</table>

5.1.1 If sulfur mortar, high strength gypsum plaster and other materials except neat cement paste are to be used to test concrete with a strength greater than 50 M Pa (7000 psi), the manufacturer or the user of the material must provide documentation:

• that the average strength of 15 cylinders capped with the materials is not less than 98 percent of the average strength of 15 companion cylinders capped with neat cement paste or 15 cylinders ground plane to within 0.05 mm (0.002 in.),
• that the standard deviation of the strengths of the capped cylinders is not greater than 1.57 times that of the standard deviation of the reference cylinders,
• that the cap thickness requirements were met in the qualification tests, and
• of the hardening time of the caps used in the qualification tests.

5.1.2 Additionally, the qualification test report must include the compressive strength of 50-mm (2-in.) cubes of the material qualified and of neat cement paste cubes, if used. Capping materials conforming to these requirements are permitted to be used for cylinders with strengths up to 20 percent greater than the concrete tested in these qualification tests. The manufacturer must requalify lots of material manufactured on an annual basis or whenever there is a change in the formulation of the raw materials. The user of the material must retain a copy of the qualification results, and the dates of manufacture of material qualified and of the material currently being used. See Table 2.

Table 2—Sample Report of Qualifications of a Capping Material

Note: Manufacturer: Testing Supplies Co.
Capping Material: Super Strong AAA-Sulfur mortar
Lot: 12a45    Date Tested: 11/3/98
Signed by: ____________________________________________
(Testing Agency and Responsible Official)
<table>
<thead>
<tr>
<th>Item</th>
<th>Capping Material</th>
<th>Control Cylinders</th>
<th>Ratio</th>
<th>Criteria</th>
<th>Pass/Fail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concrete Cylinder Test Data</td>
<td>Sulfur</td>
<td>Ground</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Type of Capping Material</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average Concrete Strength,M Pa (psi)</td>
<td>76.2 (11,061)</td>
<td>75.9 (11,008)</td>
<td>1.005</td>
<td>&gt;0.98 Xc ≤1.57 C</td>
<td>Pass</td>
</tr>
<tr>
<td>Standard Deviation M Pa (psi)</td>
<td>2.59 (376)</td>
<td>1.72 (250)</td>
<td>1.504</td>
<td></td>
<td>Pass</td>
</tr>
<tr>
<td>Number of cylinders tested</td>
<td>15</td>
<td>15</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cap age when cylinders tested</td>
<td>7 days</td>
<td>n/a</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Capping Material Test Data</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average cap thickness mm (in.)</td>
<td>2.8 (0.11)</td>
<td>n/a</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Compressive strength of 50-mm (2-in.) cubes, M Pa (psi)</td>
<td>91 (12,195)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cube age when tested</td>
<td>7 days</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum concrete strength qualified, MPa (psi)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.2 Av. Str = 91.5 (13,273)(^a)</td>
</tr>
</tbody>
</table>

\(^a\)Nominally a specified strength of 75 MPa (11,000 psi) and perhaps somewhat higher.

5.1.3 The compressive strength of capping materials shall be determined by testing 50-mm (2-in.) cubes following the procedure described in Test Method T 106. Except for sulfur mortars, molding procedures shall be as in Test Method T 106 unless other procedures are required to eliminate large entrapped air voids. See test methods for alternative compaction procedures. Cure cubes in the same environment for the same length of time as the materials used to cap specimens.

5.1.4 The strength of the capping material shall be determined on receipt of a new lot and at intervals not exceeding three months. If a given lot of the capping material fails to conform to the strength requirements, it shall not be used, and strength tests of the replacement material shall be made weekly until four consecutive determinations conform to specification requirements.

5.2 Neat Hydraulic Cement Paste:

5.2.1 Make qualification test of the neat hydraulic cement paste prior to use for capping to establish the effects of water-cement ratio and age on compressive strength of 50-mm (2-in.) cubes (Note 2).

\textbf{Note 2}—The cements used generally conform to Specification M 85 Types I, II, or III; however, Specification M 240 blended cements, calcium aluminate or other hydraulic cements producing acceptable strength may be used.

5.2.2 Mix the neat cement paste to the desired consistency at a water-cement ratio equal to or less than that required to produce the required strength, generally two to four hours before the paste is to be used (Note 3). Remix as necessary to maintain acceptable consistency (Note 4). Some re-tempering of the paste is acceptable if the required water-cement ratio is not exceeded. Optimum consistency is generally produced at water-cement ratios of 0.32 to 0.36 by mass for Type I and Type II cements and 0.35 to 0.39 by mass for Type III cements.
Note 3—Freshly mixed pastes tend to bleed, shrink, and make unacceptable caps. The two to four hour period is generally appropriate for Portland cements.

Note 4—The required consistency of the paste is determined by the appearance of the cap when it is stripped. Fluid paste results in streaks in the cap. Stiff paste results in thick caps.

5.3. High-Strength Gypsum Cement Paste:

5.3.1 No fillers or extenders may be added to neat high-strength gypsum cement paste subsequent to the manufacture of the cement (Note 5). Qualification tests shall be made to determine the effects of water-cement ratio and age on the compressive strength of 50-mm (2-in.) cubes. Retarders may be used to extend working time, but their effects on the required water-cement ratio and strength must be determined (Note 6.)

Note 5—Low-strength molding plaster, plaster of paris, or mixtures of plaster of paris and portland cement are unsuitable for capping.

Note 6—The water-gypsum cement ratio should be between 0.26 and 0.30. Use of low water-cement ratios and vigorous mixing will usually permit development of 35 MPa (5000 psi) at ages of one to two hours. Higher water-gypsum cement ratios extend working time, but reduce strength.

5.3.2 Mix the neat gypsum cement paste at the desired water-cement ratio and use it promptly since it sets rapidly.

5.4 Sulfur Mortar:

5.4.1 Proprietary or laboratory prepared sulfur mortars are permitted if allowed to harden a minimum of two hours before testing concrete with strength less than 35 MPa (5000 psi). For concrete strengths of 35 MPa (5000 psi) or greater, sulfur mortar caps must be allowed to harden at least 16 hours before testing, unless a shorter time has been shown to be suitable as specified in Section 5.1.1.

5.4.2 Determination of Compressive Strength—Prepare test specimens using a cube mold and base plate conforming to the requirements of T 106 and a metal cover plate conforming in principle to the design shown in Figure 1 (Note 7). Bring the various parts of the apparatus to a temperature of 20 to 30ºC (68 to 86ºF), lightly coat the surfaces that will be in contact with the sulfur mortar with mineral oil and assemble near the melting pot. Bring the temperature of the molten-sulfar mortar in the pot within a range of 129 to 143ºC (265 to 290ºF), stir thoroughly, and begin casting cubes. Using a ladle or other suitable pouring device, quickly fill each of the three compartments until the molten material reaches the top of the filling hole. Allow sufficient time for maximum shrinkage, due to cooling, and solidification to occur (approximately 15 minutes) and refill each hole with molten material (Note 8). After solidification is complete, remove the cubes from the mold without breaking off the knob formed by the filling hole in the cover plate. Remove oil, sharp edges, and fins from the cubes and check the planeness of the bearing surfaces in the manner described in T 106. After storage at room temperature to the desired age, but not less than two hours, test cubes in compression following the procedure described in T 106 and calculate the compressive strength in MPa (psi).
**Dimensional Equivalents**

<table>
<thead>
<tr>
<th>mm</th>
<th>6.4</th>
<th>12.7</th>
<th>22.2</th>
<th>44.5</th>
<th>102</th>
<th>254</th>
</tr>
</thead>
<tbody>
<tr>
<td>in.</td>
<td>¼</td>
<td>½</td>
<td>⅞</td>
<td>1¾</td>
<td>4</td>
<td>10</td>
</tr>
</tbody>
</table>

Note: All dimensions shown in millimeters unless otherwise noted.

**Figure 1**—Sketch of Cover for 50-mm (2-in.) Cube Mold

Notes: All dimensions shown in millimeters unless otherwise noted.

**Figure 1**—Sketch of Cover for 50-mm (2-in.) Cube Mold

**Note 7**—If desired, a plane phenol formaldehyde (bakelite) plate of 3 mm (0.125 in.) thickness, provided with three appropriately spaced filling holes, may be inserted between the cover plate and the mold to slow the rate of cooling of test specimens.
Note 8—The second filling helps to prevent the formation of a large void or shrinkage pipe in the body of a cube. However, such defects may occur no matter how much care is exercised, and it therefore is advisable to inspect the interior of tested sulfur mortar cubes for homogeneity whenever the strength values obtained are significantly lower than anticipated.

6. CAPPING PROCEDURES

6.1 Freshly Molded Cylinders — Use only neat Portland cement pastes (Note 9) to cap freshly molded cylinders. Make caps as thin as practicable. Do not apply the neat paste to the exposed end until the concrete has ceased settling in the molds, generally from 2 to 4 hours after molding. During the molding of the cylinder, strike off the upper end even with or slightly below the plane of the rim of the mold. Mix the neat paste to a stiff consistency 2 or 4 hour before it is to be used in order to allow the paste to go through its period of initial shrinkage. The strength of the paste will depend on the consistency, water-cement ratio, curing, brand, and type of cement. For Type I and Type II cement pastes, the optimum consistency is generally produced at a water-cement ratio of 0.32 to 0.36 by mass. For Type III cement, the water ratio should generally be between 0.35 to 0.39 by mass. The paste will stiffen during the 2 to 4 hours waiting period and the use of re-tempering water is not recommended. However, if re-tempering water is used, the amount should not increase the water-cement ratio by more than 0.05 by mass. Remove free water and laitance from the top of the specimen immediately before capping. Form the cap by placing a conical mound of paste on the specimen and then gently pressing a freshly oiled capping plate on the conical mound until the plate contacts the rim of the mold. A very slight twisting motion may be required to extrude excess paste and minimize air voids in the paste. The capping plate must not rock during this operation. Carefully cover the capping plate and mold with a double layer of damp burlap and a polyethylene sheet to prevent drying. Removal of the capping plate after hardening may be accomplished by tapping the edge with a rawhide hammer in a direction parallel to the plane of the cap.

Note 9: Type I neat cement caps generally require at least 6 days to develop acceptable strength and Type III neat cement caps at least 2 days. Dry concrete specimens will absorb water from freshly mixed neat cement paste and produce unsatisfactory caps. Neat cement paste caps will shrink and crack on drying and, therefore, should be used only for specimens which are to be moist-cured continuously until time of testing.

6.2 Hardened Concrete Specimens:

6.2.1 General — Caps should be about 1/8 in. (3 mm) thick, and in no instance shall any part of a cap be more than 5/16 in. (8 mm) thick. If either or both ends of a specimen have coatings or deposits of oily or waxy materials that would interfere with the bond of the cap, remove such coatings or deposits. If necessary, the ends of a specimen may be slightly roughened with a steel file or wire brush to produce proper adhesion of the cap. If desired, capping plates may be coated with a thin layer of mineral oil or grease to prevent the capping material from adhering to the surface of the plate. Form the caps as described in Section 6.1 using capping plates described in Section 4.1 to achieve the alignment required in Section 4.2 (Note 10). Generally capping plates may be removed within 45 minutes with gypsum cement pastes and after 12 hours with neat cement paste, without visibly damaging the cap.
**Note 10**—A number of methods have been used to obtain the desired perpendicularity of the cap to the axis of the cylinder. A mound of paste can be placed on a capping plate and the specimen lowered into it. A bull’s-eye level on the top of the cylinder helps obtain alignment. A mound of paste can be placed on top of the cylinder and a capping plate pressed into it, again using the bull’s-eye level. A better system is to make a half-height mold with a vertical split so that it can be slipped over the hardened cylinder. A clamp is used to position the mold and to ensure the required cap thickness. The mound of paste can then be placed either on a capping plate or on top of the cylinder and pressed until the plate contacts the mold. As noted earlier, very stiff paste may require excessive force and produce thick or defective caps.

6.2.2 **End Condition**—The distance of any point on an uncapped end from a plane that passes through the highest point of the end surface and is perpendicular to the axis of the cylinder shall not exceed 3 mm (0.125 in.) (Note 11). If the end exceeds this limit, the end of the cylinder shall be cut, lapped or ground prior to capping.

**Note 11**—This provision is to control the difference between the thickest and thinnest parts of a cap. The distance may be checked using a square with one blade touching the cylinder parallel to the cylinder axis and the other blade touching the highest point on the end of the cylinder. The distance between the blade of the square and the lowest point on the end of the cylinder is measured.

6.2.3 **Capping with High-Strength Gypsum Plaster**—Mix high-strength plaster for capping, using the same percent of mixing water as was used in making the qualification test described in Section 5.2.1 (Note 12).

**Note 12**—High-strength gypsum caps soften and deteriorate on contact with water and cannot be used on freshly mixed concrete or stored in a moist room for more than very brief periods up to four hours.

6.2.4 **Capping with Sulfur Mortar**—Prepare sulfur mortar for use by heating to about 265°F (130°C), as periodically determined by an all-metal thermometer inserted near the center of the mass. Empty the pot and recharge with fresh material at frequent enough intervals to ensure that the oldest material in the pot has not been used more than five times (Note 13). Fresh sulfur mortar must be dry at the time it is placed in the pot as dampness may cause foaming. Keep water away from molten sulfur mortar for the same reason. The capping plate or device should be warmed slightly before use to slow the rate of hardening and permit the production of thin caps. Oil the capping plate lightly and stir the molten sulfur mortar immediately prior to pouring each cap. The ends of moist-cured specimens shall be dry enough at the time of capping to preclude the formation of steam or foam pockets under or in the cap larger than 1/4 in. (6 mm) in diameter. To ensure that the cap shall be bonded to the surface of the specimen, the end of the specimen shall not be oiled prior to application of the cap. When using a vertical device, pour the mortar onto the surface of the capping plate, lift the cylinder above the plate and contact the cylinder sides with the guides; slide the cylinder down the guides onto the capping plate while keeping constant contact with the alignment guides. The cylinder end should continue to rest on the capping plate with cylinder sides in positive contact with the alignment guides until the mortar has hardened. Use sufficient material to cover the cylinder end after the sulfur mortar solidifies. The sulfur mortar cap may be tapped or rubbed with a light metal implement. If a hollow sound is produced, an unsatisfactory mortar cap is indicated. See Section 25.16 of the ASTM Manual of Aggregate and Concrete Testing.
Note 13: Reuse of material must be restricted in order to minimize loss of strength and pourability occasioned by contamination of the mortar with oil miscellaneous debris, and loss of sulfur through volatilization.

6.2.4.1 Caution: Hydrogen sulfide gas may be produced during capping when sulfur mortar is contaminated with organic materials such as paraffin or oil. The gas is colorless and has a notoriously bad odor of rotten eggs; however, the odor should not be relied upon as a warning sign, since the sensitivity to the odor disappears rapidly on exposure. High concentrations are lethal and less concentrated dosages may produce nausea, stomach ache, distress, dizziness, headache, or irritation of the eyes. For this and other reasons, it is desirable that the melting pot be located under a hood or near an exhaust fan and that the capping area be well ventilated.

6.2.4 Daily Check — During each day’s capping operation, planeness of the caps on at least three specimens representing the start, middle, and end of the run, shall be check by means of a straight-edge and feeler gage, making a minimum of three measurements on different diameters to ensure that the surfaces of the caps do not depart from a plane by more than 0.05 mm (0.002 in.).

7. PROTECTION OF SPECIMENS AFTER CAPPING

7.1 Moist-cured specimens shall be maintained in a moist condition between the completion of capping and the time of testing by returning them to moist storage or wrapping them with a double layer of wet burlap. Specimens with gypsum plaster caps shall not be immersed in water and shall not be stored in a moist room for more than 4 hours. If stored in a moist room, the plaster caps shall be protected against water dripping on their surfaces.
APPENDIX
(Nonmandatory Information)

A1. COMPOSITION AND DETERMINATION OF LOSS ON IGNITION OF SULFUR MORTAR CAPPING MATERIALS

A1.1 Composition:

A1.1.1. Loss on ignition, percent 48 to 70 Residue after ignition, percent 30 to 52.

A1.2 Determination of Loss on Ignition – Obtain samples from caps on concrete cylinders or from cast specimens similar to caps in size and thickness. Divide each cap-size specimen into eight approximately equal triangular sections, and secure test samples by breaking either two or four of the triangular sections into small pieces with the fingers. Using a balance capable of determining mass to an accuracy of 0.01 g, measure out 20 to 25 g of fragmented material in a previously ignited, cooled, and tared Coors No. 3, high-form porcelain crucible. Place the crucible on a ring approximately 2 in. (50 mm) above a Terrel-type Bunsen burner and adjust the flame so that the sulfur burns slowly without spattering (Note 14), (see Section 4). When the sulfur has been completely consumed, adjust the burner for high heat and ignite the residue for 30 minutes. Cool the crucible and residue in a desiccator and determine the mass. Continue to ignite, cool, and determine the mass of the crucible until a constant mass is obtained. Calculate the percentage of loss on ignition C, as follows (Note 15):

\[ C = \frac{X}{100} \]

where:

A = original mass of sample less mass of the residue after ignition, and

B = original mass of sample.

NOTE 14– Where the filler is known or found to be composed of carbonate minerals, the ignition test shall be made at a carefully controlled temperature in the range from 600 to 650°C, to prevent calcinations of the mineral. Small amounts of plasticizer and carbon filler will be included in the reported value for loss on ignition using the simple test herein described.

NOTE 15 – A referee procedure for the determination of the percent of sulfur contained in sulfur mortar may be found in ASTM C 287.
# Performance Exam Checklist

**Capping Cylindrical Concrete Specimens**  
**FOP for AASHTO T 231**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

## Procedure Element

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>The tester has a copy of the current procedure on hand?</td>
<td></td>
</tr>
<tr>
<td>2.</td>
<td>All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td></td>
</tr>
</tbody>
</table>

## Sulpher Only

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Is the temperature taken and the mortar stirred at the beginning of the operation?</td>
<td></td>
</tr>
<tr>
<td>2.</td>
<td>Is the capping plate lightly oiled prior to use?</td>
<td></td>
</tr>
<tr>
<td>3.</td>
<td>Are cylinder ends dried prior to capping?</td>
<td></td>
</tr>
<tr>
<td>4.</td>
<td>Is the capping plate warmed slightly?</td>
<td></td>
</tr>
</tbody>
</table>

## Renumber rest

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.</td>
<td>Are perpendicularity guides or leveling devices used effectively?</td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td>Are caps checked for planeness?</td>
<td></td>
</tr>
<tr>
<td>5a.</td>
<td>If yes, how often?</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>Are cylinders kept moist after capping?</td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner ____________________________

Comments:

_________________________________________________________________________
                                                                                      
_________________________________________________________________________
                                                                                      
_________________________________________________________________________
Reducing Samples of Aggregate to Testing Size

1. Scope

1.1 This method covers for the reduction of large samples of aggregate to the appropriate size for testing employing techniques that are intended to minimize variations in measured characteristics between the test samples so selected and the large sample.

1.2 The values stated in English units are to be regarded as the standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 AASHTO Standards:
   T 2 Sampling of Aggregate
   T 84 Specific Gravity and Absorption of Coarse Aggregate

2.2 ASTM Standards:
   C 125 Terminology Relating to Concrete and Concrete Aggregates

3. Terminology

3.1 Definitions — The terms used in this practice are defined in ASTM C 125.

4. Significance and Use

4.1 Specifications for aggregates require sampling portions of the material for testing. Other factors being equal, larger samples will tend to be more representative of the total supply. These methods provide for reducing the large sample obtained in the field or produced in the laboratory to a convenient size for conducting a number of tests to describe the material and measure its quality in a manner that the smaller test sample portion is most likely to be a representation of the larger sample, and thus of the total supply. The individual test methods provide for minimum amount of material to be tested.

4.2 Under certain circumstances, reduction in size of the large sample prior to testing is not recommended. Substantial differences between the selected test samples sometimes cannot be avoided, as for example, in the case of an aggregate having relatively few large size particles in the sample. The laws of chance dictate that these few particles may be unequally distributed among the reduced size test samples. Similarly, if the test sample is being examined for certain contaminants occurring as a few discrete fragments in only small percentages, caution should be used in interpreting results from the reduced size test sample. Chance inclusion or exclusion of only one or two particles in the selected test sample may importantly influence interpretation of the characteristics of the original sample. In these cases, the entire original sample should be tested.

\[1\]This FOP is based on AASHTO T 248-02.
4.3 Failure to carefully follow the procedures in this practice could result in providing a nonrepresentative sample to be used in subsequent testing.

5. SELECTION OF METHOD

5.1 Fine Aggregate — Samples of fine aggregate that are drier than the drier saturated-surface-dry condition or drier (Note 1) may be reduced using a mechanical splitter according to Method A. Samples having free moisture on the particle surfaces may be reduced in size by quartering according to Method B, or by treating as a miniature stockpile as described in Method C.

5.1.1 If the use of Method B or Method C is desired, and the sample does not have free moisture on the particle surfaces, the sample may be moistened to achieve this condition, thoroughly mixed, and then the sample reduction performed.

Note 1: The method of determining the saturated-surface-dry condition is described in Test Method T 84. As a quick approximation, if the fine aggregate will retain its shape when molded in the hand, it may be considered to be wetter than saturated-surface-dry.

5.1.2 If use of Method A is desired and the sample has free moisture on the particle surfaces, the entire sample may be dried to at least the saturated-surface-dry condition, using temperatures that do not exceed those specified for any of the tests contemplated, and then the sample reduction performed. Alternatively, if the moist sample is very large, a preliminary split may be made using a mechanical splitter having wide chute openings of 1½ in. (38 mm) or more to reduce the sample to not less than 5000 g. The portion so obtained is then dried, and reduction to test sample size is completed using Method A.

5.2 Coarse Aggregates and Mixtures of Coarse and Fine Aggregates — Reduce the sample using a mechanical splitter in accordance with Method A (preferred method) or by quartering in accordance with Method B. The miniature stockpile Method C is not permitted for coarse aggregates or mixtures of coarse and fine aggregates.

5.3 Untreated materials shall be prepared for testing using this procedure. Treated materials (i.e., Hot Mix Asphalt or Asphalt Treated Base) shall be prepared for testing using WSDOT Test Method No. T 712 for reduction of size of samples of Asphalt treated materials.

6. SAMPLING

6.1 The samples of aggregate obtained in the field shall be taken in accordance with T 2, or as required by individual test methods. When tests for sieve analysis only are contemplated, the size of field sample listed in T 2 is usually adequate. When additional tests are to be conducted, the user shall determine that the initial size of the field sample is adequate to accomplish all intended tests. Similar procedures shall be used for aggregate production in the laboratory.
**Method A — Mechanical Splitter**

7. APPARATUS

7.1 Sample Splitter — Sample splitters shall have an even number of equal width chutes, but not less than a total of eight for coarse aggregate, or 12 for fine aggregate, which discharge alternately to each side of the splitter. For coarse aggregate and mixed aggregate, the minimum width of the individual chutes shall be approximately 50 percent larger than the largest particles in the sample to be split (Note 2). For dry fine aggregate in which the entire sample will pass the 3/8 in. (9.5 mm) sieve, the minimum width of the individual chutes shall be at least 50 percent larger than the largest particles in the sample and the maximum width shall be 3/4 in. (19 mm). The splitter shall be equipped with two receptacles to hold the two-halves of the sample following splitting. It shall also be equipped with a hopper or straight edge pan which has a width equal to or slightly less than the overall width of the assembly of chutes, by which the sample may be fed at a controlled rate to the chutes. The splitter and accessory equipment shall be so designed that the sample will flow smoothly without restriction or loss of material (Figure 1).

![Figure 1: Sample Dividers (Riffles)](image)
Note 2: Mechanical splitters are commonly available in sizes adequate for coarse aggregate having the largest particle not over 1 1/2 in. (37.5 mm).

8. PROCEDURE

8.1 Place the original sample in the hopper or pan and uniformly distribute it from edge to edge, so that when it is introduced into the chutes, approximately equal amounts will flow through each chute. The rate at which the sample is introduced shall be such as to allow free flowing through the chutes into the receptacles below. Reintroduce the portion of the sample in one of the receptacles into the splitter as many times as necessary to reduce the sample to the size specified for the intended test. The portion of the material collected in the other receptacle may be reserved for reduction in size for other tests.

Method B — Quartering

9. APPARATUS

9.1 Apparatus shall consist of a straightedge, scoop, shovel, or trowel; a broom or brush; and a canvas blanket approximately 6 by 8 ft. (2 by 2.5 m).

10. PROCEDURE

10.1 Use either the procedure described in 10.1.1 or 10.1.2 or a combination of both.

10.1.1 Place the original sample on a hard clean, level surface where there will be neither loss of material nor the accidental addition of foreign material. Mix the material thoroughly by turning the entire sample over three times. With the last turning, shovel the entire sample into a conical pile by depositing each shovelful on top of the preceding one. Carefully flatten the conical pile to a uniform thickness and diameter by pressing down the apex with a shovel so that each quarter sector of the resulting pile will contain the material originally in it. The diameter should be approximately four to eight times the thickness. Divide the flattened mass into four equal quarters with a shovel or trowel and remove two diagonally opposite quarters, including all fine material, and brush the cleared spaces clean. Successively mix and quarter the remaining material until the sample is reduced to the desired size (Figure 2).

Figure 2: Quartering on a Hard, Clean Level Surface
10.1.2 As an alternative to the procedure in 10.1.1 when the floor surface is uneven, the field sample may be placed on a canvas blanket and mixed with a shovel as described in 10.1.1, or by alternatively lifting each corner of the canvas and pulling it over the sample toward the diagonally opposite corner causing the material to be rolled. Flatten the pile as described in 10.1.1. Divide the sample as described in 10.1.1 or if the surface beneath the blanket is uneven, insert a stick or pipe beneath the blanket and under the center of the pile, then lift both ends of the stick, dividing the sample into two equal parts. Remove the stick leaving a fold of the blanket between the divided portions. Insert the stick under the center of the pile at right angles to the first division and again lift both ends of the stick, dividing the sample into four equal parts. Remove two diagonally opposite quarters, being careful to clean the fines from the blanket. Successively mix and quarter the remaining material until the sample is reduced to the desired size (Figure 3).

Figure 3: Quartering on a Canvas Blanket
Method C — Miniature Stockpile Sampling (Damp Fine Aggregate Only)

11. APPARATUS

11.1 Apparatus shall consist of a straight-edged scoop, shovel, or trowel for mixing the aggregate, and either a small sampling thief, small scoop, or spoon for sampling.

12. PROCEDURE

12.1 Place the original sample of damp fine aggregate on a hard clean, level surface where there will be neither loss of material nor the accidental addition of foreign material. Mix the material thoroughly by turning the entire sample over three times. With the last turning, shovel the entire sample into a conical pile by depositing each shovelful on top of the preceding one. If desired, the conical pile may be flattened to a uniform thickness and diameter by pressing the apex with a shovel so that each quarter sector of the resulting pile will contain the material originally in it. Obtain a sample for each test by selecting at least five increments of material at random locations from the miniature stockpile, using any of the sampling devices described in 11.1.
Performance Exam Checklist

Reducing Samples of Aggregates to Testing Size
FOP for AASHTO T 248

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Selection of Method

1. Fine Aggregate
   A. Saturated surface dry or drier: Method A (Splitter) used?                     |     |    |
   B. Free moisture present: Method B (Quartering) used?                           |     |    |
2. Coarse Aggregate and Mixtures of Fine and Coarse Aggregates
   A. Method A used (preferred)?                                                   |     |    |
   B. Method B used?                                                              |     |    |

Method A — Splitting

1. Material spread uniformly on feeder?                                           |     |    |
2. Rate of feed slow enough so that sample flows freely through chutes?           |     |    |
3. Material in one pan re-split until desired mass is obtained?                   |     |    |
4. Chutes are set correctly for material being split?                            |     |    |

Method B — Quartering

1. Sample placed on clean, hard, and level surface?                               |     |    |
2. Mixed by turning over 3 times with shovel or by raising canvas and pulling over pile? |     |    |
3. Conical pile formed?                                                          |     |    |
4. Diameter equal to about 4 to 8 times thickness?                                |     |    |
5. Pile flattened to uniform thickness and diameter?                             |     |    |
6. Divided into 4 equal portions with shovel or trowel?                           |     |    |
7. Two diagonally opposite quarters, including all fine material, removed?      |     |    |
<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>8. Cleared space between quarters brushed clean?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Process continued until desired sample size is obtained when two opposite quarters combined?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The sample may be placed upon a blanket and a stick or pipe may be placed under the blanket to divide the pile into quarters.*

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner ________________________________

Comments:

_________________________________________________
_________________________________________________
_________________________________________________
_________________________________________________
_________________________________________________
_________________________________________________
_________________________________________________
_________________________________________________
_________________________________________________
WSDOT FOP for AASHTO T 255

Total Evaporable Moisture Content of Aggregate by Drying

1. SCOPE

1.1 This test method covers the determination of the percentage of evaporable moisture in a sample of aggregate by drying, both surface moisture and moisture in the pores of the aggregate. Some aggregate may contain water that is chemically combined with the minerals in the aggregate. Such water is not evaporable and is not included in the percentage determined by this test method.

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

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use. For specific precautionary statements, see 5.3.1, 7.2.1, and 7.3.1.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:
- M 92 Wire-Cloth Sieves for Testing Purposes
- M 231 Weighing Devices Used in Testing Materials
- R 16 Regulatory Information for Chemicals Used in AASHTO Tests
- T 2 Sampling of Aggregate
- T 19/T 19M Bulk Density (“Unit Weight”) and Voids in Aggregate
- T 84 Specific Gravity and Absorption of Coarse Aggregate
- T 85 Specific Gravity and Absorption of Fine Aggregate

2.2 ASTM Standards:
- C 125 Terminology Relating to Concrete and Concrete Aggregates
- C 670 Practice for Preparing Precision Statements for Test Methods for Construction Materials

3. TERMINOLOGY

3.1 Definitions:

3.1.1 For definitions of terms used in this test method, refer to ASTM C 125.

4. Significance and Use

4.1 This test method is sufficiently accurate for usual purposes, such as adjusting batch quantities of ingredients for concrete. It will generally measure the moisture in the test sample more reliably than the sample can be made to represent the aggregate supply. In rare cases where the aggregate itself is altered by heat, or where more refined measurement is required, the test should be conducted using a ventilated, controlled temperature oven.

This FOP is based on AASHTO T 255-00.
4.2 Large particles of coarse aggregate, especially those larger than 2 in. (50 mm), will require greater time for the moisture to travel from the interior of the particle to the surface. The user of this test method should determine by trial if rapid drying methods provide sufficient accuracy for the intended use when drying large-size particles.

5. APPARATUS

5.1 Balance — The balances shall have sufficient capacity, be readable to 0.1 percent of the sample mass, or better, and conform to the requirements of M 231.

5.2 Source of Heat — A ventilated oven capable of maintaining the temperature surrounding the sample at 110 ± 5°C (230 ± 9°F). Where close control of the temperature is not required (see Section 4.1), other suitable sources of heat may be used, such as an electric or gas hot plate, electric heat lamps, or a ventilated microwave oven.

5.3 Sample Container — A container not affected by the heat, and of sufficient volume to contain the sample without danger of spilling, and of such shape that the depth of sample will not exceed one fifth of the least lateral dimension.

5.3.1 Precaution — When a microwave oven is used, the container shall be nonmetallic.

   Note 1: Except for testing large samples, an ordinary frying pan is suitable for use with a hot plate, or any shallow flat-bottomed metal pan is suitable with heat lamps or oven. Note Precaution in Section 5.3.1.

5.4 Stirrer — A metal spoon or spatula of convenient size.

6. SAMPLING

6.1 Sampling shall generally be accomplished in accordance with T 2, except for the sample size may be as stated in Table 1.

6.2 Secure a sample of the aggregate representative of the moisture content in the supply being tested and having a mass not less than the amount listed in Table 1. Protect the sample against loss of moisture prior to determining the mass.
Table 1
Sample Size for Aggregate

<table>
<thead>
<tr>
<th>Nominal Maximum Size* of Aggregate, in. (mm)</th>
<th>Mass of Normal Weight Aggregate Sample, min, kg*</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 4 (4.75)</td>
<td>0.5</td>
</tr>
<tr>
<td>3/8 (9.5)</td>
<td>1.5</td>
</tr>
<tr>
<td>1/2 (12.5)</td>
<td>2</td>
</tr>
<tr>
<td>3/4 (19.0)</td>
<td>3</td>
</tr>
<tr>
<td>1 (25.0)</td>
<td>4</td>
</tr>
<tr>
<td>1 1/2 (37.5)</td>
<td>6</td>
</tr>
<tr>
<td>2 (50)</td>
<td>8</td>
</tr>
<tr>
<td>2 1/2 (63)</td>
<td>10</td>
</tr>
<tr>
<td>3 (75)</td>
<td>13</td>
</tr>
<tr>
<td>3 1/2 (90)</td>
<td>16</td>
</tr>
<tr>
<td>4 (100)</td>
<td>25</td>
</tr>
<tr>
<td>6 (150)</td>
<td>50</td>
</tr>
</tbody>
</table>

* For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.

Note: For an aggregate specification having a generally unrestricted gradation (i.e., wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.

Note: When determining moisture content for T99 samples, use approximately 100 grams, and approximately 500 grams for T180 samples.

Based on sieves with square openings.

To determine the minimum sample weight for lightweight aggregate, multiply the value by the approximate dry-loose unit mass of the aggregate in kg/m³ and dividing by 1600.

7. PROCEDURE

7.1 Determine the mass of the sample to the nearest 0.1 percent or better of the total sample mass.

7.2 Dry the sample thoroughly in the sample container by means of the selected source of heat, exercising care to avoid loss of any particles. Very rapid heating may cause some particles to explode, resulting in loss of particles. Use a controlled temperature oven when excessive heat may alter the character of the aggregate, or where more precise measurement is required. If a source of heat other than the controlled temperature oven is used, stir the sample during drying to accelerate the operation and avoid localized overheating. When using a microwave oven, stirring of the sample is optional.

7.2.1 Caution — When using a microwave oven, occasionally minerals are present in aggregates that may cause the material to overheat and explode. If this occurs it can damage the microwave oven.
7.3 When a hot plate is used, drying can be expedited by the following procedure. Add sufficient anhydrous denatured alcohol to cover the moist sample. Stir and allow suspended material to settle. Decant as much of the alcohol as possible without losing any of the sample. Ignite the remaining alcohol and allow it to burn off during drying over the hot plate.

7.3.1 Warning — Exercise care to control the ignition operation to prevent injury or damage from the burning alcohol.

7.4 The sample is thoroughly dry when further heating causes, or would cause, less than 0.1 percent additional loss in mass.

WSDOT NOTE: When weighing hot samples, use a heat sink so the balance is not damaged from excessive overheating.

7.5 Determine the mass of the dried sample to the nearest 0.1 percent or better of the total sample mass after it has cooled sufficiently not to damage the balance.

WSDOT NOTE: When weighing hot samples, use a heat sink so not to damage the balance.

8. CALCULATION

8.1 Calculate total evaporable moisture content as follows:

\[ p = 100 \frac{(W - D)}{D} \]

where:

- \( p \) = total evaporable moisture content of sample, percent;
- \( W \) = mass of original sample, g; and
- \( D \) = mass of dried sample, g

8.2 Surface moisture content is equal to the difference between the total evaporated moisture content and the absorption, with all values based on the mass of a dry sample. Absorption may be determined in accordance with T 85, Test for Specific Gravity and Absorption of Coarse Aggregates, or T 84, Test for Specific Gravity and Absorption of Fine Aggregates

9. PRECISION AND BIAS

See AASHTO T-255

10. REPORT

Report results using WSDOT Form 422-020, or other report approved by the State Materials Engineer.
Performance Exam Checklist

Total Moisture Content of Aggregate by Drying
FOP for AASHTO T 255

Participant Name ___________________________________________ Exam Date ________________

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required,</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Representative sample of appropriate mass obtained?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Mass of clean, dry container determined?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Sample placed in container and mass determined?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Test sample mass conforms to the required mass?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Sample mass determined to 0.1 percent?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Loss of moisture avoided prior to mass determination?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. Sample dried by a suitable heat source?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. Sample cooled prior to mass determination?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>11. If aggregate heated by means other than a controlled oven, is sample</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>stirred to avoid localized overheating?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>12. Mass determined and compared to previous mass – showing less than 0.1 percent loss?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>13. Calculations performed properly and results reported to the nearest 0.1 percent?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt:  Pass ☐  Fail ☐  Second attempt:  Pass ☐  Fail ☐

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
AASHTO T 265 (Modified)

Laboratory Determination of Moisture Content of Soils

AASHTO T 265 has been adopted by WSDOT with the following definition for “Constant Mass.”

“Constant Mass — Test samples dried at a temperature of 230 ± 9º F (110 ± 5º C) to such a condition that it will not lose more than 0.1 per cent moisture after 2 hours of drying. Such a condition of dryness can be verified by determining the mass of the sample before and after successive 2 hour drying periods. In lieu of such determination, samples may be considered to have reached constant mass when they have been dried at a temperature of 230 ± 9º F (110 ± 5º C) for an equal or longer period than that previously found adequate for producing the desired constant mass condition under equal or heavier loading condition of the oven.”
WSDOT FOP for AASHTO T 272

Family of Curves — One-point Method

1. SCOPE

1.1 These methods of tests are for the rapid determination of the maximum density and optimum moisture content of a soil sample utilizing a family of curves and a one-point determination.

1.2 One-point determinations are made by compacting the soil in a mold of a given size with a 5.5-lb (2.5-kg) rammer dropped from a height of 12 in. (305 mm). Four alternate procedures are provided as follows:

   Method A — A 4-in. (101.6 mm) mold; soil material passing a No. 4 (4.75-mm) sieve. Sections 4 and 5.

   Method B — A 6-in. (152.4-mm) mold; soil material passing a No. 4 (4.75-mm) sieve. Sections 6 and 7.

   Method C — A 4-in. (101.6 mm) mold; soil material passing a 3/4 in. (19.0-mm) sieve. Sections 8 and 9.

   Method D — A 6-in. (152.4-mm) mold; soil material passing a 3/4 in. (19.0-mm) sieve. Sections 10 and 11.

   The preferred method of WSDOT is to use method A.

1.3 The methods described herein correspond to the methods in T 99 and must be chosen accordingly; i.e., when moisture-density relationships as determined by T 99 Method C are used to form the family of curves, then Method C described in this procedure must be used for the one-point determination (Note 1).

   Note 1: Direct reference to T 99 is made throughout these test methods and most terminology, apparatus and procedures are the same.

1.4 In addition, the concepts described herein are applicable to one-point determinations and moisture-density relationships as specified in T 180 with appropriate apparatus and method used as required.

1.5 The following applies to all specified limits in this standard: For the purposes of determining conformance with these specifications, an observed value or a calculated value shall be rounded off “to the nearest unit” in the last right-hand place of figures used in expressing the limiting value, in accordance with the rounding-off method of R 11, Recommended Practice for Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values.

1.6 The values stated in English units are to be regarded as the standard.

¹This FOP is based on AASHTO T 272-04
FIGURE 1  Example of Curves

ONE POINT VALUES
Dry Density = 1871 kg/m$^3$ (116.8 PCF)
Moisture Content = 11.2%

Maximum Density = 1906 kg/m$^3$ (119.0 PCF)
Optimum Moisture Content = 12.7%
2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:
   • R 11, Indicating Which Places of Figures Are to Be Considered Significant in Specified Limiting Values
   • T 19/T 19M, Bulk Density ("Unit Weight") and Voids in Aggregate
   • T 99, Moisture-Density Relations of Soils Using a 2.5-kg (5.5-lb) Rammer and a 305-mm (12-in.) Drop
   • T 180, Moisture-Density Relations of Soils Using a 4.54-kg (10-lb) Rammer and a 457-mm (18-in.) Drop

3. DEFINITION

3.1 A family of curves is a group of typical soil moisture-density relationships determined using T 99, which reveal certain similarities and trends characteristic of the soil type and source. Soils sampled from one source will have many different moisture-density curves, but if a group of these curves are plotted together certain relationships usually become apparent. In general it will be found that higher unit mass soils assume steeper slopes with maximum dry densities at lower optimum moisture contents, while the lower unit mass soils assume flatter more gently sloped curves with higher optimum moisture contents (Figure 1).

4. APPARATUS

4.1 See T 99, Section 3.

METHOD A

5. SAMPLE

5.1 See T 99, Section 4.

6. PROCEDURE

6.1 Thoroughly mix the selected representative sample with sufficient water to dampen approximately 4 percentage points below optimum moisture content. Greater accuracy in the determination of the maximum density will result as the moisture content used approaches optimum moisture content. Moisture content of the sample should never exceed the optimum water content. When doing a one-point determination in the field, use the sample as obtained and determine the moisture after the test.

6.2 Form a specimen by compacting the prepared soil in the 4-in. (101.6-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5 in. (125 mm). Compact each layer by 25 uniformly distributed blows from the rammer dropping free from a height of 12 in. (305 mm) above the elevation of the soil when a sleeve-type rammer is used, or from 12 in. (305 mm) above the approximate elevation of compacted soil when a stationary mounted type of rammer is used. During compaction, the mold shall rest firmly on a dense uniform, rigid and stable foundation (Note 2).

Note 2: Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: A block of concrete, with a mass not less than 200 lb (91 kg) supported by a relatively stable foundation; a sound concrete floor; and for field application, such surfaces as are found in concrete box culverts, bridges, and pavements.
6.2.1 Following compaction, remove the extension collar, carefully trim the compacted soil even with the top of the mold by means of the straightedge, and determine the mass of the mold and moist soil in kilograms to the nearest 5 grams, or determine the mass in pounds to the nearest 0.01 pounds. For molds conforming to tolerances given in T 99 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 1060, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in T 99 and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 30, and record the result as the wet density, \( W_1 \), in pounds per cubic foot, of compacted soil. For used molds out of tolerance by not more than 50 percent (T 99), use the factor for the mold as determined in accordance with AASHTO T 19.

6.3 Remove the material from the mold and slice vertically through the center. Take a representative sample of the material from one of the cut faces, determine the mass immediately, and dry in an oven at 110 ± 5°C (230 ± 9°F), for at least 12 hours, or to a constant mass to determine the moisture content in accordance with AASHTO T 255 or T 217. The moisture sample shall have a mass not less than 100 g.

**WSDOT Note** — When developing a compaction curve for free draining soils, such as uniform sands and gravels, where seepage occurs at the bottom of the mold and base plate, taking a representative moisture content sample from the mixing bowl may be preferred in order to determine the amount of moisture available for compaction.

**METHOD B**

7. SAMPLE

7.1 Select the representative sample in accordance with Section 4, except that it shall have a mass of approximately 16 lb (7 kg).

8. PROCEDURE

8.1 Follow the same procedure as described for Method A in Section 5, except for the following: Form a specimen by compacting the prepared soil in the 6-in. (152.4-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5-in. (125 mm), each layer being compacted by 56 uniformly distributed blows from the rammer. For molds conforming to tolerances given in T 99, and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 471, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter of compacted soil. For molds conforming to tolerances given in T 99, and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 13.33, and record the result as the wet density, \( W_1 \), in pounds per cubic foot, of the compacted soil. For used molds out of tolerance by not more than 50 percent (T 99), use the factor for the mold as determined in accordance with AASHTO T 19.

**METHOD C**

9. SAMPLE

9.1 If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of drying apparatus such that the temperature does not exceed 140°F (60°C). Then thoroughly break up the aggregations in such a manner as to avoid reducing the natural size of individual particles.
9.2 Sieve an adequate quantity of the representative pulverized soil over the 3/4 in. (19.0-mm) sieve. Discard the coarse material, if any, retained on the 3/4 in. (19.0-mm) sieve (Note 3).

**Note 3:** The use of a replacement method, where the oversized particles are replaced with finer particles to maintain the same percentage of coarse material, is not considered appropriate to compute the maximum density.

9.3 Select a representative sample having a mass of approximately 12 lb (5 kg) or more of the soil prepared as described in Sections 9.1 and 9.2.

10. **PROCEDURE**

10.1 Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately 4 percentage points below optimum moisture content. Greater accuracy in the determination of the maximum density will result as the moisture content used approaches the optimum moisture content.

10.2 Form a specimen by compacting the prepared soil in the 4-in. (101.6-mm) mold (with collar attached) in three approximately equal layers to give total compacted depth of about 5 in. (125 mm). Compact each layer by 25 uniformly distributed blows from the rammer dropping free from a height of 12 in. (305 mm) above the elevation of the soil when a sleeve-type rammer is used or from 12 in. (305 mm) above the approximate elevation of each finely compacted layer when a stationary mounted type rammer is used. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation (Note 2).

10.2.1 Following compaction, remove the extension collar and carefully trim the compacted soil even with the top of the mold by means of the straightedge. Holes developed in the surface by removal of coarse material shall be patched with smaller size material. Determine the mass of the mold and moist soil in kilograms to the nearest 5 grams, or determine the mass in pounds to the nearest 0.01 pounds. For molds conforming to tolerances given in T 99 and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 1060, and record the result as the wet density, \( W_1 \), in kilograms per cubic meter of compacted soil. For molds conforming to tolerances given in T 99 and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 30, and record the result as the wet density, \( W_1 \), in pounds per cubic foot of compacted soil. For used molds out of tolerance by not more than 50 percent (T 99), use the factor for the mold as determined in accordance with T-19.

10.3 Remove the material from the mold and slice vertically through the center. Take a representative sample of the material from one of the cut faces, determine the mass immediately and dry to a constant mass using a drying apparatus described in T 99 to determine the moisture content. The moisture sample shall have a mass not less than 500 g.

**METHOD D**

11. **SAMPLE**

11.1 Select the representative sample in accordance with Section 8.3 except that it shall have a mass of approximately 25 lb (11 kg).
12. PROCEDURE

12.1 Follow the same procedure as described for Method C in Section 9, except for the following:
Form a specimen by compacting the prepared soil in the 6-in. (152.4-mm) mold (with collar attached) in three approximately equal layers to give a total compacted depth of about 5 in. (125 mm), each layer being compacted by 56 uniformly distributed blows from the rammer. For molds conforming to tolerances given in T 99, and masses recorded in kilograms, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 471, and record the result as the wet density, $W_1$, in kilograms per cubic meter, of compacted soil. For molds conforming to tolerances given in T 99, and masses recorded in pounds, multiply the mass of the compacted specimen and the mold, minus the mass of the mold, by 13.33, and record the result as the wet density, $W_1$, in pounds per cubic foot, of the compacted soil. For used molds out of tolerance by not more than 50 percent (T 99), use the factor for the mold as determined in accordance with AASHTO T 19.

CALCULATIONS AND REPORT

13. CALCULATIONS

12.1 See T 99, Section 12.

14. MAXIMUM DENSITY AND OPTIMUM MOISTURE CONTENT DETERMINATION

14.1 The calculations in Section 12.1 shall be made to determine the moisture content and corresponding over-dry density (mass) in pounds per cubic foot (kilograms per cubic meter) of the compacted specimen. The dry density (unit mass) of the soil shall be plotted as ordinate and the corresponding moisture content as the abscissa to define one-point within or on the family of curves (Figure 1).

14.2 If the one-point falls on one of the curves in the family of curves the maximum dry density and optimum moisture content defined by that curve shall be used (Note 4).

14.3 If the one-point falls within the family but not on a curve, a new curve shall be drawn through the plotted one-point parallel and in character with the nearest existing curve in the family of curves. The maximum dry density and optimum moisture content as defined by the new curve shall be used (Note 4).

**Note 4:** If the one-point plotted within or on the family of curves does not fall in the 80 to 100 percent of optimum moisture range, compact another specimen, using the same material, at an adjusted moisture content that will place the one-point within this range.

14.3.1 If the family of curves is such that the profile of a new curve to be drawn through a one-point is not well defined or in any way questionable, then a full moisture-density relationship shall be made for the soil in question to correctly define the new curve and verify the applicability of the family of curves (Note 5).

**Note 5:** New curves drawn through plotted one-point determinations shall not become a permanent part of the family of curves until verified by a full moisture-density relationship.
16. REPORT

16.1 The report shall include the following:

16.1.1 The method used (Method A, B, C, or D).

16.1.2 The optimum moisture content as a percentage to the nearest whole number.

16.1.3 The maximum density to the nearest 1.0 lb/ft³ (0.5 kg/m³).

16.1.4 In Methods C and D indicate if the material retained on the 3/4-in. (19.0-mm) sieve was removed or replaced.

16.1.5 Type of face if other than 2-in. (50.8-mm) circular.

**Note 6:** Inherent variability of soils places limitations on this method of test. The person using this test method must realize this and become thoroughly familiar with the material being tested. Knowledge of the AASHTO Soil Classification System and ability to recognize the gradation of soils are requirements for this work.
APPENDIX

DEVELOPING A MOISTURE-DENSITY FAMILY OF CURVES

The purpose of the family of curves is to represent the average moisture-density characteristics of the material. The family must, therefore, be based on moisture-density relationships which adequately represent the entire mass range and all types of material for which the family is to be used. It may be that particular soil types have moisture-density relationships that differ considerably and cannot be represented on one general family of curves; in this case a separate family may be developed. Also, moisture-density relationships for material of widely varying geologic origins should be carefully examined to determine if separate families are required.

When a small number of moisture-density relationships are being used to develop a family of curves, plot the point representing the maximum density and optimum moisture content for each relationship on a single sheet of graph paper. Draw a smooth curve which as closely as possible connects all these points. This line will define the maximum density and optimum moisture content of the material represented by this family of curves. At 2-lb (1-kg) increments draw moisture-density curves with slopes similar to the slopes of the original moisture-density relationships. Slopes should gradually steepen going from low to high maximum density material.

When a great number of moisture-density relationships are available, the above procedure can be modified by using average values. Tabulate the maximum density, optimum moisture content, and slope for all moisture-density relationships in each 2-lb (1-kg) increment of density. Average the maximum densities and optimum moisture contents for each increment and plot these values. As before, draw a smooth curve which as closely as possible connects all these points. Determine the average slope for each increment, and at each 2-lb (1-kg) increment draw a moisture-density curve using this average slope value. A computer, if available, may be used to accomplish this work.

The accuracy of a family of curves can be checked by comparing the maximum density and optimum moisture content from an individual moisture-density relationship with that obtained using the One-Point Method and family of curves. A point representing 80 percent of optimum moisture content is taken from the individual moisture-density relationship and used as described in the One-Point Method to determine the maximum density and optimum moisture content from the family of curves. These values are compared with the values from the individual moisture-density relationship. The difference represents the maximum variance expected when the One-Point Method and family of curves are used for material represented by that individual moisture-density relationship. This comparison should be made for all types of material over the mass range of the family. Based on these results some adjustments may be necessary to the family and/or it may be recognized that the family is not applicable to some types of material. Families based on relatively few moisture-density relationships will generally require the closest scrutiny since it can be expected that a larger number of relationships will give better average conditions.
Performance Exam Checklist

*Family of Curves — One-point Method
FOP for AASHTO T 272*

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Procedure Element

<table>
<thead>
<tr>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. The tester has a copy of the current procedure on hand?

2. One-point determination of dry density and corresponding moisture content made in accordance with the FOP for AASHTO T 99, or AASHTO T 180?
   a. Correct size mold used?
   b. Correct number of blows per layer used (25 or 56)?
   c. Correct number of layers used (3, 4, or 5)?
   d. Moisture content determined in accordance with FOP for AASHTO T255/T265 or AASHTO T 217?

3. One-point plotted on family of curves supplied?

4. One-point falls within 80 to 100 percent of optimum moisture content in order to be valid?

5. If one-point does not fall within 80 to 100 percent of optimum moisture content, another one-point determination with an adjusted water content is made?

6. Maximum dry density and corresponding optimum moisture content correctly estimated?

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
### Performance Exam Checklist

#### AASHTO T 288 Checklist

**Determining Minimum Laboratory Soil Resistivity**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Laboratory method of Determining Minimum Resistivity</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Sample dried at 140 F, and screened through # 10 sieve?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Quartered or split out 1500 grams of passing #10 material?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. 150 ml of distilled water added to the 1500 gram and thoroughly mixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Sample covered with a wet cloth and allow to stabilize or cure for 12 hours?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Sample placed &amp; compacted in soil box in layers and the excess trimmed off with a straightedge?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Resistivity measured with the instrument?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Soil removed and retained from box and 100 ml of distilled water added and thoroughly mixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Soil box cleaned with distilled water?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. Repeat procedure by increasing moisture content by 100 ml until minimum resistivity can be established?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. Record the lowest value during the repeated measurements?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt:  Pass ☐  Fail ☐  Second attempt:  Pass ☐  Fail ☐

Signature of Examiner ________________________________

**Comments:**

________________________________________________________________________________________

________________________________________________________________________________________

________________________________________________________________________________________

________________________________________________________________________________________
WSDOT Test Method for AASHTO T 304

Uncompacted Void Content of Fine Aggregate

1. SCOPE

1.1. This method describes the determination of the loose uncompacted void content of a sample of fine aggregate. When measured on any aggregate of a known grading, void content provides an indication of that aggregate’s angularity, sphericity, and surface texture compared with other fine aggregates tested in the same grading. When void content is measured on an as-received fine aggregate grading, it can be an indicator of the effect of the fine aggregate on the workability of a mixture in which it may be used.

1.2. Three procedures are included for the measurement of void content. Two use graded fine aggregate (standard grading or as-received grading), and the other uses several individual size fractions for void content determinations:

1.2.1. Standard Graded Sample (Method A) – This method uses a standard fine aggregate grading that is obtained by combining individual sieve fractions from a typical fine aggregate sieve analysis. See the section on Preparation of Test Samples for the Grading.

Note WSDOT Specifications require Method A

1.2.2. Individual Size Fractions (Method B) – This method uses each of three fine aggregate size fractions: (a) 2.36-mm (No. 8) to 1.18-mm (No.16); (b) 1.18-mm (No.16) to 600-um (No. 30); and (c) 600-um (No.30) to 300 um (No. 50). For this method, each size is tested separately.

1.2.3. As-Received Grading (Method C) – This method uses that portion of the fine aggregate finer than a 4.75-mm (No. 4) sieve.

1.2.4. See the section on Significance and Use for guidance on the method to be used.

1.3. The values stated in English units shall be regarded as the standard.

1.4. This standard does not purport to address all of the safety problems, 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. REFERENCES DOCUMENTS

AASHTO Standards

T 2, WSDOT FOP for AASHTO for the Sampling of Aggregates
T 248, WSDOT FOP for AASHTO for Reducing Field Samples of Aggregates to Testing Size
T 27/11, WSDOT FOP for WAQTC /AASHTO for the Sieve Analysis of Fine & Coarse Aggregates
T 84 Specific Gravity and Absorption of Fine Aggregate

1This test method is based on AASHTO T304-96 (2000).
2.1. ASTM Standards:
   • B 88, Specification for Seamless Copper Water Tube
   • B 88M, Specification for Seamless Copper Water Tube (Metric)
   • C 29/29M, Test Method for Bulk Density ("Unit Weight") and Voids in Aggregate
   • C 117, Test Method for Materials Finer than 75-um (No. 200) Sieve in Mineral Aggregates by Washing
   • C 125, Terminology Relating to Concrete and Concrete Aggregates
   • C 128, Test Method for Specific Gravity and Absorption of Fine Aggregate
   • C 136, Test Method for Sieve Analysis of Fine and Coarse Aggregates
   • C 702, Practice for Reducing Samples of Aggregate to Testing Size
   • C 778, Specification for Standard Sand
   • D 75, Practice for Sampling Aggregates

2.2. ACI Document:
   • ACI 116R, Cement and Concrete Terminology

3. TERMINOLOGY
3.1. Terms used in this standard are defined in ASTM C 125 or ACI 116R.

4. SUMMARY OF TEST METHOD
4.1. A nominal 100-mL calibrated cylindrical measure is filled with fine aggregate of prescribed grading by allowing the sample to flow through a funnel from a fixed height into the measure. The fine aggregate is struck off, and its mass is determined by weighing. Uncompacted void content is calculated as the difference between the volume of the cylindrical measure and the absolute volume of the fine aggregate collected in the measure. Uncompacted void content is calculated using the bulk dry specific gravity of the fine aggregate. Two runs are made on each sample and the results are averaged.

4.1.1. For a graded sample (Method A or Method C), the percent void content is determined directly, and the average value from two runs is reported.

4.1.2. For the individual size fractions (Method B), the mean percent void content is calculated using the results from tests of each of the three individual size fractions.

5. SIGNIFICANCE AND USE
5.1. Methods A and B provide percent void content determined under standardized conditions which depend on the particle shape and texture of a fine aggregate. An increase in void content by these procedures indicates greater angularity, less sphericity, or rougher surface texture, or some combination of the three factors. A decrease in void content results is associated with more rounded, spherical, smooth surfaced fine aggregate, or a combination of these factors.

5.2. Method C measures the uncompacted void content of the minus No. 4 (4.75-mm) portion of the as-received material. This void content depends on grading as well as particle shape and texture.
5.3. The void content determined on the standard graded sample (Method A) is not directly comparable with the average void content of the three individual size fractions from the same sample tested separately (Method B). A sample consisting of single size particles will have a higher void content than a graded sample. Therefore, use either one method or the other as a comparative measure of shape and texture, and identify which method has been used to obtain the reported data. Method C does not provide an indication of shape and texture directly if the grading from sample to sample changes.

5.3.1. The standard graded sample (Method A) is most useful as a quick test which indicates the particle shape properties of a graded fine aggregate. Typically, the material used to make up the standard graded sample can be obtained from the remaining size fractions after performing a single sieve analysis of the fine aggregate.

5.3.2. Obtaining and testing individual size fractions (Method B) is more time consuming and requires a larger initial sample than using the graded sample. However, Method B provides additional information concerning the shape and texture characteristics of individual sizes.

5.3.3. Testing samples in the as-received grading (Method C) may be useful in selecting proportions of components used in a variety of mixtures. In general, high void content suggests that the material could be improved by providing additional fines in the fine aggregate or more cementitious material may be needed to fill voids between particles.

5.3.4. The bulk dry specific gravity of the fine aggregate is used in calculating the void content. The effectiveness of these methods of determining void content and its relationship to particle shape and texture depends on the bulk specific gravity of the various size fractions being equal, or nearly so. The void content is actually a function of the volume of each size fraction. If the type of rock or minerals, or its porosity, in any of the size fractions varies markedly it may be necessary to determine the specific gravity of the size fractions used in the test.

5.4. Void content information from Methods A, B, or C will be useful as an indicator of properties such as: the mixing water demand of hydraulic cement concrete; flowability, pumpability, or workability factors when formulating grouts or mortars; or, in bituminous concrete, the effect of the fine aggregate on stability and voids in the mineral aggregate; or the stability of the fine aggregate portion of a base course aggregate.

6. APPARATUS

6.1. Cylindrical Measure – A right cylinder of approximately 100 mL capacity having an inside diameter of approximately 39 mm and an inside height of approximately 86 mm made of drawn copper water tube meeting ASTM Specification B 88 Type M, or B 88 M Type C. The bottom of the measure shall be metal at least 6 mm thick, shall be firmly sealed to the tubing, and shall be provided with means for aligning the axis of the cylinder with that of the funnel. (See Figure 1.)

6.2. Funnel – The lateral surface of the right frustum of a cone sloped 60 ± 4º from the horizontal with an opening of 12.7 ± 0.6 mm diameter. The funnel section shall be a piece of metal, smooth on the inside and at least 38 mm high. It shall have a volume of at least 200 mL or shall be provided with a supplemental glass or metal container to provide the required volume. (See Figure 2.)
Figure 1 – Nominal 100-ml Cylindrical Measure

Figure 2 – Suitable Funnel Stand Apparatus with Cylindrical Measure in Place
Note 1 – Pycnometer top C9455 sold by Hogentogler and Co., Inc., 9515 Gerwig, Columbia, MD 21045, 410-381-2390 is satisfactory for the funnel section, except that the size of the opening has to be enlarged and any burrs or lips that are apparent should be removed by light filing or sanding before use. This pycnometer top must be used with suitable glass jar with the bottom removed (Figure 2).

6.3. Funnel stand – A three or four legged support capable of holding the funnel firmly in position with the axis of the funnel colinear (within a 4° angle and a displacement of 2 mm) with the axis of the cylindrical measure. The funnel opening shall be 115 ± 2 mm above the top of the cylinder. A suitable arrangement is shown in Figure 2.

6.4. Glass Plate – A square glass plate approximately 60 mm by 60 mm with a minimum 4-mm thickness used to calibrate the cylindrical measure.

6.5. Pan – A metal or plastic pan of sufficient size to contain the funnel stand and to prevent loss of material. The purpose of the pan is to catch and retain fine aggregate particles that overflow the measure during filling and strike off.

6.6. Metal spatula with a blade approximately 100 mm long, and at least 20 mm wide, with straight edges. The end shall be cut at a right angle to the edges. The straight edges. The straight edge of the spatula blade is used to strike off the fine aggregate.

6.7. Scale or balance accurate and readable to ±0.1 g within the range of use, capable of weighing the cylindrical measure and its contents.

7. SAMPLING

7.1. The sample(s) used for this test shall be obtained using FOP for AASHTO T 2 ASTM D 75 and FOP for AASHTO T 248 ASTM C 702, or from sieve analysis samples used for FOP for WAQTC/AASHTO T 27/11 ASTM C 136, or from aggregate extracted from a bituminous concrete specimen. For Methods A and B, the sample is washed over a 150-um (No. 100) or 75-um (No. 200) sieve in accordance with FOP for WAQTC/AASHTO T 27/11 ASTM C 136 and then dried and sieved into separate size fractions according to FOP for WAQTC/AASHTO T 27/11 ASTM C 136 procedures. Maintain the necessary size fractions obtained from one (or more) sieve analysis in a dry condition in separate containers for each size. For Method C, dry a split of the as-received sample in accordance with the drying procedure in FOP for AASHTO T 27/11 ASTM C 136.

8. CALIBRATION OF CYLINDRICAL MEASURE

8.1. Apply a light coat of grease to the top edge of the dry, empty cylindrical measure. Weigh the measure, grease, and glass plate. Fill the measure freshly boiled, deionized water at a temperature of 18 to 24°C. Record the temperature of the water. Place the glass plate on the measure, being sure that no air bubbles remain. Dry the outer surfaces of the measure and determine the combined mass of measure, glass plate, grease, and water by weighing. Following the final weighing, remove the grease, and determine the mass of the clean, dry, empty measure for subsequent test.
8.2. Calculate the volume of the measure as follows:

\[ V = \frac{1000 M}{D} \]

where:

- \( V \) = volume of cylinder, mL,
- \( M \) = net mass of water, g, and
- \( D \) = density of water (see table in ASTM C 29/C 29M for density at the temperature used), Kg/m³.

Determine the volume to the nearest 0.1 mL.

**Note 2** – If the volume of the measure is greater than 100.0 mL, it may be desirable to grind the upper edge of the cylinder until the volume is exactly 100.0 mL, to simplify subsequent calculations.

9. PREPARATION OF TEST SAMPLES

9.1. *Method A – Standard Graded Sample* – Weigh out and combine the following quantities of fine aggregate which has been dried and sieved in accordance with FOP for AASHTO T 27/11 ASTM C 136

<table>
<thead>
<tr>
<th>Individual Size Fraction</th>
<th>Mass, g</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 8 (2.36mm) to No. 16 (1.18 mm)</td>
<td>44</td>
</tr>
<tr>
<td>No. 16 (1.18 mm) to No. 30 (600 um)</td>
<td>57</td>
</tr>
<tr>
<td>No. 30 (600 um) to No. 50 (300 um)</td>
<td>72</td>
</tr>
<tr>
<td>No. 50 (300 um) to No. 100 (150 um)</td>
<td>17</td>
</tr>
<tr>
<td>Total</td>
<td>190</td>
</tr>
</tbody>
</table>

The tolerance on each of these amounts is ±0.2 g.


WSDOT has deleted this section they use Method A

9.3. *Method C – As Received Grading*:

WSDOT has deleted this section they use Method A

9.4. Specific Gravity of Fine Aggregate—If the bulk dry specific gravity of fine aggregate from the source is unknown, determine it on the minus No. 4 (4.75 mm) material according to AASHTO T 84 ASTM C 128. Need to add SG from Mix Design. Use this value in subsequent calculations unless some size fractions differ by more than 0.05 from the specific gravity typical of the complete sample, in which case the specific gravity of the fraction (or fractions) being tested must be determined. An indicator of differences in specific gravity of various particle sizes is a comparison of specific gravities run on the fine aggregate in different gradings. Specific gravity can be run on gradings with and without specific size fractions of interest. If specific gravity differences exceed 0.05, determine the specific gravity of the individual 2.36 mm (No. 8) to 150 um (No. 100) sizes for use with Method A or the individual size fractions for use with Method B either by direct measurement or by calculation using the specific gravity data on gradings with and without the size fraction of interest. A difference in specific gravity of 0.05 will change the calculated void content about one percent.
10. Procedure

10.1. Mix each test sample with the spatula until it appears to be homogeneous. Position the jar and funnel section in the stand and center the cylindrical measure as shown in Figure 2. Use a finger to block the opening of the funnel. Pour the test sample into the funnel. Level the material in the funnel with the spatula. Remove the finger and allow the sample to fall freely into the cylindrical measure.

10.2. After the funnel empties, strike-off excess heaped fine aggregate from the cylindrical measure by a single pass of the spatula with the width of the blade vertical using the straight part of its edge in light contact with the top of the measure. Until this operation is complete, exercise care to avoid vibration or any disturbance that could cause compaction of the fine aggregate in the cylindrical measure. (Note 3) Brush adhering grains from the outside of the container and determine the mass of the cylindrical measure and contents to the nearest 0.1 g. Retain all fine aggregate particles for a second test run.

Note 3 – After strike-off, the cylindrical measure may be tapped lightly to compact the sample to make it easier to transfer the container to scale or balance without spilling any of the sample.

10.3. Recombine the sample from the retaining pan and cylindrical measure and repeat the procedure. The results of two runs are averaged. See the Calculation section.

10.4. Record the mass of the empty measure. Also, for each run, record the mass of the measure and fine aggregate.

11. Calculation

11.1. Calculate the uncompacted voids for each determination as follows:

\[ U = \frac{V - (F/G)}{V} \times 100 \]

\[ V = \text{volume of cylindrical measure, mL;} \]

\[ F = \text{net mass, g, of fine aggregate in measure (gross mass minus the mass of the empty measure);} \]

\[ G = \text{Bulk dry specific gravity of fine aggregate; and} \]

\[ U = \text{uncompacted voids, percent, in the material.} \]

11.2. For the standard Graded Sample (Method A) calculate the average uncompacted voids for the two determinations and report the result as \( U_s \).

11.3. For the Individual Size Fractions (Method B) calculate:

11.3.1. First, the average uncompacted voids for the determination made on each of the three-size-fraction samples:

\[ U_1 = \text{Uncompacted Voids, No. 8 (2.36 mm) to No. 16 (1.18 mm), percent;} \]

\[ U_2 = \text{Uncompacted Voids, No. 16 (1.18 mm) to No. 30 (600 um), percent;} \]

\[ U_3 = \text{Uncompacted Voids, No. 30 (600 um) to No. 50 (300 um), percent;} \]

11.3.2. Second, the mean uncompacted voids (\( U_m \)) including the results for all three sizes:

\[ U_m = \frac{(U_1 + U_2 + U_3)}{3} \]

11.4. For the As-Received grading (Method C) calculate the average uncompacted voids for the two determinations and report the result as \( U_r \).
12. REPORT

12.1 For the Standard Graded Sample (Method A) report:

12.1.1. The Uncompacted Voids ($U_0$) in percent to the nearest one-tenth of a percent (0.1%).

12.1.2. The specific gravity value used in the calculations.

12.2 For the Individual Size Fractions (Method B) report the following percent voids to the nearest one-tenth of a percent (0.1%):

12.2.1. Uncompacted Voids for size fractions: (a) No. 8 (2.36 mm) to No. 16 (1.18 mm) ($U_1$); (b) No. 16 (1.18 mm) to No. 30 (600 um) ($U_2$); and (c) No. 30 (600 um) to No. 50 (300 um) ($U_3$).

12.2.2. Mean Uncompacted Voids ($U_m$).

12.2.3. Specific gravity value(s) used in the calculations, and whether the specific gravity value(s) were determined on a graded sample or the individual size fractions used in the test.

12.3 For the As-Received Sample (Method C) report:

12.3.1. The uncompacted voids ($U_R$) in percent to the nearest one-tenth of a percent (0.1%).

12.3.2. The specific gravity value used in the calculation.

12.4 Report Results using WSDOT Form 350-161, or other report approved by the State Materials Engineer.

13. PRECISION AND BIAS

See AASHTO T 304 for Precision and bias

14. KEYWORDS

Angularity; fine aggregate; particle shape; sand; surface texture; void content.

Copies may be obtained from the American Concrete Institute, Box 19150, Detroit, MI 48219.
Performance Exam Checklist

**FOP AASHTO T-304**

**UNCOMPACTED VOID CONTENT OF FINE AGGREGATE**

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**SAMPLE PREPARATION** (Method A)

Note: If Bulk Dry Specific Gravity is unknown, determine it on the minus No. 4-(4.75 mm) material according to AASHTO T-84.

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Field sample obtained per FOP for AASHTO T-2?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Sample reduced to testing size per FOP for AASHTO T-248?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Sample washed over No. 100 or No. 200 sieve in accordance with FOP for WAQTC/AASHTO T-27/11?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Sample dried to constant weight?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Standard Graded sample achieved per FOP for WAQTC/AASHTO T-27/11?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Necessary size fractions obtained, maintained in a dry condition in separate containers for Each size?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Standard Graded sample-weighed out and combined per Section 9.1, FOP for AASHTO T-304?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Procedure Element

PROCEDURE (Method A)

Note: If Bulk Dry Specific Gravity is unknown, determine it on the minus No. 4- (4.75 mm) material according to AASHTO T-84.

1. Test sample mixed until it appears to be homogeneous?  
   - Yes □  No □

2. Jar and funnel section positioned in stand and cylindrical measure centered on stand?  
   - Yes □  No □

3. Finger used to block the opening of the funnel?  
   - Yes □  No □

4. Test sample poured into the funnel and leveled?  
   - Yes □  No □

5. Finger removed and sample allowed to fall freely into cylindrical measure?  
   - Yes □  No □

6. After funnel empties, is excess material struck off with single pass of upright spatula?  
   - Yes □  No □

7. Was care taken to avoid any vibration or disturbance that could cause compaction of material?  
   - Yes □  No □

8. All adhering grains brushed off before weighing the cylindrical measure?  
   - Yes □  No □

9. Mass of the cylindrical measure and contents weighed to nearest 0.1 gram?  
   - Yes □  No □

10. All fine aggregate particles retained and re-homogenized for a second test run?  
    - Yes □  No □

11. Percent (%) of Uncompacted Voids calculated for each run, as per FOP for AASHTO T-304, Method A?  
    - Yes □  No □

12. Were the results for each run averaged for a final result?  
    - Yes □  No □

13. Was the (%) percent of Uncompacted voids reported to the nearest one-tenth of a percent (0.1%)?  
    - Yes □  No □

14. All calculations performed correctly?  
    - Yes □  No □

First attempt:  Pass □  Fail □
Second attempt: Pass □  Fail □

Signature of Examiner __________________________________________

Comments:

____________________________________________________________________________________
____________________________________________________________________________________
____________________________________________________________________________________
____________________________________________________________________________________
AASHTO T 307 (Modified)
Determining the Resilient Modulus of Soils and Aggregate Materials

AASHTO T 307 has been adopted by WSDOT with the following changes and/or additions:

Section 3.3 and Section 3.4:
Any material that is cohesive enough that it can form a cylinder and remain intact through a Resilient Modulus test shall be considered Type 2 material. All other material shall be considered Type 1.

Section 3.8:
The Contact Load is 5.0 psi.

Section 6.1:
The LVDTs will be clamped to the center of the specimen.

Section 6.3.1:
The load cell will have a capacity of 1000 pound-force, and an accuracy of ± 2.5 1000 pound-force.

Section 6.3.3:
The LVDTs will be inside the test chamber.

Section 6.3.2 through Section 6.3.3:
Note 2 Following Section 6.3.1, Section 6.3.2, and Section 6.7:
The load cell will be calibrated by outside services and the LVDTs will be verified by WSDOT Verification Procedure VP-68.

Section 6.4:
Specimens will be compacted by impact or vibration.

Section 6.6:
Remove bubble chamber, nor a membrane expander, and porous stones nor porous bronze discs from list of equipment.

Section 7.1.1:
Use 4-inch diameter specimens for all types of materials. Particles retained on the 19 mm sieve will be scalped. The sample gradation will be per Appendix C of the WSDOT Lab. Manual.**
Section 7.3 Replace in its entirety with:

The target moisture content of the sample is to be the amount of moisture, which will cause exudation from the sample at 300 psi.

The target density of cohesive material is that produced by impact compaction per the section on compaction.

The target density of granular materials is that produced by vibratory compaction per the section on vibratory compaction.

Section 7.4:

See comments under Section 6.4.

Section 7.4.3:

Samples will be prepared and used the same day.

8. RESILIENT MODULUS SPECIMEN SET UP FOR SUBGRADE, AND BASE MATERIALS

Section 8.1.

WSDOT uses the Resilient Modulus test only for laboratory compacted samples. All samples will be approximately 4 inches in diameter.

Section 8.2.

Cohesive specimens are placed in the triaxial chamber and loading apparatus in the following steps:

Place a filter paper on the base, place the specimen on top of the filter paper, and put the load cell on top of the specimen.

Section 8.3.

Granular specimens are compacted on the base of the triaxial chamber inside a rubber membrane, with a vacuum applied to the membrane.

When compaction is finished, the vacuum line will be moved to produce a vacuum inside the membrane. The membrane is checked for leaks.

If leaks are found they can usually be sealed for the duration of the test with masking tape or the equivalent. If leaks cannot be sealed, remove the specimen from the rubber membrane, and prepare a new specimen using another rubber membrane.

Once a specimen has been prepared in an airtight membrane, the O-ring sealing the membrane at the base is checked. The load cell is placed on top of the specimen, and the membrane is extended over the load cell. An O-ring or other seal is placed on the load cell.

Section 8.4.

For both cohesive and granular specimens, the LVDTs will be placed at about the middle of the specimen, and adjusted so that the LVDTs make good contact with the clamps.

Section 8.5.

Place the chamber on the base plate and the cover plate on the chamber. Insert the loading piston and obtain a firm connection with the load cell. Tighten the chamber tie rods firmly.
Section 8.6.
Slide the assembly apparatus into position under the axial loading device. Position the piston rod precisely under the loading device. Apply a small (1 to 3 psi) contact stress to the test specimen then put the triaxial chamber under pressure.

Check that there are no air leaks. If air leaks from the edges of the chamber, the chamber must be disassembled, the edges cleaned, and possibly moistened. Repeat until chamber is airtight. (A very small amount of air leak around the piston rod is acceptable.)

Section 8.7.
Connect the air pressure supply line to the triaxial chamber and apply the pre-conditioning confining pressure of 4 psi to the test specimen. Raise the contact stress to 5 psi.

9. RESILIENT MODULUS TEST PROCEDURE- FOR SUBGRADE, AND BASE MATERIALS

Section 8.8.
Begin the test by sample conditioning with a minimum of 1000 repetitions of a load equivalent to a cyclic Stress of 8 psi.

If the total vertical permanent strain exceeds 5 percent during conditioning, stop the test and report the result on the appropriate worksheet. Recreate the sample, using extra care to assure adequate compaction. If the sample again reaches 5 percent total vertical strain during conditioning terminate the test and report on the worksheet.

Apply 200 load applications between each item on the testing sequence. Record the average recovered deformations for each LVDT separately for one cycle on the report form.

Modified table 5 and 6 are used for the conditioning and testing sequences. **

If at any time the total vertical permanent strain exceeds 5 percent, stop the test and report the result on the appropriate worksheet.

10. CALCULATIONS
Use the WSDOT computer generated stress-strain curve for the specimen.

11 REPORT
Fill out the RESILIENT MODULUS WORKSHEET, and the SOILS Software Worksheet.
The report will be generated by the SOILS Software.
Appendix A SAMPLE PREPARATION

Section A.1.2.2. and A.1.2.2.3.

Omit

Section A.1.1.2.7

Cure sample only if necessary according the judgment of the operator.

Appendix B VIBRATORY COMPACTION

Section B.2.3:

The compactor head diameter is approximately 50mm.

Section B.3.2.

Omit

Section B.3.8. through Section B.3.10

Omit

Section B.3.14.

Insert the vibrator and vibrate the soil, being careful that the only pressure applied is from the weight of the vibrator. Vibrate for approximately one minute, moving vibrator head to different parts of the layer.

Appendix C COMPACTION OF TYPE 2 SOILS

Type 2 soils are compacted as for T-99, except that the specimen shall be compacted in eight lifts to a height of eight inches.
WSDOT FOP for AASHTO T 308

Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method

1. SCOPE

1.1 This test method covers the determination of asphalt binder content of HMA mixtures by ignition at temperatures that reach the flashpoint of the binder in a furnace. The means of sample heating may be the convection method or the direct infrared (IR) irradiation method. The aggregate remaining after burning can be used for sieve analysis using FOP for WAQTC/AASHTO T 27/T11.

1.2 The values in English units are to be regarded as the standard.

1.3 This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety problems associated with its use. It is the responsibility of whoever uses this standard to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards

<table>
<thead>
<tr>
<th></th>
<th>AASHTO Standards</th>
</tr>
</thead>
<tbody>
<tr>
<td>M 231</td>
<td>Weighing Devices Used in the Testing of Materials</td>
</tr>
<tr>
<td>T 2</td>
<td>Sampling of Aggregates</td>
</tr>
<tr>
<td>T 30</td>
<td>Mechanical Analysis of Extracted Aggregate</td>
</tr>
<tr>
<td>T 40</td>
<td>Sampling Bituminous Materials</td>
</tr>
<tr>
<td>T 110</td>
<td>Moisture or Volatile Distillates in Hot-Mix Asphalt (HMA)</td>
</tr>
<tr>
<td>T 168</td>
<td>Sampling Bituminous Paving Mixtures</td>
</tr>
<tr>
<td>T 248</td>
<td>Reducing Samples of Aggregate to Testing Size</td>
</tr>
</tbody>
</table>

2.2 Manufacturer’s Instruction Manual

2.3 WSDOT Standards

<table>
<thead>
<tr>
<th></th>
<th>WSDOT Standards</th>
</tr>
</thead>
<tbody>
<tr>
<td>FOP for AASHTO T 329</td>
<td>Moisture Content of Asphalt (HMA) by Oven</td>
</tr>
<tr>
<td>FOP for WAQTC/AASHTO T 168</td>
<td>Sampling Bituminous Paving Materials</td>
</tr>
<tr>
<td>WSDOT T 712</td>
<td>Reducing Samples of Hot Mix Asphalt to Testing Size</td>
</tr>
<tr>
<td>SOP 728</td>
<td>Method for Determining Ignition Furnace Calibration Factor</td>
</tr>
</tbody>
</table>

3. SUMMARY OF TEST METHOD

3.1 The asphalt binder in the paving mixture is ignited using the furnace equipment applicable to the particular method. The asphalt binder content is calculated as the difference between the initial mass of the asphalt mixture and the mass of the HMA residual aggregate, with adjustments for the calibration factor, and the moisture content. The asphalt content is expressed as mass percent of moisture-free mixture.

This FOP is based on AASHTO T 308-05 and has been modified per WSDOT standards. To view the redline modifications, contact WSDOT Quality Systems Manager at (360) 709-5497.
4. SIGNIFICANCE AND USE

4.1 This method can be used for quantitative determinations of asphalt binder content and gradation in HMA mixtures and pavement samples for quality control, specification acceptance, and mixture evaluation studies. This method does not require the use of solvents. Aggregate obtained by this test method may be used for gradation analysis according to FOP for WAQTC/AASHTO T 27/11.

5. SAMPLING

5.1 Obtain samples of aggregate in accordance with T 2. or

5.2 Obtain samples of asphalt binder in accordance with T 40. or

5.3 Obtain samples of freshly produced hot-mix asphalt in accordance with FOP for WAQTC/ AASHTO T 168.

5.4 The test specimen shall be the end result of quartering a larger sample taken in accordance with T 248. The test specimen for asphalt content determination shall be the end result of a larger sample taken in accordance with FOP for WAQTC/AASHTO T 168.

5.5 If the mixture is not sufficiently soft to separate with a spatula or trowel, place it in a large flat pan in an oven at 12°C + 5°C (25°F + 9°F) until it is workable. If the mixture is not sufficiently soft to separate for testing, carefully heat the mixture in an oven until sufficiently soft, not to exceed 350°F or the recommended mixing temperature from the mix design verification report. Do not leave the sample in the oven for an extended period of time.

5.6 The size of the test sample shall be governed by the nominal maximum aggregate size of the mixture and shall conform to the mass requirement shown in Table 1. When the mass of the test specimen exceeds the capacity of the equipment used, the test specimen may be divided into suitable increments, tested, and the results appropriately combined for calculation of the asphalt binder content (weighted average). Specimen sizes shall not be more than 500 g greater than the minimum recommended specimen mass. The maximum sample size including basket shall not exceed the capacity of the balance.

**Note 1:** Large samples of fine mixes tend to result in incomplete ignition of asphalt binder.

<table>
<thead>
<tr>
<th>Nominal Max. Agg. Size</th>
<th>Class of HMA</th>
<th>Minimum Mass of Specimen, g</th>
<th>Maximum Mass of Specimen, g</th>
</tr>
</thead>
<tbody>
<tr>
<td>US No. 4</td>
<td>Superpave</td>
<td>1200</td>
<td>1700</td>
</tr>
<tr>
<td>3/8 in.</td>
<td>3/8 In.</td>
<td>Class G &amp; D</td>
<td>1200</td>
</tr>
<tr>
<td>1/2 in.</td>
<td>1/2 In.</td>
<td>Class A, B, &amp; ATB</td>
<td>1500</td>
</tr>
<tr>
<td>3/4 in.</td>
<td>3/4 In.</td>
<td>2000</td>
<td>2500</td>
</tr>
<tr>
<td>1 in.</td>
<td>1 In.</td>
<td>Class E</td>
<td>3000</td>
</tr>
<tr>
<td>1 1/2 in.</td>
<td></td>
<td>4000</td>
<td>4500</td>
</tr>
</tbody>
</table>

* For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.
Note: For an aggregate specification having a generally unrestricted gradation (i.e. wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.

6. CALIBRATION

6.1 Note 2 and this section has been replaced with the following:

The Calibration is to be performed according to WSDOT Standard Operating Procedure SOP 728.

Test Method A

7. APPARATUS

7.1 Ignition Furnace — A forced air ignition furnace that heats the samples by either convection method or direct IR direct irradiation method. The convection-type furnace must be capable of maintaining the temperature at 1072°F (578°C). The furnace shall have an internal balance thermally isolated from the furnace chamber accurate to 0.1 g. The balance shall be capable of weighing a 3500 gram sample in addition to the sample baskets. A data collection system will be included so that the weight can be automatically determined and displayed during the test. The furnace shall have a built in computer program to calculate change in mass of the sample baskets and provide for the input of a correction factor for aggregate loss. The furnace shall provide a printed ticket with the initial specimen mass, specimen mass loss, temperature compensation, correction factor, corrected asphalt content (percent), test time, and test temperature. The furnace chamber dimensions shall be adequate to accommodate a sample size of 3500 grams. The furnace shall provide an audible alarm and indicator light when the sample mass loss does not exceed 0.0 percent of the total sample mass for three consecutive minutes. The furnace door shall be equipped so that the door cannot be opened during the ignition test. A method for reducing furnace emissions shall be provided. The furnace shall be vented into a hood or to the outside and, when set up properly, shall have no noticeable odors escaping into the laboratory. The furnace shall have a fan with capability to pull air through the furnace to expedite the test and to reduce the escape of smoke into the laboratory.

Note 3: The furnace shall also allow the operator to change the ending mass loss percentage to 0.02 percent, WSDOT uses 0.01%.

7.2 Sample Basket(s) — of appropriate size that allows the sample(s) to be thinly spread and allows air to flow through and around the sample particles. Sets with two or more baskets shall be nested. The sample shall be completely enclosed with screen mesh, perforated stainless steel plate, or other suitable material.

7.2.1 Sample Basket Assembly — consisting of sample basket(s) (7.2), catch pan (7.3), and an assembly guard to secure sample basket(s) to catch pan.

Note 4: Screen mesh or other suitable material with maximum and minimum opening of No. 8 (2.36 mm) and No. 30 (600 microns) respectively has been found to perform well.

7.3 Catch Pan — of sufficient size to hold the sample basket(s) so that aggregate particles and melting asphalt binder falling through the screen mesh are caught.

7.4 Oven or suitable devise — Capable of maintaining 325 ± 25°F (163 ± 14°C).

7.5 Balance — of sufficient capacity and conforming to the requirements of M231, Class G2, for weighting specimen in basket(s).
7.6 Safety Equipment — safety glasses or face shield, high temperature gloves, long sleeve jacket, a heat resistant surface capable of withstanding 1202°F (650°C) and a protective cage capable of surrounding the sample baskets during the cooling period.

7.7 Miscellaneous Equipment — a pan larger than the sample basket(s) for transferring sample after ignition, spatulas, bowls, and wire brushes.

8. TEST PROCEDURES

8.1 Test Initiation

8.1.1 For the convection-type furnace, Preheat the ignition furnace to 538°C (1000°F) or as determined in Section 6.9. Manually record the furnace temperature (set point) prior to the initiation of the test if the furnace does not record automatically. Preheat the ignition furnace to 1000°F (538°C). Manually record the furnace temperature (set point) prior to the initiation of the test if the furnace does not record automatically.

8.1.2 For the direct irradiation-type furnace, use the same burn profile as used during the calibration.

8.2 Oven dry the HMA sample to a constant mass at a temperature of 105 ± 5°C (221 ± 9°F) or determine the moisture content of the samples according to T 0. Determine the moisture content of the samples according to FOP for AASHTO T 329.

8.3 Enter the calibration factor for the specific mix to be tested as determined in Section 6 in the ignition furnace.

8.4 Weigh and record the mass of the sample basket(s) and catch pan (with guards in place).

8.5 Prepare the sample as described in Section 5. Evenly distribute this sample in the sample basket(s) that have been placed in the catch pan, taking care to keep the material away from the edges of the basket. Use a spatula or trowel to level the specimen.

8.6 Weigh and record the total mass of the sample, basket(s), catch pan, and basket guards. Calculate and record the initial mass of the specimen (total mass minus the mass of the specimen basket assembly).

8.7 Input the initial mass of the specimen in whole grams into the ignition furnace controller. Verify that the correct mass has been entered.

8.8 Tare or zero furnace balance, open the chamber door, and gently set the sample baskets in the furnace. Close the chamber door, and verify that the sample mass (including the basket(s)) displayed on the furnace scale equals the total mass recorded in Section 8.6 within ±5 g ± 6 g. Differences greater than 6 g or failure of the furnace scale to stabilize may indicate that the sample basket(s) are contacting the furnace wall. Initiate the test by pressing the start/stop button. This will lock the sample chamber and start the combustion blower.

Note 5: The furnace temperature will drop below the setpoint when the door is opened, but will recover with the door closed and when ignition occurs. Sample ignition typically increases the temperature well above the setpoint, depending on sample size and asphalt content.

WSDOT Note: Operator should wear safety equipment – high temperature gloves, face shield, fire-retardant shop coat – when opening the door to load or unload the sample.

WSDOT Safety Note: Do not attempt to open the furnace door until the binder has been completely burned off.
8.9 Allow the test to continue until the stable light and audible stable indicator indicate the test is complete (the change in mass does not exceed 0.01 percent for three consecutive minutes). Press the start/stop button. This will unlock the sample chamber and cause the printer to print out the test results.

Note 6: An ending mass loss percentage of 0.02 may be substituted when aggregate that exhibits an excessive amount of loss during ignition testing is used. The precision and bias statement was developed using 0.01 percent. Both precision and accuracy may be adversely affected by using 0.02 percent.

8.10 Use the corrected asphalt binder content (0.01 percent) from the printed ticket. If a moisture content (0.01 percent) has been determined, subtract the percent moisture from the printed ticket corrected asphalt content, and report the resultant value as the corrected asphalt binder content to 0.1 percent.

8.11 Open the chamber door, remove the sample basket assembly and place on heat resistance surface and cover with the protective cage. Allow sample to cool to room temperature (approximately 30 minutes).

Test Method B

WSDOT does not use Method B and has deleted it from the procedure.

11. GRADATION

11.1 Allow the specimen to cool to room temperature in the sample baskets.

11.2 Empty the contents of the baskets into a suitable container. Use a small wire sieve brush to ensure that any residual fines are removed from the baskets.

11.3 Perform the gradation analysis according to FOP for WAQTC/AASHTO T-30 T 27/T11.

12. REPORT

12.1 Report the test method (A or B), corrected asphalt binder content, calibration factor, temperature compensation factor (if applicable), total percent loss, sample mass, moisture content (if determined) and the test temperature. Attach the original printed tickets to the report for units with internal balances.

12.2 The asphalt percentage and aggregate gradation shall be reported on WSDOT Form 350-560 or other report approved by the State Materials Engineer.

13. PRECISION AND BIAS

See AASHTO T-308 Precision and Bias
Performance Exam Checklist

Determining the Asphalt Cement Content of Hot Mix Asphalt (HMA) by the Ignition Method for AASHTO T 308

Participant Name ___________________________________________ Exam Date ______________

Procedure Element Yes No
1. The tester has a copy of the current procedure on hand? □ □
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? □ □

Procedure
1. Oven at correct temperature 538 C? □ □
2. Mass of sample baskets and catch pan recorded? □ □
3. Samples evenly distributed in basket? □ □
4. Mass of sample recorded? □ □

Method A
5. Enter calibration factor for specific mix design? □ □
6. Initial mass entered into furnace controller? □ □
7. Sample correctly placed into furnace? □ □
8. Test continued until stable indicator signals? □ □
9. Binder content obtained on printed ticket? □ □
10. Binder content corrected for moisture? □ □
11. All calculations performed correctly? □ □

First attempt: Pass □ Fail □ Second attempt: Pass □ Fail □

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
WSDOT FOP for AASHTO T 309¹

*Temperature of Freshly Mixed Portland Cement Concrete*

1. SCOPE

1.1 This test method covers the determination of temperature of freshly mixed portland cement concrete.

1.2 The values stated in English units are to be regarded separately as standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:

T 141 Sampling Freshly Mixed Concrete

2.2 ASTM Standards:

C 172 Practice for Sampling Freshly Mixed Concrete

3. SIGNIFICANCE AND USE

3.1 This test method provides a means for measuring the temperature of freshly mixed concrete. It may be used to verify conformance to a specified requirement for temperature of concrete.

3.2 Concrete containing aggregate of a nominal maximum size greater than 3 in. [75 mm] may require up to 20 min for the transfer of heat from aggregate to mortar. (See ACI Committee 207.1R Report.3)

4. APPARATUS

4.1 Container, shall be made of nonabsorptive material and large enough to provide at least 3 in. [75 mm] of concrete in all directions around the sensor of the temperature measuring device; concrete cover must also be at least three times the nominal maximum size of the coarse aggregate.

4.2 Temperature Measuring Device — The temperature measuring device shall be capable of measuring the temperature of the freshly mixed concrete to ±1°F (± 0.5°C) throughout the entire temperature range likely to be encountered in the fresh concrete. Liquid-in-glass thermometers having a range of 0 to 120°F (-18 to 49°C) are satisfactory. Other thermometers of the required accuracy, including the metal immersion type, are acceptable. Liquid-in-glass thermometers having a range of 0 to 120°F (-18 to 49°C) are satisfactory. Other thermometers of the required accuracy, including the metal immersion type, are acceptable.

4.3 Partial immersion liquid-in-glass thermometers (and possibly other types) shall have a permanent mark to which the device must be immersed without applying a correction factor.

¹This procedure is based on AASHTO T 309-05
4.4 Reference Temperature Measuring Device — The reference temperature measuring device shall be a liquid-in-glass thermometer readable to 0.5°F (0.2°C) that has been verified and calibrated. The calibration certificate or report indicating conformance to the requirements of ASTM E 77 shall be available for inspection.

5. CALIBRATION OF TEMPERATURE MEASURING DEVICE

5.1 Each temperature measuring device used for determining temperature of freshly mixed concrete shall be calibrated annually, or whenever there is a question of accuracy. This calibration shall be performed by comparing the readings of the temperature measuring device at two temperatures at least 27°F (15°C) apart.

5.2 Calibration of the temperature measuring devices may be made in oil or other suitable baths having uniform density if provision is made to:

5.2.1 Maintain the bath temperature constant within 0.5°F (0.2°C) during the period of the test.

5.2.2 Have both the temperature and reference temperature measuring devices maintained in the bath for a minimum of 5 min before reading temperatures.

5.2.3 Continuously circulate the bath liquid to provide a uniform temperature.

5.2.4 Slightly tap thermometers containing liquid to avoid adhesion of the liquid to the glass if the temperature exposure is being reduced.

6. SAMPLING CONCRETE

6.1 The temperature of freshly mixed concrete may be measured in the transporting equipment provided the sensor of the temperature measuring device has at least 3 in. [75 mm] of concrete cover in all directions around it.

6.2 Temperature of the freshly mixed concrete may be obtained following concrete placement using the forms as the container.

6.3 If the transporting equipment or placement forms are not used as the container, a sample shall be prepared as follows:

6.3.1 Immediately, prior to sampling the freshly mixed concrete, dampen (with water) the sample container.

6.3.2 Sample the freshly mixed concrete in accordance with Practice C 172, except that composite samples are not required if the only purpose for obtaining the sample is to determine temperature.

6.3.3 Place the freshly mixed concrete into the container.

6.3.4 When concrete contains a nominal maximum size of aggregate greater than 3 in. (75 mm), it may require 20 min before the temperature is stabilized after mixing.

7. PROCEDURE

7.1 Place the temperature measuring device in the freshly mixed concrete so that the temperature sensing portion is submerged a minimum of 3 in. (75 mm). Gently press the concrete around the temperature measuring device at the surface of the concrete so that ambient air temperature does not affect the reading.

7.2 Leave the temperature measuring device in the freshly mixed concrete for a minimum period of 2 min or until the temperature reading stabilizes, then read and record the temperature.
7.3 Complete the temperature measurement of the freshly mixed concrete within 5 min after obtaining the sample.

8. REPORT
8.1 Record the measured temperature of the freshly mixed concrete to the nearest 1°F (0.5°C).
8.2 Report results on concrete delivery ticket (i.e., Certificate of Compliance).
8.3 The signature of the tester who performed the field acceptance test is required on concrete delivery tickets containing test results.

9. PRECISION AND BIAS
9.1 The precision and bias of this test method have not been determined. A precision and bias statement will be included when sufficient test data have been obtained and analyzed.
Performance Exam Checklist

**Temperature of Freshly Mixed Concrete**
**FOP for AASHTO T-309**

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First attempt:  Pass ☐  Fail ☐
Second attempt: Pass ☐  Fail ☐

Signature of Examiner __________________________________________

Comments:

_________________________________________________________________
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_________________________________________________________________
WSDOT FOP for AASHTO T 310
In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)

1. SCOPE

1.1 This test method describes the procedure for determining the in-place density and moisture of soil and soil-aggregate by use of nuclear equipment. The density of the material may be determined by either direct transmission, backscatter, or backscatter/air-gap ratio method. The WSDOT standard method for determining density is by direct transmission.

1.2 Density — The total or wet density of soil and soil-rock mixtures is determined by the attenuation of gamma radiation where the source or detector is placed at a known depth up to 300 mm (12 in.) while the detector(s) or source remains on the surface (Direct Transmission Method) or the source and detector(s) remain on the surface (Backscatter Method).

1.2.1 The density in mass per unit volume of the material under test is determined by comparing the detected rate of gamma radiation with previously established calibration data.

1.3 Moisture — The moisture content of the soil and soil-rock mixtures is determined by thermalization or slowing of fast neutrons where the neutron source and the thermal neutron detector both remain at the surface.

1.3.1 The water content in mass per unit volume of the material under test is determined by comparing the detection rate of thermalized or slow neutrons with previously established calibration data.

1.4 SI Units — The values stated in SI units are to be regarded as the standard.

1.5 This standard does not purport to address all of the safety problems, 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. See Section 6. Hazards.

2. REFERENCED DOCUMENTS

2.1 AASHTO Standards:

T 99 Moisture-Density Relations of Soils Using a 2.5-kg (5.5-lb) Rammer and a 305-mm (12-in.) Drop

T 180 Moisture-Density Relations of Soils Using a 4.54-kg (10-lb) Rammer and a 457-mm (18-in.) Drop

T 191 Density of Soil In-Place by the Sand-Cone Method

T 217 Determination of Moisture in Soils by Means of a Calcium Carbide Gas Pressure Moisture Tester

T 224 Correction for Coarse Particles in the Soil Compaction Test

This FOP is based on AASHTO T 310-03 and has been modified per WSDOT standards. To view the redline modifications, contact WSDOT Quality Systems Manager at (360) 709-5411.
2.2 **ASTM Test Method:**

- D 2216, Laboratory Determination of Moisture Content of Soil
- D 2487, Classification of Soils for Engineering Purposes (Unified Soil Classification System)
- D 2488, Description and Identification for Soils (Visual-Manual Procedure)
- D 2937, Density of Soil in Place by the Drive-Cylinder Method
- D 4253, Maximum Index Density and Unit Weight of Soils Using a Vibratory Table
- D 4254, Maximum Index Density and Unit Weight of Soils and Calculation of Relative Density

**WSDOT Standards:**

- T 606 Method of Test for Compaction Control of Granular Materials
- SOP 615 Determination of the % Compaction for Embankment & Untreated Surfacing Materials using the Nuclear Moisture-Density Gauge

3. **SIGNIFICANCE**

3.1 The test method described is useful as a rapid, nondestructive technique for the in-place determination of the wet density and water content of soil and soil-aggregate.

3.2 The test method is used for quality control and acceptance testing of compacted soil and rock for construction and for research and development. The non-destructive nature allows repetitive measurements at a single test location and statistical analysis of the results.

3.3 **Density** — The fundamental assumptions inherent in the methods are that Compton scattering is the dominant interaction and that the material under test is homogeneous.

3.4 **Moisture** — The fundamental assumptions inherent in the test method are that the hydrogen present is in the form of water as defined by ASTM D 2216, and that the material under test is homogeneous.

3.5 Test results may be affected by chemical composition, sample heterogeneity, and, to a lesser degree, material density and the surface texture of the material being tested. The technique also exhibits spatial bias in that the gauge is more sensitive to water contained in the material in close proximity to the surface and less sensitive to water at deeper levels.
4. INTERFERENCES

4.1 In-Place Density Interferences

4.1.1 The chemical composition of the sample may affect the measurement, and adjustments may be necessary.

4.1.2 The gauge is more sensitive to the density of the material in close proximity to the surface in the Backscatter Method.

**Note 1:** The nuclear gauge density measurements are somewhat biased to the surface layers of the soil being tested. This bias has largely been corrected out of the Direct Transmission Method and any remaining bias is insignificant. The Backscatter Method is still more sensitive to the material within the first several inches from the surface. Density measurements with direct transmission is the WSDOT standard method.

4.1.3 Oversize rocks or large voids in the source-detector path may cause higher or lower density determination. Since there is lack of uniformity in the soil due to layering, rock or voids, the test site beneath the gauge will be excavated and a representative sample will be taken to determine the gradation per WSDOT SOP 615.

4.1.5 Keep all other radioactive sources at least the minimum distance recommended by the manufacture away from the gauge to avoid affecting the measurement.

4.2 In-Place Moisture Content Interferences

4.2.1 The chemical composition of the sample may dramatically affect the measurement and adjustments may be necessary. Hydrogen in forms other than water, as defined by ASTM D 2216, and carbon will cause measurements in excess of the true value. Some chemical elements such as boron, chlorine, and minute quantities of cadmium will cause measurements lower than the true value.

4.2.2 The water content determined by this test method is not necessarily the average water within the volume of the sample involved in the measurement. The measurement is heavily influenced by the water content of the material closest to the surface. The volume of soil and rock represented in the measurement is indeterminate and will vary with the water content of the material. In general, the greater the water content of the material, the smaller the volume involved in the measurement. At 10 lbs/ft.\(^3\) (160 kg/m\(^3\)), approximately 50 percent of the typical measurement results from the water content of the upper 2 to 3 in. (50 to 75 mm).

4.2.3 Keep all other neutron sources at least the minimum distance recommended by the manufacture away from the gauge to avoid affecting the measurement.
5. APPARATUS

5.1 Nuclear Density/Moisture Gauge — While exact details of construction of the gauge may vary, the system shall consist of:

5.1.1 A sealed source of high energy gamma radiation such as cesium or radium.

5.1.2 Gamma Detector — Any type of gamma detector such as a Geiger-Mueller tube(s).

5.2 Fast Neutron Source — A sealed mixture of a radioactive material such as americium, radium, or californium-252 and a target material such as beryllium.

5.3 Slow Neutron Detector — Any type of slow neutron detector such as boron trifluoride or helium-3 proportional counter.

5.4 Reference Standard — A block of material used for checking instrument operation, correction of source decay, and to establish conditions for a reproducible reference count rate.

5.5 Site Preparation Device — A plate, straightedge, or other suitable leveling tool which may be used for planing the test site to the required smoothness, and in the Direct Transmission Method, guiding the drive pin to prepare a perpendicular hole.

5.6 Drive Pin — A pin not to exceed the diameter of the rod in the Direct Transmission Gauge by more than ¼ in (6mm) or as recommended by the gauge manufacturer used to prepare a hole in the material under test for inserting the rod.

5.6.1 A slide hammer, with a drive pin attached, may also be used both to prepare a hole in the material to be tested and to extract the pin without distortion to the hole. In place of a slide hammer a hammer of significant size and weight for preparing a hole in the material to be tested using the drive pin along with an extraction tool.

5.7 Drive Pin Extractor — A tool that may be used to remove the drive pin in a vertical direction so that the pin will not distort the hole in the extraction process.

6. HAZARDS

6.1 This gauge utilizes radioactive materials that may be hazardous to the health of the users unless proper precautions are taken. Users of this gauge must become familiar with applicable safety procedures and government regulations.

6.2 Effective user instructions together with routine safety procedures, such as source leak tests, recording and evaluation of film badge data, etc., are a recommended part of the operation and storage of this gauge.

7. CALIBRATION

WSDOT has removed this section.
8. STANDARDIZATION

8.1 All nuclear density/moisture gauges are subject to long-term aging of the radioactive sources, detectors, and electronic systems, which may change the relationship between count rates and the material density and water content. To offset this aging, gauge are calibrated as a ratio of the measurement count rate to a count rate made on a reference standard or to an air-gap count (for the backscatter/air-gap ratio method). The reference count rate should be in the same or higher order of magnitude than the range of measurement count rates over the useful range of the gauge.

8.2 Standardization of equipment on the reference standard is required at the start of each day’s use and a permanent record of these data shall be retained. The standardization shall be performed with the equipment away from other radioactivity devices, large masses of water and large vertical surfaces at the manufacture’s recommended distance. Standard counts should be taken in the same environment as the actual measurement counts.

8.2.1 Turn on the gauge and allow for stabilization according to the manufacturer’s recommendations. If the gauge is to be used either continuously or intermittently during the day, it is best to leave it in the “power on” condition to prevent having to repeat the stabilization (refer to manufacturer recommendations). This will provide more stable, consistent results.

8.2.2 Using the reference standard block supplied with the density/moisture gauge perform standardization using manufacture’s recommendations.

Use the procedure recommended by the gauge manufacturer for determining compliance with the gauge calibration curves. Without specific recommendations for the gauge manufacturer, use the procedure in 8.2.3.

8.2.3 If the mean of the four repetitive readings is outside the limits set by Equation 1, repeat the standardization check. If the second standardization check satisfies Equation 1, the gauge is considered in satisfactory operating condition. If the second standardization check does not satisfy Equation 1, the gauge should be checked and verified according to Appendices A1 and A2, sections A1.8 and A2.5. If the verification shows that there is no significant change in the calibration curve, a new reference standard count, \( N_o \), should be established. If the verification check shows that there is a significant difference in the calibration curve, repair and recalibrate the gauge.

\[ N_s = N_o \pm 1.96 \sqrt{\frac{N_o}{F}} \]  
(Eq. 1)

where:

\( N_s \) = value of current standardization count,

\( N_o \) = Average of the past four values of \( N_s \), taken for prior usage, and

\( F \) = factory pre-scale factor (contact gauge manufacturer for the factor).
9. PROCEDURE

9.1 Turn on and allow the equipment to stabilize (warm up) according to the manufacturer’s recommendations (see 8.2.1). Prior to performing density test verify that today’s Standardization Count has been performed.

Select a test location per WSDOT SOP 615.

9.2 Prepare the test site in the following manner:

9.2.1 Remove all loose and disturbed material and additional material as necessary to expose the top of the material to be tested.

**Note 2:** The spatial bias should be considered in determining the depth at which the gauge is to be seated.

9.2.2 Select a horizontal area sufficient in size to accommodate four gauge readings that will be 90° to each other, by planing the area to a smooth condition so as to obtain maximum contact between the gauge and material being tested.

9.2.3 The maximum void beneath the gauge shall not exceed 1/8 in. (3 mm). Use native fines or fine sand to fill the voids and smooth the surface with a rigid plate or other suitable tool. The depth of the filler shall not exceed approximately 1/8 in. (3 mm).

9.4 This Section has been deleted because WSDOT does not use this method

9.5 Direct Transmission Method of In-Place Nuclear Density & Moisture Content

9.5.1 Select a test location where the gauge in test position will be at least the minimum distance recommended by the manufacture away from any vertical projection. If gauge will be within the minimum distance recommended by the manufacture follow instructions outlined by manufactures instruction manual.

9.5.2 Make a hole perpendicular to the prepared surface using the guide and the hole-forming device (Section 5). The hole shall be a minimum of 2 in. (50 mm) deeper than the desired measurement depth and of an alignment that insertion of the probe will not cause the gauge to tilt from the plane of the prepared area.

9.5.3 Mark the test area to allow the placement of the gauge over the test site and to allow the alignment of the source rod to the hole. Follow manufacturer recommendations if applicable.

WSDOT Note: For alignment purposes, the user may expose the source rod for a maximum of ten seconds.

9.5.4 Remove the hole forming device carefully to prevent the distortion of the hole, damage to the surface, or loose material to fall into the hole.

**Note:** If the hole cannot be maintained contact Regional Materials Laboratory for directions on how to proceed.

9.5.5 Place the instrument on the material to be tested, making sure of maximum surface contact as described above.

9.5.6 Lower the source rod into the hole to the desired test depth. Pull gently on the gauge in the direction that will bring the side of the probe to face the center of the gauge so that the probe is in intimate contact with the side of the hole in the gamma measurement path.
9.5.7 Select a test location(s) in accordance with the project specifications. Test sites should be relatively smooth and flat. The gauge should not be used within 1.5 ft. (450 mm) of any vertical mass. Keep all other radioactive sources at least the minimum distance recommended by the manufacture away from the gauge to avoid affecting the measurement. The test location should be at least 33 ft (10 m) away from other sources of radioactivity and at least 10 ft (3 m) away from large objects.

9.5.8 If the gauge is so equipped, set the depth selector to the same depth as the probe before recording the automated (gauge computed densities, moisture contents, and weights) values.

9.5.9 Secure and record one, one minute dry density and moisture content readings, then turn the gauge 90º and perform another set of readings. If the two dry density readings are not within 3 lbs/cf (50 kg/m³) of each other see note 5.

9.5.10

**Note 5:** If two readings are not within tolerances stated, rotate gauge 90º and retest. Again compare both readings. If these reading are still not within tolerances stated move to another location to perform test.

10. **CALCULATION OF RESULTS**

10.1 If dry density is required, the in-place water content may be determined by using the nuclear methods described herein; gravimetric samples and laboratory determination; or other approved instrumentation.

10.1.1 If the water content is determined by nuclear methods, use the gauge readings directly.

10.1.2 If the water content is determined by other methods, and is in the form of percent, proceed as follows:

\[
\frac{d}{m} = \frac{100}{100+W} (m) \tag{Eq. 2}
\]

where:

- \(d\) = dry density in lb/ft³ (kg/m³),
- \(m\) = wet density in lb/ft³ (kg/m³),
- \(W\) = water as a percent of dry mass.

10.2. Percent Compaction

WSDOT has deleted this section refer to WSDOT SOP 615 for determining the percent compaction.
11. REPORT
WSDOT has deleted this section refer to WSDOT SOP 615 for reporting.

12. Precision and Bias
This section has been deleted by WSDOT. Refer to AASHTO T310 for this information.

13. KEYWORDS
13.1 Compaction test; construction control; density; moisture content; nuclear methods; quality control; water content.

APPENDIX
A1. WET DENSITY CALIBRATION AND VERIFICATION
A2. WATER CONTENT CALIBRATION AND VERIFICATION
A3. GAUGE COUNT PRECISION
WSDOT has removed these section as WSDOT used manufacturer’s software to calibrate
Performance Exam Checklist

In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)

FOP FOR AASHTO T 310

Participant Name __________________________________________ Exam Date ____________________

**Procedure Element**

1. The tester has a copy of the current procedure on hand?  
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?  
3. Gauge turned on and allowed to stabilize per manufacturer’s recommendations?  
4. Gauge calibrated and standard count recorded in accordance with manufacturer’s instructions?  
5. Test location selected per WSDOT SOP 615?  
6. Loose, disturbed material removed?  
7. Flat, smooth area prepared?  
8. Surface voids filled with native fines (1/8 in. (3 mm) maximum thickness)?  
9. Hole driven 2 in. (50 mm) deeper than material to be tested?  
10. Gauge placed, probe placed, and source rod lowered without disturbing loose material?  
11. For alignment purposes, did not expose the source rod for more than 10 seconds.  
12. Method B:
   a. Gauge firmly seated, and gently pulled back so that source rod is against hole?  
   b. A one minute count taken; dry density and moisture data recorded?  
   c. Gauge turned 90° (180° in trench)?  
   d. Gauge firmly seated, and gently pulled back so that source rod is against hole?  
   e. A second one-minute count taken; dry density and moisture data recorded?  
   f. Density counts within 3 lb/ft³ (50 kg/m³)?  
   g. Average of two tests?  
13. A minimum 9 lbs. (4 kg) sample obtained from below gauge?  
14. Oversize determined following WSDOT SOP 615?  
15. All calculations performed correctly?  
16. Nuclear Gauge secured in a manner consistent with current DOH requirements?  

First attempt: Pass □ Fail □  Second attempt: Pass □ Fail □

Signature of Examiner __________________________________________
Preparing Hot-Mix Asphalt (HMA) Specimens by Means of the Superpave Gyratory Compactor

1. SCOPE

1.1. This standard covers the compaction of cylindrical specimens of hot-mix asphalt (HMA) using the Superpave gyratory compactor.

1.2. This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety problems 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. AASHTO Standards:

- M 231, Weighing Devices Used in Testing of Materials
- PP 28, Superpave Volumetric Design for Hot-Mix Asphalt (HMA)
- PP 48, Evaluation of the Superpave Gyratory Compactor (SGC) Internal Angle of Gyration
- R 30, Mixture Conditioning of Hot-Mix Asphalt (HMA)
- R 35, Superpave Volumetric Design for Hot-Mix Asphalt (HMA)
- T 166, Bulk Specific Gravity of Compacted Bituminous Mixtures Using Saturated Surface-Dry Specimens
- T 168, Sampling Bituminous Paving Mixtures
- T 209, Theoretical Maximum Specific Gravity and Density of Bituminous Paving Mixtures
- T 275, Bulk Specific Gravity of Compacted Bituminous Mixtures Using Paraffin-Coated Specimens
- T 316, Viscosity Determination of Asphalt Binder Using Rotational Viscometer

2.2. Other Standards:

- WSDOT SOP 731, Method for determining volumetric properties of asphalt concrete pavement class superpave
- WSDOT SOP 732, Superpave Volumetric Design for Hot-Mix Asphalt (HMA)

3. SIGNIFICANCE AND USE

3.1. This standard is used to prepare specimens for determining the mechanical and volumetric properties of HMA. The specimens simulate the density, aggregate orientation, and structural characteristics obtained in the actual roadway when proper construction procedure is used in the placement of the paving mix.

3.2. This test method may be used to monitor the density of test specimens during their preparation. It may also be used for field control of an HMA production process.

Based on AASHTO T 312-04
4. APPARATUS

4.1. Superpave Gyratory Compactor - An electrohydraulic or electromechanical compactor with a ram and ram heads as described in Section 4.3. The axis of the ram shall be perpendicular to the platen of the compactor. The ram shall apply and maintain a pressure of 600 ± 18 kPa perpendicular to the cylindrical axis of the specimen during compaction (Note 1). The compactor shall tilt the specimen molds at an angle of 1.25 ± 0.02º (22 ± 0.35 mrad) or an average internal angle of 1.16 ± 0.02º (20.2 ± 0.35 mrad), determined in accordance with AASHTO PP 48. The compactor shall gyrate the specimen molds at a rate of 30.0 ± 0.5 gyrations per minute throughout compaction.

Note 1—This stress calculates to 10,600 ± 310 N total force for 6 inches (150 mm) specimens.

4.1.1 Specimen Height Measurement and Recording Device - When specimen density is to be monitored during compaction, a means shall be provided to continuously measure and record the height of the specimen to the nearest 0.1 mm during compaction once per gyration.

4.2. Specimen Molds - Specimen molds shall have steel walls that are at least 0.3 inches (7.5 mm) thick and are hardened to at least a Rockwell hardness of C48. The initial inside finish of the molds shall have a root mean square (rms) of 1.60 um or smoother (Note 2). Molds shall have an inside diameter of 5.9 to 6.0 inches (149.90 to 150.00 mm) and be at least 9.8 inches (250 mm) high at room temperature.

Note 2—Smoothness measurement is in accordance with ANSI B 46.1. One source of supply for a surface comparator, which is used to verify the rms value of 1.60 um, is GAR Electroforming, Danbury, Connecticut.

4.3. Ram Heads and Mold Bottoms - Ram heads and mold bottoms shall be fabricated from steel with a minimum Rockwell hardness of C48. The ram heads shall stay perpendicular to its axis. The platen side of each mold bottom shall be flat and parallel to its face. All ram and base plate faces (the sides presented to the specimen) shall be flat to meet the smoothness requirement in Section 4.2 and shall have a diameter of 5.88 to 5.90 inches (149.50 to 149.75 mm).

4.4. Thermometers - Armored, glass, or dial-type thermometers with metal stems for determining the temperature of aggregates, binder, and HMA between 18 to 418ºF (10 and 232ºC).

4.5. Balance - A balance meeting the requirements of M 231, Class G5, for determining the mass of aggregates, binder, and HMA.

4.6. Oven - An oven, thermostatically controlled to ±5.4ºF (±3ºC), for heating aggregates, binder, HMA, and equipment as required. The oven shall be capable of maintaining the temperature required for mixture conditioning in accordance with R 30.

4.7. Miscellaneous - flat-bottom metal pans for heating aggregates, scoop for batching aggregates, containers (grill-type tins, beakers, containers for heating asphalt), large mixing spoon or small trowel, large spatula, gloves for handling hot equipment, paper disks, mechanical mixer (optional), lubricating materials recommended by the compactor manufacturer.
4.8 Maintenance—In addition to routine maintenance recommended by the manufacturer, check the Superpave gyratory compactor’s mechanical components for wear, and perform repair, as recommended by the manufacturer.

5. HAZARDS

5.1. Use standard safety precautions and protective clothing when handling hot materials and preparing test specimens.

6. STANDARDIZATION

6.1. Items requiring periodic verification of calibration include the ram pressure, angle of gyration, gyration frequency, LVDT (or other means used to continuously record the specimen height), and oven temperature. Verification of the mold and platen dimensions and the inside finish of the mold are also required. When the computer and software options are used, periodically verify the data processing system output using a procedure designed for such purposes. Verification of calibration, system standardization, and quality checks may be performed by the manufacturer, other agencies providing such services, or in-house personnel. Frequency of verification shall follow the manufacturer’s recommendations.

6.2 The angle of gyration may refer to either the external angle (tilt of mold with respect to a plane external to the gyratory mold) or the internal angle (tilt of mold with respect to end plate surface within the gyratory mold). Procedures used to verify the calibration of the angle of gyration must be appropriate for measuring the angle desired.

6.2.1 Method A—The calibration of the external angle of gyration should be verified using the manufacturer’s recommendations for the appropriate SGC.

6.2.2. Method B—The calibration of the internal angle of gyration should be verified in accordance with AASHTO PP 48.

6.2.3. The two methods (Method A—external and Method B—internal) of verifying the calibration of the gyration angle should NOT be considered equivalent. The gyration angle for all SGCs in a group for which compaction results are to be compared should be verified using the same method.

7. PREPARATION OF APPARATUS

7.1. Immediately prior to the time when the HMA is ready for placement in the mold, turn on the main power for the compactor for the manufacturer’s required warm-up period.

7.2. Verify the machine settings are correct for angle, pressure, and number of gyrations.

7.3. Lubricate any bearing surfaces as needed per the manufacturer’s instructions.

7.4. When specimen height is to be monitored, the following additional item of preparation is required. Immediately prior to the time when the HMA is ready for placement in the mold, turn on the device for measuring and recording the height of the specimen, and verify the readout is in the proper units, mm, and the recording device is ready. Prepare the computer, if used, to record the height data, and enter the header information for the specimen.
8. HMA MIXTURE PREPARATION

8.1. Weigh the appropriate aggregate fractions into a separate pan, and combine them to the desired batch weight. The batch weight will vary based on the ultimate disposition of the test specimens. If a target air void level is desired, as would be the case for Superpave mix analysis and performance specimens, batch weights will be adjusted to create a given density in a known volume. If the specimens are to be used for the determination of volumetric properties, the batch weights will be adjusted to result in a compacted specimen having dimensions of 6 inches (150 mm) in diameter and 4.53 ± 0.12 inches (115 ± 5 mm) in height at the desired number of gyrations.

Note 3—It may be necessary to produce a trial specimen to achieve this height requirement. Generally, 4500 – 4700 g of aggregate are required to achieve this height for aggregates with combined bulk specific gravities of 2.55—2.70, respectively.

8.2. Place the aggregate and binder container in the oven, and heat them to the required mixing temperature.

8.2.1. The mixing temperature range is defined as the range of temperatures where the unaged binder has a kinematic viscosity of 170 ± 20 mm²/s (approximately 0.17 ± 0.02 Pa·s for a binder density of 1.00 g/cm³) measured in accordance with ASTM D 4402.

Note 4—Modified asphalts may not adhere to the equi-viscosity requirements noted, and the manufacturer’s recommendations should be used to determine mixing and compaction temperatures.

Note 5—The SI unit kinematic viscosity is m²/s; for practical use, the submultiple mm²/s is recommended. The more familiar centistokes is a cgs unit of kinematic viscosity; it is equal to 1 mm²/s. The kinematic viscosity is the ratio of the viscosity of the binder to its density. For a binder with a density equal to 1.000 g/cm³, a kinematic viscosity of 170 mm²/s is equivalent to a viscosity of 0.17 Pa·s measured in accordance with T 316.

8.3. Charge the mixing bowl with the heated aggregate from one pan, and dry-mix thoroughly. Form a crater in the dry blended aggregate, and weigh the required amount of binder into the mix. Immediately initiate mixing.

8.4. Mix the aggregate and binder as quickly and thoroughly as possible to yield HMA having a uniform distribution of binder. As an option, mechanical mixing may be used.

8.5. After completing the mixture preparation perform the required mixture conditioning in accordance with R 30.

8.6. Place a compaction mold and base plate in an oven above the required compaction temperature for a minimum of 60 minutes prior to the estimated beginning of compaction (during the time the mixture is being conditioned in accordance with R 30).

8.7. Following the mixture conditioning period specified in R 30, if the mixture is at the compaction temperature, proceed immediately with the compaction procedure as outlined in Section 9. If the compaction temperature is different from the mixture conditioning temperature used in accordance with R 30, place the mix in another oven at the compaction temperature for a brief time (maximum of 30 minutes) to achieve the required temperature.

8.7.1. The compaction temperature is the mid-point of the range of temperatures where the unaged binder has a kinematic viscosity of 280 ± 30 mm²/s (approximately 0.28 ± 0.03 Pa·s) measured in accordance with T 316 (Note 4).
8.8. If loose HMA plant mix is used, the sample should be obtained in accordance with T 168. The mixture shall be brought to the compaction temperature range by careful, uniform heating in an oven immediately prior to molding.

9. COMPACTION PROCEDURE

9.1. When the temperature of the HMA is five degrees above the compaction temperature as shown on the “Mix Design Verification Report,” remove the heated mold, base plate, and upper plate (if required) from the oven. Place the base plate and a paper disk in the bottom of the mold.

9.2. Remove the pan of HMA from the oven and in one motion invert the pan onto the construction paper, vinyl mat, etc. Quickly remove any material that remains in the pan and include it with the HMA sample to be compacted. Grasp opposing edges of the paper and roll them together to form the HMA into a cylindrical shape. Insert one end of the paper roll into the bottom of the compaction mold and remove the paper as the HMA slides into the mold. This process needs to be accomplished in approximately 60 seconds. Place the mixture into the mold in one lift. Care should be taken to avoid segregation in the mold. After all the mix is in the mold, level the mix, and place another paper disk and upper plate (if required) on top of the leveled materials.

9.3. Load the charged mold into the compactor, and center the loading ram.

9.4. Apply a pressure of 600 ± 18 kPa on the specimen.

9.5. Apply a 1.25 ± 0.02° (22.0 ± 0.35 mrad) external angle or a 1.16 ± 0.02° (20.2 ± 0.35 mrad) average internal angle, as appropriate, to the mold assembly, and begin

9.6. Allow the compaction to proceed until the desired number of gyrations specified in PP 28 is reached and the gyratory mechanism shuts off.

9.7. Remove the angle from the mold assembly; retract the loading ram; remove the mold from the compactor (if required); and extrude the specimen from the mold.

**Note 6**—No additional gyrations with the angle removed are required unless specifically called for in another standard referencing T 312 (as in R 30 Section 7.3.2.1.2). The extruded specimen may not be a right angle cylinder. Specimen ends may need to be sawed to conform to the requirements of specific performance tests.

**Note 7**—The specimens can be extruded from the mold immediately after compaction for most HMA. However, a cooling period of 5 to 10 minutes in front of a fan may be necessary before extruding some specimens to insure the specimens are not damaged.

9.8. Remove the paper disks from the top and bottom of the specimens.

**Note 8**—Before reusing the mold, place it in an oven for at least 5 minutes. The use of multiple molds will speed up the compaction process.

10. DENSITY PROCEDURE

10.1 Determine the maximum specific gravity ($G_{max}$) of the loose mix in accordance with T 209 using a companion sample. The companion sample shall be conditioned to the same extent as the compaction sample.

10.2 Determine the bulk specific gravity ($G_{mb}$) of the specimen in accordance with T 166 or T 275 as appropriate.
10.3. When the specimen height is to be monitored, record the specimen height to the nearest 0.1 mm after each revolution in addition to those specified in Section 8.

11. DENSITY CALCULATIONS
   WSDOT has removed this section refer to WSDOT SOP 731.

12. REPORT
   WSDOT has removed this section refer to WSDOT SOP 731.

   12.2 Report results on WSDOT form 350-162 or other report approved by the State Materials Engineer.

13. PRECISION AND BIAS
   See AASHTO T 312 for Precision and Bias

14. KEYWORDS
   14.1. Gyratory; compaction; density
Performance Exam Checklist

*Determining Density of Hot Mix Asphalt (HMA)*
*Specimens by Means of the SHRP Gyratory Compactor*
*FOP For AASHTO T 312*

Participant Name ___________________________ Exam Date __________

**Procedure Element**

1. The tester has a copy of the current procedure on hand?  
   ![Yes][No]

2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?  
   ![Yes][No]

3. Main power for compactor turned on for manufacturer’s required warm-up period if applicable?  
   ![Yes][No]

4. Angle, pressure and number of gyrations set?  
   ![Yes][No]

5. Bearing surfaces, rotating base surface, and rollers lubricated?  
   ![Yes][No]

**Preparation of Mixtures**

1. Is mixture 5 degrees above compaction temperature shown on “Mix Design Verification Report?” If not, was mixture placed in an oven and heated to 5 degrees above compaction temperature?  
   ![Yes][No]

2. Mold and base plate heated for a minimum of 60 minutes in an oven at a temperature not to exceed the compaction temperature by 25 F?  
   ![Yes][No]

**Plant mix – Loose mix brought to compaction temperature by uniform heating immediately prior to molding.**

1. Mold, base plate and upper plate (if required) removed from oven and paper disk placed on bottom of mold?  
   ![Yes][No]

2. Mixture placed into mold in one lift, mix leveled, and paper disk and upper plate (if required) placed on top of material?  
   ![Yes][No]

3. Mixture removed from oven and mold charged within approximately 60 seconds?  
   ![Yes][No]

4. Mold loaded into compactor and a pressure of 600 ± 18 kPa applied?  
   ![Yes][No]

5. Angle of 1.25 ± 0.02° (22 ± 0.35 mrad) applied to the mold assembly and gyratory compaction started?  
   ![Yes][No]

6. Compactor shuts off when appropriate gyration level is reached?  
   ![Yes][No]

7. Mold removed and specimen extruded?  
   ![Yes][No]

8. Paper disks removed?  
   ![Yes][No]
<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>9. If specimens are used for determination of volumetric properties, are the</td>
<td></td>
<td></td>
</tr>
<tr>
<td>heights of the specimens 115 ± 5mm?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. All calculations performed correctly?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________
________________________________________________________________________
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________________________________________________________________________
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________________________________________________________________________
WSDOT Test Method T 313

Method of Test for Cement-Latex Compatibility

1. SCOPE

This method tests the compatibility of cement and latex additives when combined.

2. EQUIPMENT

   a. Two brass cylindrical unit weight cups having an inside diameter of 3 in. (76 mm) and a depth of approximately 3 15/32 in. (88 mm).

   b. Mixer, bowl, and paddle conforming to AASHTO 162.

   c. Straightedge — A steel straight edge, not less than 4 in. (102 mm) long, and not less than 1/16 in. (1.6 mm) nor more than 1/8 in. (3.2 mm) in thickness.

   d. Glass Graduated Cylinder — Shall have 250 ml capacity, graduations at 2 ml intervals, made to deliver indicated volume at 20° C.

   e. Glass Beaker — Shall have at least a 300 ml capacity.


   g. Tapping Stick — Made of hardwood, a diameter of 1/8in. (3.2 mm), and a length of 6 in. (152 mm).

   h. Tamper — Made of hardwood, a diameter of 7/16in. (11 mm), and a length of 6 in. (152 mm).

   i. Scoop — Approximately 8 in. (203 mm) long, and 3 in. (76 mm) wide.

   j. Scraper — Shall consist of a semirigid rubber blade attached to a handle about 6 in. (152 mm) long. Blade is about 3 1/2 in. (90 mm) long and 2 in. (50 mm) wide. (Kitchen tool called a plate and bowl scraper meets these requirements.)

3. PROCEDURE

   a. Weigh out a batch of washed and dried Steilacoom aggregate (B-1), using the following table:

<table>
<thead>
<tr>
<th>U.S. No. Screen Size</th>
<th>Batch Weights (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Individual Wt.</td>
</tr>
<tr>
<td>3/8 in. - #4 (9.5 mm - 4.75 mm)</td>
<td>14</td>
</tr>
<tr>
<td>#4 - #8 (4.75 mm - 2.36 mm)</td>
<td>205</td>
</tr>
<tr>
<td>#8 - #16 (2.36 mm - 1.18 mm)</td>
<td>186</td>
</tr>
<tr>
<td>#16 - #30 (1.18 mm - 600 µm)</td>
<td>230</td>
</tr>
<tr>
<td>#30 - #50 (600 µm - 300 µm)</td>
<td>561</td>
</tr>
<tr>
<td>#50 - minus (600 µm - minus)</td>
<td>278</td>
</tr>
</tbody>
</table>

   b. Weigh out 1 lb. (454 g) of the cement candidate. Measure 4.80 oz. (142 ml) of the latex candidate into a beaker. 2.976 oz. (Measure 88 ml) of water into the graduated cylinder.

   c. Put the aggregate and cement into the mixer bowl. (Adjust the blade to bowl clearance for the largest aggregate size first.) Mix the dry ingredients for 45 seconds at speed 1. Stop the mixer.

   d. Pour the latex sample into the mixer bowl. Rinse the latex beaker with the measured mixing water, then pour the solution into the mixing bowl.
e. Mix for one minute at speed 1.

f. Stop mixer for three minutes. Scrape down the sides and check the bottom of the bowl for sand pockets using the bowl scraper.

g. Mix for ten seconds at speed 1.

h. Tare the first unit weight (mass) cup.

i. Use the scoop to fill the unit weight (mass) cup in three lifts. For each lift, tamp with the small wood dowel 25 times, just penetrating the layer below it, and strike the side of the unit weight (mass) cup sharply five times with the large wood dowel, at equal intervals around its circumference.

j. Cut off the mortar to a plane surface, flush with the top of the unit weight (mass) cup, by drawing the straight edge, held at 90 degrees, with a sawing motion across the top, making two passes over the entire surface, the second pass being made at right angles to the first. Wipe off all mortar and water from the sides of the unit weight (mass) cup. Record the weight (mass). Tare the second unit weight (mass) cup and fill with the remaining mortar from the bowl, and repeat the procedure. Record the second weight (mass).

4. CALCULATIONS

a. Calculate the average weight (mass) of the two samples.

5. SIGNIFICANCE

If a latex additive and concrete mix are incompatible, they will react chemically, changing the specific gravity and density, which will adversely affect the final strength of the concrete/latex. In this test method, an adverse reaction is determined by examining the weight (mass) of the two wet products when combined. A very lightweight means there is probably a problem, and a different latex or cement should be submitted for testing.
WSDOT Test Method T 314

Method of Test for Photovolt Reflectance

1. SCOPE
   a. This method covers the determination of the 45-degree, 0-degree directional reflectance factor of nonfluorescent opaque specimens by means of a filter photometer.

2. APPARATUS
   a. Photovoltmeter conforming to ASTM E 1347.
   b. Calibrated standard plaques.
   c. Black plastic film canister.
   d. Filters.

3. PROCEDURE
   a. Reflectometer Calibration
      1. Warm up the reflectometer for 30 minutes before use.
      2. Turn the REF control knob until neither LO nor HI lights are lit.
      3. Place the black plastic film canister over the sensor unit opening, and turn the COARSE sensitivity control fully clockwise, and the FINE sensitivity control to its midrange (approximately five turns). Turn the ZERO knob until 00.00 appears on the PERCENT REFLECTANCE display.
      4. Put the required filter into the sensor unit. Set the sensor unit on top of the appropriate standard plaque and turn the COARSE sensitivity knob until a value close to the standard appears in the display. The standard value is printed on the reverse side of the plaque. Then, using the FINE sensitivity knob, dial in the exact value of the standard plaque.
   ii. Photovolt Model 577
      1. Warm up the reflectometer for 30 minutes before use.
      2. Select the appropriate filter channel by depressing the “G” button on the front panel.
      3. Place the black plastic film canister over the sensor unit opening, depress the CHANGE button, the display will go blank.
      4. Press the ZERO button, the display will read 0.0 (+/- 0.1)
      5. Set the standard plaque on the search unit, depress the CHANGE button, the display will go blank again.
      6. Depress the STD button, the display will read 75.0. Using the buttons A, B, and G set the display digits to read the value printed on the back of the standard plaque.
      7. Depress the STD button again. The display will display the standard plaque value +/- 0.2%.
b. Reflectance Determination

1. Remove the sensor unit from the standard plaque and place it on the specimen to be measured.

2. The PERCENT REFLECTANCE reading of the specimen will now appear in the display.

3. When testing Curing Compounds, take three readings for each specimen in the “brightest” locations. When testing lane markers, move the unit to determine an average reading. Recheck the standard plaque reading and the zero reading between each sample measurement.

4. REPORT

   a. Report the average result of all readings for a sample to the nearest percent.
WSDOT FOP for AASHTO T 329

Moisture Content of Asphalt (HMA) by Oven Method

1. SCOPE

1.1. This method is intended for the determination of moisture content of hot mix asphalt (HMA) by drying in an oven.

1.2. The values stated in SI units are to be regarded as the standard.

1.3. This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user of this standard to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. REFERENCED DOCUMENTS

2.1. AASHTO Standards:
   • M 231, Weighing Devices Used in the Testing of Materials
   • T 168, Sampling Bituminous Paving Mixtures
   • T 248, Reducing Samples of Aggregate to Testing Size

3. SUMMARY OF TEST METHOD

3.1. A sample of HMA is dried in a forced-air, ventilated, or convection oven at 325 ± 25°F (163 ± 14 °C).

3.2. The moisture content of the HMA is expressed for the HMA being in either a moist or dry condition, depending upon agency standards for reporting the asphalt binder content of the HMA:

   3.2.1. When the asphalt binder content is reported as a percent of the HMA, the moisture content is reported as a percent of the moist mass of the HMA, as shown in Section 7.1.1.

   3.2.2. When the asphalt binder content is reported as a percent of the aggregate in the HMA, the moisture content is reported as a percent of the dry mass of the HMA, as shown in Section 7.1.2.

4. APPARATUS

4.1. Balance or Scale—4.4-lb (2-kg) capacity, readable to at least 0.1 g and conforming to the requirements of M 231.

4.2. Forced-Air, Ventilated, or Convection Oven—capable of maintaining the temperature surrounding the sample at 325 ± 25°F (163 ± 14 °C).

4.3. Sample Container—the container in which the sample is dried shall be of sufficient size to contain the sample without danger of spilling and to allow the sample to be evenly distributed in a manner that will allow completion of the test in an expeditious manner.

This SOP is based on AASHTO T 329-05.
5. **SAMPLE**

5.1. A sample of HMA shall be obtained in accordance with WAQTC FOP for AASHTO T 168.

5.2. The sample shall be reduced in size in accordance with WSDOT T 712 T-248, Method B. The size of the test sample shall be a minimum of 500 g.

6. **PROCEDURE**

6.1. Determine and record the mass of the sample container to the nearest 0.1 g.

6.2. Place the test sample in the sample container. Determine and record the temperature of the test sample. To facilitate drying, evenly distribute the test sample in the sample container.

6.3. Determine and record the total mass of the sample container and moist test sample to the nearest 0.1 g.

6.4. Calculate the mass of the initial, moist test sample by subtracting the mass of the sample container determined in Section 6.1 from the total mass of the sample container and moist test sample determined in Section 6.3.

6.5. Dry the test sample to a constant mass in the sample container.

   **Note 1**—Constant mass shall be defined as the mass at which further drying at 325 ± 25°F (163 ± 14 °C) does not alter the mass by more than 0.1 percent. The sample shall initially be dried for 90 minutes and its mass determined, and then at 30 minute intervals until a constant mass is reached.

   **Note 2**—The moisture content of test samples and the number of test samples in the oven will affect the rate of drying at any given time. Placing wet test samples in the oven with nearly dry test samples could affect the drying process.

6.6. Cool the sample container and test sample to approximately the same temperature as determined in Section 6.2.

6.7. Determine and record the total mass of the sample container and dry test sample to the nearest 0.1 g.

   **Note 3**—Do not attempt to remove the test sample from the sample container for the purposes of determining the dry mass of the test sample.

6.8. Calculate the mass of the final, dry test sample by subtracting the mass of the sample container determined in Section 6.1 from the total mass of the sample container and dry test sample determined in Section 6.7.
7. CALCULATIONS

7.1. Moisture content is determined as described in either Sections 7.1.1 or 7.1.2, depending upon agency standards:

7.1.1. When the asphalt binder content is reported as a percent of the HMA, the moisture content is determined and reported as a percent of the mass of the initial, moist test sample as follows.

\[
\text{Moisture Content, } \% = \left( \frac{M_f - M_i}{M_i} \right) \times 100
\]

where:

- \( M_i \) = mass of the initial, moist test sample; and
- \( M_f \) = mass of the final, dry test sample.

Example: \( M_i = 541.2 \text{ g} \)

\[
M_f = 536.0 \text{ g}
\]

\[
\text{Moisture Content } = \left( \frac{541.2 \text{ g} - 536.0 \text{ g}}{541.2} \right) \times 100 = 0.96\%
\]

7.1.2. When the asphalt binder content is reported as a percent of the aggregate in the HMA, the moisture content is determined and reported as a percent of the mass of the final, dry test sample as follows.

\[
\text{Moisture Content, } \% = \left( \frac{M_f - M_i}{M_f} \right) \times 100
\]

where:

- \( M_i \) = mass of the initial, moist test sample; and
- \( M_f \) = mass of the final, dry test sample.

Example: \( M_i = 541.2 \text{ g} \)

\[
M_f = 536.0 \text{ g}
\]

\[
\text{Moisture Content } = \left( \frac{541.2 \text{ g} - 536.0 \text{ g}}{536.0} \right) \times 100 = 0.97\%
\]

8. REPORT

8.1. Report the moisture content to the nearest 0.01 percent.

8.2. Results shall be reported on standard forms approved for use by the agency.
# Performance Exam Checklist

**Moisture Content of Asphalt (HMA) by Oven Method**  
**WSDOT FOP for AASHTO T 329**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

## Procedure Element

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

## Test for Moisture

<table>
<thead>
<tr>
<th>Test for Moisture</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Representative sample obtained; 500 g minimum?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Mass of sample determined to nearest 0.1 g?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Initial temperature recorded?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Sample placed in drying oven for a minimum of 90 minutes?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Sample dried to a constant weight at 325 ±25°F?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Samples checked for additional loss?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Sample and container cooled to approximately the initial temperature before mass determined?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Calculation of moisture content performed correctly?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[
\text{% Moisture as percent of Wet Mass} = \left( \frac{M_i - M_r}{M_i} \right) \times 100
\]

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner ____________________________

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T 329  
January 2006  
Page 5 of 6  
T 329
WSDOT Test Method T 405

Method of Test for Diagonal Shear Strength of Epoxy Resins
(Arizona Composite Cylinder Test)

1. SCOPE
   This method covers a procedure for determining the shear strength of epoxy resins used to bond freshly mixed Portland cement concrete to hardened Portland cement concrete, by compression testing of a specially prepared, diagonally split, 6 in. by 12 in. (150 mm by 300 mm) standard concrete test cylinder.

2. APPARATUS
   b. Cylinder Molds — Standard 6 in. (152.4 mm) diameter by 12 in. (304.8 mm) tall, made from plastic.
   c. Elliptical diagonal half-cylinder filler, made from nondeforming material such as wood, epoxy or plastic, that fits inside a standard 6 in. × 12 in. (152.4 mm by 304.8 mm) cylinder mold such that the mold diameter remains round. Filler dimensions are approximated by drawing a line starting at 1 in. (25.4 mm) above the bottom left of the cylinder and traveling diagonally up and across the cylinder ending 1 in. (25.4 mm) below the top right of the cylinder. See drawing below (Figure 1).
   d. Tamper or vibrator.
   e. Small trowel.
   f. Measuring Cups — 250 to 300 ml graduated plastic tri-pour beakers are one type.
   g. Mixing Container — 21 polyethylene is one type of container.
   h. Stir Rod — Wooden tongue depressors are one type.
   i. Applicator Brush — 1 in. (25.4 mm) disposable painters chip brush is one type.

3. REAGENTS
   a. Hydrochloric acid, 10 percent.
   b. Portland cement concrete, WSDOT Class 4000 (28 MPa), enough to make two standard 6 in. × 12 in. (152.4 mm by 304.8 mm) cylinders per sample.
4. SPECIMENS

Make Portland cement concrete half-cylinder specimens as follows:

a. Place elliptical filler in mold such that elliptical surface is up.

b. Put bond breaker mold release on elliptical surface if necessary (i.e., for wood. If fillers are made from epoxy or plastic a mold release agent may not be necessary).

c. Fill mold to top with Portland cement concrete mixture, using mixture as above. Vibrate and smooth top even with top of mold.

d. Cure first half of specimen 28 days minimum at 100 percent humidity. Molds can be stripped from the composite cylinders as per WSDOT Test Method 809 Section 5 Method 1. Store in a manner such that the elliptical surface is up, i.e., not in contact with storage racks.

Figure 1
Diagram of composite cylinder
5. PROCEDURE

a. The epoxy is tested in duplicate, so two half-cylinder specimens are needed for each sample. Remove cured concrete half-cylinder specimen from 100 percent humidity storage room and place in a sink, preferably in a chemistry laboratory hood, with the elliptical surface up. Acid etch the elliptical surface with the 10 percent hydrochloric acid solution. After the reaction ceases, wash surface thoroughly with running tap water and allow to air dry, approximately ½ hour.

b. Place the cured, acid etched, concrete half-cylinder partially into cylinder mold with elliptical surface up.

c. Tilt mold so that the elliptical surface is horizontal and place a support under the mold to hold it at this position.

d. Mix the epoxy test sample using manufacturer’s mix ratio and directions. Stir thoroughly for proper mixing (three minutes minimum is usually sufficient). Using a brush, apply the epoxy in an even layer to the entire elliptical surface. The thickness of the epoxy layer should be 10 to 15 mils (0.3 mm to 0.4 mm).

e. Place a layer of plastic Portland cement concrete, same mixture as described above, over the epoxy bonding compound, taking care not to allow large aggregate to become lodged in the toe of the mold. Return mold to a vertical position. The half-cylinder should slide into the mold. Seat the half-cylinder with the epoxy on it into the mold by picking up the mold and tapping the bottom of it once or twice onto the table top. (Caution, watch out for splashing epoxy.)

f. Fill remainder of mold with concrete mixture as described above. Vibrate and smooth the top.

g. Cure the composite cylinder (old concrete, epoxy, and fresh concrete) for seven days at 100 percent humidity, then test in compression machine as indicated in AASHTO T 22. If the first cylinder fails to meet specifications, the second cylinder is cured another seven days and then tested on the compression machine.

6. CALCULATION

The bond strength is determined as follows:

\[
\text{Bond Strength, psi (MPa)} = \frac{T}{A}
\]

\[
T = \text{Total Load to failure, pounds (Newton)}
\]

\[
A = \text{Area of elliptical surface, square inches (square meters)}
\]

The area of the elliptical surface is determined as follows:

\[
\text{Area of ellipse} = \pi \left( \frac{D}{2} \right) \left( \frac{d}{2} \right)
\]

\[
D = \text{diameter of ellipse at greatest length, inches (mm)}
\]

\[
d = \text{diameter of ellipse at greatest width, inches (mm) = diameter of cylinder}
\]

\[
\pi = 3.1416
\]
D = hypotenuse of right triangle formed by diameter of cylinder, \( a = 6 \text{ in.} \ (152.4 \text{ mm}) \) and height of diagonal slant normal to the cylinder base, \( b = 10 \text{ in.} \ (254 \text{ mm}) \); Therefore, \( D = c = (a^2 + b^2)^{1/2} = 11.66 \text{ in.} \ (296.2 \text{ mm}) \)

\[ d = 6 \text{ in.} \ (152.4 \text{ mm}) \]

\[ A = (3.1416)(11.66 \text{ in/2})(6 \text{ in/2}) = 54.95 \text{ sq. in.} \]

or

\[ A = (3.1416)(296.2 \text{ mm/2})(152.4 \text{ mm/2}) = 35,454 \text{ square mm} = 0.0355 \text{ sq. m} \]

7. REPORT

The bond strength reported shall be the average of the two composite cylinders tested, except if the first cylinder fails to meet specifications after the initial seven-day cure and the second cylinder was tested after another seven days of curing, report both bond strength and days cured for each cylinder.
1. SCOPE  
   a. This method is intended for laboratory use in determining the quality of water to be used in mixing concrete.

2. APPARATUS  
   a. Porcelain evaporating dish with 3.4 oz. (100 ml) minimum capacity.  
   b. 3.4 oz. (100 ml) pipet.  
   c. Drying oven maintained at 212°F (100°C).  
   d. Analytical balance.

3. PROCEDURE  
   a. Pipet 3.4 oz. (100 ml) of sample into a weighed porcelain evaporating dish and record total weight (mass). Evaporate to dryness, cool in a desiccator and reweigh, using analytical balance for all weighing.

4. CALCULATION  
   a. \[ \frac{g \text{ of residue}}{g \text{ of sample}} \times 10^6 = \text{ppm total solids} \]

5. REPORT  
   a. All test results will be reported on Test Data Form WSDOT 350-034.  

   Note: The determination of the composition of the mineral matter in the water requires a complete chemical analysis and is not generally undertaken except when the percentage of total solids is above 1000 ppm. When the mineral analysis is desired, the procedure starting on page 2388 of Scott’s Standard Methods of Chemical Analysis, Sixth Edition (1963), Volume II, should be used. The results should be reported as the separate constituents in parts per million. If the hypothetical combination into salts is desired, the method given by Scott, or the method given on page 336, Volume V, Number 5, Industrial and Engineering Chemistry, should be used.
WSDOT Test Method No. 411

Method of Test for Water Absorption and Moisture Vapor Transpiration

1. SCOPE
   a. This method is intended for laboratory uses in determining the efficiency of water repellant compounds.

2. APPARATUS
   a. Test blocks 1 1/2in. (38.1 mm) thick by 3 1/2 in. (88.9 mm) wide by 6 in. (152.4 mm) long.
   b. Oven maintained at 100°F (37.8°C).

3. TEST SPECIMENS
   a. Three specimens per test shall be made with one part of Type 1 or Type II Portland cement, one part hydrated lime, six parts 8- to 28-mesh sound natural sand, six parts 28- to 100-mesh sound natural sand, and 1.94 parts of water on a weight basis. These materials shall be thoroughly mixed for two minutes by hand or mechanical mixer in a suitable bowl after the water has been added. Following this the mix shall be cast in molds that will make specimens approximately 1 1/2in. (38.1 mm) thick, 3 1/2 in. (88.9 mm) wide, and 6 in. (152.4 mm) long.
   b. Specimens shall then be cured one day in the molds at 73.4 ± 3°F (23°C ± 1.7°C) and six days in 95 percent relative humidity at 73.4 ± 3°F (23°C ± 1.7°C). At the end of the moist curing period, the specimens shall be dried in 20 to 30 percent relative humidity at 100°F (37.8°C) to constant weight (mass).

4. PROCEDURE FOR WATER ABSORPTION TEST
   a. When the specimens reach constant weight the water repellant compound shall be applied in two applications at the rate of 150 sq. ft. per gallon (3.7 m² per liter) for each application to the bottom sand side surfaces of the specimens as cast. A minimum of 12 hours air drying shall be allowed between applications. The specimens shall be allowed to air dry for 24 hours after final application before starting the test.
   b. The specimens shall then be weighed to the nearest gram and placed in 1/4 in. (6.4 mm) of 73.4 ± 3°F (23°C ± 1.7°C) water with the treated 3 1/2 in. (88.9 mm) by 6 in. (152.4 mm) surface face down. The water shall be maintained at a constant level and the units shall be supported so as to permit the free circulation of water around and under them. The specimens, after 72 hours immersion, shall be surface dried and weighed to the nearest gram. The water absorption shall be calculated as a percentage of the dry weight (mass). If the average of three specimens shows more then 2 percent moisture absorption, the water repellant compound is unsatisfactory.
5. PROCEDURE FOR MOISTURE VAPOR TRANSPERSION TEST
   a. Specimens meeting the water absorption test specification, they shall then be tested for transpiration (breathing).
   b. Immediately following the absorption test, the three test specimens shall be placed, untreated face down, in the water storage until they reach saturation as determined by a constant weight (mass). Total water absorbed shall then be recorded. At this point, the specimens shall be surface dried and the untreated face completely covered with an oversized piece of wax paper. The wax paper shall be applied by placing the untreated face of the specimen down on the wax paper and then pouring a hot wax fillet between the wax paper and the vertical sides of the specimen. The weight (mass) of each specimen shall then be recorded and the specimen placed, treated side up, in a suitable location where there is free circulation of air at 73.4 ± 3°F (23° ± 1.7°C) and 40 to 50 percent relative humidity. The specimens shall be weighed after seven days drying to determine the moisture loss. The loss shall be expressed as a percentage of the total water absorbed and shall be not less than 50 percent at seven days.

6. REPORT
   a. Test results will be reported on Test Data Form WSDOT 350-034.
WSDOT Test Method T 412
Bond Test for Joint Sealants

1. SCOPE
This method covers the bond test for joint sealers.

2. BOND TEST APPARATUS
   a. Bond Extension Apparatus — The apparatus shall consist of an extension machine, similar to that described in Federal Specifications SS-R-406, so designed that the specimen can be extended for four hours at a uniform rate of approximately 1/8 in. (3.175 mm) per hour. It shall consist essentially of one or more screws rotated by an electric motor through suitable gear reductions. Self-aligning plates or grips, one fixed and the other carried by the rotating screw or screws, shall be provided for holding the test specimen in position during testing. The machine must be capable of reversing itself for the recompression cycle. The bond extension machine shall be housed in a refrigeration unit capable of maintaining a temperature of 0°F ± 5°F (–18°C ± 2°C).
   b. Concrete Blocks — 1 in. × 3 in. × 8 in. (25.4 mm × 76.2 mm × 203.2 mm), sawed from 28-day cured WSDOT Class 4000 (28 MPa) concrete. **Note:** Do not use kerosene as a cutting solvent, use water.
   c. Spacers made of brass or steel.
      (1) 6 in. × 1 in. × 1/2 in. (152.4 mm × 25.4 mm × 12.7 mm)
      (2) 2 in. × 1 in. × 1/2 in. (50.8 mm × 25.4 mm × 12.7 mm)
   d. Release Agent — Release agent consisting of 50 percent talc (baby powder), 35 percent glycerin and 15 percent water-soluble medical lubricant (KY jelly) blended into a smooth paste. (Percentages by weight.)

3. BOND TEST PREPARATION
   Prior to use, the cured blocks shall be removed from storage in 98 percent relative humidity, and acid etched with hydrochloric acid. The blocks shall be rinsed with water, wiped and allowed to surface dry before using.
   Two prepared blocks shall then be placed on a glass plate and a spacer 6 in. × 1 in. × 1/2 in. (152.4 mm × 25.4 mm × 12.7 mm) placed between them. At each end of the spacer block, a spacer block 2 in. × 1 in. × 1/2 in. (50.8 mm × 25.4 mm × 12.7 mm) shall be placed on end. Prior to using, the spacer blocks shall be coated with a release agent. When the blocks are pushed together and secured with either clamps or rubber bands, a joint is formed 6 in. × 1 in. × 1/2 in. (152.4 mm × 25.4 mm × 12.7 mm) in size.
   a. Hot poured joint sealer, prepared as per ASTM D-5167, is poured in the formed joint to the top of the spacer blocks. Allow the specimen to cure for 24 hours.
   b. Two component joint sealers shall be prepared according the manufacturers recommendations and poured in the formed joint to the top of the spacer blocks. Allow to cure for 96 hours.
   Two samples are required for each test.
4. **BOND TEST PROCEDURE**

Remove the spacers and insert both test specimens in the bond extension machine. The bond extension machine must be at 0°F ± 5°F (–18°C ± 2°C). Allow approximately one hour for the samples to attemperate to test temperature. The blocks shall then be extended to 1/2 in. (12.7 mm) in approximately 4 hours. The refrigeration unit shall then be turned off, the door opened and the motor reversed for four hours to recompress the specimens to their original size at the same rate. This completes one cycle.

After one cycle, one of the samples is selected as the “soaked” sample. This sample is soaked for 16 hours in room temperature water before each of the next two cycles. The table below shows the cycles:

<table>
<thead>
<tr>
<th>Cycle</th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Dry</td>
<td>Dry</td>
</tr>
<tr>
<td>2</td>
<td>Dry</td>
<td>Soaked</td>
</tr>
<tr>
<td>3</td>
<td>Dry</td>
<td>Soaked</td>
</tr>
<tr>
<td>4</td>
<td>Dry</td>
<td>(Finished)</td>
</tr>
</tbody>
</table>

On the fourth cycle, the “soaked” sample is left in the machine for balance.

The development of any crack, separation, or other opening in the joint sealer or between the joint sealer and the block shall be considered as a failure to pass the bond extension test.

5. **REPORT**

The results of the bond test shall be reported on the appropriate report sheet.
WSDOT Test Method T 413

Method of Test for Evaluating Waterproofing Effectiveness of Membrane and Membrane-Pavement Systems

1. SCOPE
   a. This method describes a procedure for evaluating the waterproofing effectiveness of membranes and membrane-pavement systems as applied to bridge decks. The tests can be performed on the membrane alone or on the membrane-pavement combination.

2. APPARATUS
   a. Ohmmeter — Simpson VOM Model 313 or equal; driving voltage for the range of “R × 10K” should be 1.56 ± 0.05 V; the range of “R × 100K” should be 1.43 ± 0.05 V.
   b. No. 18 Insulated Wire — Belden test probe wire or equivalent, two spools, 250 ft. (76.2 m) each, with connectors.
   c. Metal Contact Plate — 12 × 12 × ½ in. (304.8 mm × 304.8 mm by 3.175 mm), with a connection for the ohmmeter and a 36-in. (914.4 mm) nonmetallic handle (see Figure 1, below).
   d. Polyurethane Sponge — 12 × 12 × 1 in. (304.8 mm × 304.8 mm by 25.4 mm), medium density (see Figure 1, below).

![Figure 1](image-url)

**Figure 1**
e. Wetting Agent — Aerosol OT (10 percent solution as manufactured by the American Cyanamid Company) or equal; mix 0.4 cup (100 ml) of wetting agent with 5 gal. (18.9 L) of water to make up wetting solution.

f. Pressure Spray Can — 3 gal. (11.4 L) capacity.

g. ¾-in. Stone Cutter’s Chisel.

h. Hammer.

i. Eye Protection.

Note: Items g, h, and i are used for connection to reinforcing steel in deck (negative pole).

3. PROCEDURE

The following procedures shall be used only when the surface of the membrane-only system is dry or when the membrane-pavement system is judged to be dry internally. The latter condition is an important consideration since moisture will conduct electricity throughout an entire asphalt concrete overlay producing erroneous results at the individual test locations.

a. Membrane Only

(1) When the membrane has cured sufficiently to allow foot traffic on it, divide the bridge deck into a grid pattern similar to that illustrated in Figure 2. The grid spacing shall be as ordered by the Engineer, but it is suggested that the grid lines be spaced at 5 ft. (1.5 m) intervals to provide adequate test coverage. A definite connection to the top mat of reinforcing steel in the bridge deck is desirable; however, if this is not feasible, the bridge railing, expansion joints, light standards, drainage scoupers, or other exposed steel should provide the necessary connection. New bridges are supplied with connection to the reinforcing steel along each edge of the bridge via a junction box.

(2) Uncoil an ample length of wire to reach the areas to be tested and attach the negative (–) connection of the ohmmeter to the reinforcing steel and the positive (+) connection to the 12 in. × 12 in. by 1/8 in. (304.8 mm × 304.8 mm × 3.175 mm) metal contact plate. Check ohmmeter battery for satisfactory charge, then zero the ohmmeter dial indicator.

(3) Attach the polyurethane sponge to the metal plate with heavy-duty rubber bands and saturate the sponge with the wetting solution.

(4) In order to check for proper electrical connections and overall equipment operation, prewet two or three areas along the exposed concrete curb and place the measuring device at each location; read and record the resistance readings displayed on the ohmmeter. These readings will normally range from 500 to 8,000 ohms, if the connection to the reinforcing steel is good.

(5) Once the connections and equipment show proper operation, continue testing at each grid intersection and record the resistance values on photocopies of Figure 3, using blank sheets where additional space is needed.
b. Membrane-Pavement

(1) New pavement — Procedure steps 3.a.(1) through 3.a.(5) may be used in measuring the resistance of the membrane-pavement system, except that in step 3.a.(5) the test locations should be prewetted with the wetting solution. Only enough wetting solution should be used to saturate the test zone. Surface runoff should be kept to a minimum to prevent interconnecting the test zones electrically.

(2) Old Pavement — Procedure steps 3.a.(1) through 3.a.(5) may be used in measuring the resistance of the membrane-pavement system, except that in step 3.a.(5) a checkpoint should be selected at a grid intersection that appears dense-graded and well compacted. Saturate the check point with the wetting solution. Keep surface runoff to a minimum to prevent interconnecting the test zones electrically. Take resistance readings at the checkpoint until the value has stabilized at its lowest point. The penetration process should not required more than 15 to 20 minutes. If it takes more than 20 minutes, select a reasonable penetration time and saturate each grid intersection thereafter for that length of time.

4. REPORTING

a. Report the resistance values for the exposed membrane and the membrane-pavement system on photostatic copies of Figure 3, using blank sheets where additional space is needed.

b. Copies of the report are to be submitted to the following offices:

   (1) Project Engineer (for final records).
   (2) FOSSC Materials Laboratory (master file).

5. ACCEPTANCE LEVEL

Refer to the Contract Special Provisions under the section entitled “Membrane Waterproofing (Deck Seal).”
Figure 2: Example of Test Grid and Resistance Tabulation

7K = 7,000 OHMS
2M = 2,000,000 OHMS
∞ = INFINITE RESISTANCE
ELECTRICAL RESISTIVITY MEASUREMENTS  
(WDOT Test Method No. 413)

SR __ CONTRACT ____ BRIDGE ___________________________ BR. NO. ___ / ___
TEST PHASE: (Check one) ___________________________ DATE ____ / ____ / ___
   ____ MEMBRANE ONLY; TYPE ______________________
   ____ MEMBRANE-PAVEMENT; PAVT. TYPE ______ THCK ___ ft.
GROUND CHECK RDG. ______ ohms (avg.) OHMMETER RANGE USED R x _________
GRID SPACING ______ ft. CREW ______________________

***BRIDGE & GRID LAYOUT AND MEASUREMENTS***

SCALE: VERT. ___ in. = ___ ft.; HORIZ. ___ in. = ___ ft. NOTE: Use additional sheets (blank) if necessary.

Figure 3: Example of Electrical Resistivity Measurements
WSDOT Test Method No. 414

Method of Test for Total Chloride Ion in Concrete

NOTE 1: WSDOT HAS ADOPTED THE PROCEDURE DESCRIBED IN AASHTO T 260, PROCEDURE WITH THE FOLLOWING MODIFICATIONS: WSDOT considers this adoption to be as a Laboratory Operating procedure consistent with the technical practice as defined in T 260, Procedure A.

5. PROCEDURE-

Modify section 5.1 to read:

5.1. Determine the mass to the nearest milligram of a 3 g powdered sample representative of the material under test. Weigh sample as received, not oven dried.

Replace Section 5.2.1 & 5.2.2 with the following:

5.2.1. Transfer the sample quantitatively to a 400 ml beaker, add 50 ml of boiling distilled water and swirl to suspend sample. [CAUTION: wear goggles and acid proof gloves when handling Nitric Acid.] Add 5 ml of HNO3 and swirl, cover with watch glass and boil for one minute on hot plate in hood, [CAUTION: acid fumes, don’t boil outside of hood or bring out of hood while hot, cool in hood.] Cool, rinse watch glass into beaker with distilled water. Add cool distilled water to 200 mls total in beaker.

Added note: AASHTO T-260-94 procedure calls for a filtering process before titrating. This is a time consuming step which was eliminated at the expense of wearing out glass beakers and Teflon stir bars sooner. It has no effect on the accuracy or precision of chloride determinations.

Replace Section 5.4.1. and 5.4.1.1 with the following:

5.4.1 Sample Preparation

a. Calibrate the burette according to the calibration procedure in appendix B.

b. Prepare the electrodes as in appendix C.

c. To the cool sample, pipette 5.00 mls of 0.010 N standard NaCl solution to the beaker. This is called the standard spike.

d. Place, without splashing, a Teflon stir bar into beaker and place beaker onto stirrer. Turn on stirrer until the sample solution is gently stirring such that a vortex is barely discernible, avoid any splashing.

e. Rinse the two electrodes with distilled water and then immerse them into the sample solution about 15-25 mm deep. Turn on mV meter. The display should read a negative number. As the titration progresses, the display should become more positive.

f. Turn on computer and call up chloride program from the LIMS Technical System for Chemistry Section. Choose “C-Standardize” from program menu.

g. Standardize the silver nitrate solution according to the procedure in appendix A. After standardization of the silver nitrate and calibration of the burette, the normality and volume per increment of the silver nitrate solution determined are entered into the program at the appropriate prompts.
5.4.1.1 Titration of Sample and Computation of Results

a. Choose the ‘A-Run Routine Sample’ from program menu. Enter information asked for by the program prompts, i.e. sample #, bridge name, date, etc.

b. Record the initial mV reading and the initial burette reading (it is convenient to set burette to zero at beginning of each titration).

c. Add silver nitrate until approximately 20 mV below endpoint, record the volume added and mV reading at this point. (the endpoint, i.e. when there is no uncomplexed chloride ion present, can be approximated by adding 5 ml HNO₃ to 200 ml distilled water and observing the mV reading.)

d. The titration involves adding a known volume of standardized silver nitrate from the burette (i.e. 5 drop increment) to the sample and recording the millivolt reading, adding another 5 drops and recording the millivolt reading, and continuing in this pattern until the endpoint of the titration is reached. After each increment, when the mV reading is entered into the computer, the program prints the volume of silver nitrate added, the mV value and the difference between this value and the previously entered mV value, called delta mV. As the titration progresses, the change in mV becomes successively larger, reaches a maximum and then becomes successively smaller. The endpoint of the titration is the point where the change in mV reading is the largest, and the volume of silver nitrate at that point can be read from the printout.

Note: For the best results, the same number of increments and mV data points are recorded above the endpoint as below the endpoint. (i.e., if there are 5 increments above the endpoint, continue titrating 5 increments below the endpoint.)

e. When the titration is finished, the chloride program prompts for the end key. (F10). The amount of the spike (the 5.00 ml of 0.010 N NaCl added to each sample before titration began) must now be subtracted from the total. Enter 5 when it asks for the amount of the spike. (note, if more than a 5 ml spike was added, enter the actual amount of the spike.) If the original sample was not 3 g, enter the actual weight of the sample at the prompt.

f. The computer program then calculates the endpoint and prints the answer in lb. of chloride ion per cubic yard of Portland concrete cement.

\[
\text{% Cl}^- = \left[ \frac{\text{mls AgNO}_3 \times \text{N AgNO}_3 \times 3.5453}{\text{weight sample}} \right] \\
\text{kg Cl}^- / \text{m}^3 \ (\text{or lb. Cl}^- / \text{yd}^3) = \left[ \% \text{ Cl}^- \right] \times \left[ \frac{\text{UW}}{100} \right] \\
\text{UW} = \text{unit mass of concrete} = 2483 \text{ kg/m}^3 = 4185 \text{ lb./yd}^3
\]

Example:

Chloride ion = \left[ \frac{(0.93 \text{ mls AgNO}_3 \times 0.01 \text{ N AgNO}_3 \times 0.035453 \times 4.185 \text{ lb./yd}^3) }{3 \text{ g sample weight}} \right] = 0.46 \text{ lb. Cl}^- / \text{yd}^3 \text{ Portland concrete cement.}
Add the following Section:

5.4.4 Chloride Ions in Admixtures: When testing admixtures, accelerators, water reducers, etc., add reagents to sample in this order: 50 ml cool distilled water, carefully add 5 ml H₂SO₄, 8 ml of 30 percent H₂O₂, 5 ml HNO₃. Then continue as above in 5.2.1...'cover with watch glass'. Some mixtures will boil rapidly, so careful attention is needed to prevent boil over. Admixtures are digested using the sulfuric acid/peroxide/nitric acid mixture but otherwise treated the same as a PCC chloride sample. Choose “D-Use Admixture Routine” from the program menu. The computer program can calculate results by weight or volume, choose either depending on the state of the sample. If by volume, the density of the sample must be determined or known for the calculations.

\[
\% \text{Cl}^- = \frac{[\text{mls AgNO}_3] \times (N \text{AgNO}_3) \times (3.5453) \times (\text{aliquot})}{(\text{weight}) \text{ or } ((\text{volume of sample})(\text{density of sample}))}
\]

To convert % to PPM or mg/kg:

\[
[\% \text{Cl}^- /100][1000 \text{ mg/g }][1000 \text{g/kg}] = [\% \text{Cl}^-] \times 10,000 \text{ PPM or mg/kg}
\]

Add Appendices A, B, C and D as follows:

**APPENDIX A**

**Standardization of Secondary Standard**

1) To a 400 ml beaker containing around 100 ml distilled water, add 5.0 ml HNO₃
2) Add distilled water to bring the volume to 200 ml.
3) Add 10.0 ml of primary standard 0.010 N NaCl solution.
4) Proceed with titration as per 5.4.1 (4)-(6) and (9)-(13)
5) Run 2 to 4 standardization’s and find the mean. Each standardization should agree within 0.01 ml of titrant AgNO₃).
6) Calculate the Normality of the AgNO₃ solution.

\[
N \text{AgNO}_3 = \frac{[\text{mls NaCl}] \times (N \text{NaCl})}{[\text{mls AgNO}_3]}
\]

Note: If reagents are prepared as in AASHTO T-260-94 (3.3) and (3.4), then the normality of the silver nitrate solution should approximate 0.01N

**APPENDIX B**

**Calibration of Burette**

1) Fill burette to zero mark with titrating solution.(or distilled water).
2) Let burette deliver solution one drop at a time while counting the drops.
3) When a convenient number of drops have fallen (50, 100, etc.), record this number and the final volume of the burette.
4) Calculate the volume per drop.

\[
\text{Volume/drop} = (\text{ending burette reading} - \text{initial burette reading})/# \text{ drops}
\]
5) Repeat until volume/drop is reproducible. Calculate average volume of drop.
6) Determine volume of increment.

\[ \text{volume/increment} = \left( \frac{\text{volume/drop}}{\# \text{ of drops/increment}} \right) \]

Example: 50 drops used 2.49 mls.

\[ \text{volume/drop} = \frac{2.49}{50} = 0.0498 \text{ ml/drop} \]

\[ \text{volume/increment} = 0.0498 \text{ ml} \times 5 \text{ drops} = 0.249 \text{ ml or approx. 0.25 ml} \]

**APPENDIX C  Electrodes**

A) Selective Ion Electrode:

1) Clean with soft tissue to remove any adhering particles.

2) To polish membrane; place polishing cloth on hard smooth surface and put a drop of distilled water on the cloth. Use a circular motion polishing the electrode sending unit for about 30 seconds exerting medium pressure. Rinse and soak the electrode in a standard solution for about 5 minutes before use. Standard solution: Make a solution that is 1,000 PPM with respect to chloride ion. Place 1.65 g reagent grade NaCl in a 1 liter volumetric flask, brought up to volume with distilled water.

3) Storage: Store selective ion electrode in distilled water.

B) Reference Electrode

1) Inner chamber: Fill with bridge solution. Drain and replace as needed to correct erratic meter readings. Bridge solution: Add 52.77 grams acetate trihydrate (MUST BE TRHYDRATE) and 2.88 grams sodium nitrate to a 100 ml volumetric flask. Dilute with distilled water to mark. Mix well. CAUTION: When adding this solution to reference electrode. Do not get any of it into samples or standards, as it will affect results drastically.

2) Storage: Store selective ion electrode upright in distilled water, standard solution or sample solution.

C) Electrode Measuring Hints:

1) Allow all solutions to come to room temperature.

2) Stir all solutions with a magnetic stirrer at a rate that will not cause a vortex. Placing a piece of insulating material such as cork, cardboard or Styrofoam, between the stirrer and sample beaker to reduce heat transfer from stirrer to beaker.

3) Rinse electrodes and blot dry with a tissue (Kim Wipes) between measurements to prevent solution carryover.

4) After determining Cl content in an organic mixture (i.e., admixtures), clean both electrodes, polish chloride ion selective electrode.

D) To Check Overall Performance of Double Junction Reference Electrode and Specific Ion Electrode System.

1) Add 100 ml standard salt solution to 400 ml beaker (1,007 PPM chloride). (1.66 g NaCl/1000 ml)

2) Add 2 ml ionic strength adjuster, mix and record millivolt reading.

Ionic Strength Adjuster: Add 50.5 g KNO₃ to a 100 ml vol. flask, diluting and mixing well. Solution will be saturated so let stand 10 minutes before pipeting off the top.
3) Dilute 10 ml standard salt solution to 100 ml and record millivolt reading. (This solution yields 100.7 PPM chloride)

4) Difference between two readings (step 2 and step 3) should be 56 +/- 3 millivolts.

APPENDIX D  Treatment of Teflon Stirring Bars
Magnetic stir bars gather iron filings on their outer surface. If these are not removed, the filings tend to work themselves into the Teflon coating. Brushing with a stiff brush under running water and then towel drying with a paper towel removes the larger particles. When the bars become dirty looking due to the buildup of iron filings, soak in concentrated HCl for 1 to 2 hours (or overnight). the iron filings will dissolve. Rinse in running tap water, then rinse in distilled water before use to avoid chloride ion carryover from the acid. Boiling in 1:1 aqueous HNO₃ will dissolve FeCl₃ present which is then rinsed away with distilled water. Air dry stir bars and store in chloride free environment. Rinse in distilled water before use.

Glass stirring bars are also acceptable and will solve the problem of the iron filings working into the Teflon. After much use the Teflon bars become flat and the glass bars become frosted. The resulting irregular surfaces can be responsible for solution carryover and contamination of samples. Make sure the stir bars are well leaned. Periodically inspect the stir bars and discard any that show excessive wear.
WSDOT Test Method T 415

Method of Test for Fertilizer

1. SCOPE
   This method is used to determine percentage of nitrogen, phosphorus, and potassium in fertilizer.

2. EQUIPMENT
   ECO FP 428 Nitrogen Analyzer
   Thermo-Jarrell Ash Inductively Coupled Plasma Spectrophotometer (ICP)

3. TOTAL NITROGEN
   a. Reagents
      (1) Ethylenediaminetetraacetic acid (EDTA).
   b. Procedure
      (1) Run blanks and standards (EDTA) and samples as required following the methods outlined in the OPERATION section of the LECO FP 428 Nitrogen Analyzer INSTRUCTION MANUAL.
      (2) Record Fertilizer results as % Nitrogen.

4. TOTAL PHOSPHORUS and TOTAL POTASSIUM
   a. Reagents
      (1) Nitric Acid (HNO3).
      (2) Hydrochloric Acid (HCl).
      (3) Standard Phosphorus solution (100 ppm) prepared from 10,000 ppm certified standard (traceable to NIST).
      (4) Standard Potassium solution (100 ppm) prepared from 10,000 ppm certified standard (traceable to NIST).
   b. Sample Preparation
      (1) Weigh approximately 0.25 grams of fertilizer sample to the nearest milligram. Transfer to a 250 ml beaker. Add 30 ml HNO3 and 3 to 5 ml HCl and boil until organic matter is destroyed. Boil to white fumes, cool the solution, transfer to a 250 volumetric flask and bring up to volume.
      (2) Prepare a blank in the same manner using 30 ml HNO3 and 3 to 5 ml HCl.
   c. Phosphorus Analysis
      (1) Following procedures outlined in the Thermo Jarrell-Ash ICP Instruction Manual, standardize the ICP for Phosphorus and run samples to obtain ppm Phosphorus.
      (2) Calculate % P2O5 by the following formula:
          \[ \% \text{ P2O5} = \frac{(\text{ppm Phosphorus} \times 57.29)}{(\text{mg sample})} \]
d. Potassium Analysis

(1) Following procedures outlined in the Thermo Jarrell-Ash ICP Instruction Manual, standardize the ICP for Potassium and run sample to obtain ppm Potassium.

(2) Calculate % K2O by the following formula:

\[ \% \text{ K2O} = \frac{(\text{ppm Potassium} \times 30.12)}{\text{mg sample}} \]
WSDOT Test Method T 417

*Method of Test for Determining Minimum Resistivity and pH of Soil and Water*

1. **SCOPE**
   
a. This method covers the procedure for determining the minimum resistivity and pH of soil or water samples at metal culvert locations. These values are used to assist in determining the type of metal culvert materials and protective coating that are permissible at each location.

b. This test method is divided into the following parts:
   1. Method of field resistivity survey and sampling for laboratory tests.
   2. Method of determining pH of water.
   4. Laboratory method of determining minimum resistivity.

2. **METHOD OF FIELD RESISTIVITY SURVEY AND SAMPLING FOR LABORATORY TESTS**
   
a. **Scope**

   The field resistivity test is an indication of the soluble salts in the soil or water; it is used primarily as a guide for selecting samples that will be tested in the laboratory. The natural soil in each channel or culvert location and the structural backfill material are tested by a portable earth resistivity meter, and samples are selected on the basis of these tests. These samples are tested in the laboratory using a soil box to determine the minimum resistivity that will be used in the culvert-type determination.

b. **Apparatus and Materials**

   1. Portable earth resistivity meter, suitable for rapid in-place determination of soil resistivity.
   2. Field probe(s).
   3. Steel starting rod, for making hole (in hard ground) for inserting probe(s).
   4. Sledge hammer 4 lbs. (1.8 kg).
   5. Distilled, deionized, or other clean water that has a resistivity greater than 20,000 ohm-cm.

c. **Recording Data**

   Record test data in a field record book for use in selecting samples and also for use in analyzing laboratory test data.

d. **Test Procedures**

   1. In the channel of a proposed culvert site, insert the field probe into the soil between 6 in. (152.4 mm) and 12 in. (304.8 mm) and measure the resistivity. Follow the manufacturer’s instructions for use of the meter. Remove the field probe and pour about 2 oz. (59 ml) of distilled water into the hole.

   2. Reinsert the probe while twisting to mix the water and soil, then measure the resistivity.
(3) Withdraw the probe and add an additional 2 oz. (59 ml) of distilled water.

(4) Reinsert the probe and again measure the resistivity of the soil.

(5) Multiply the lowest probe reading by ten to determine the minimum field soil resistivity and record this result. Note the multiplication factor of ten for soil resistivity readings when using the field probe. The multiplication factor for the soil box is one.

(6) In addition to the single probe method described above another method is available for determination of soil resistivity in the field. Refer to the manufacturer’s instructions as well as ASTM G 57 if the 4 probe “Wenner” method is being employed to determine the soil resistivity in the field.

e. Selection of Soil Samples for Laboratory Tests

(1) Make sufficient resistivity determinations at various locations in the channel or culvert site area to adequately represent the entire area. Should the soil appear consistent at a test site, take two resistivity determinations to verify. Additional readings should be taken if different soils are present.

(2) If the resistivity is reasonably uniform within the limits of the project, soil samples from three different locations will be sufficient. If, however, some locations show resistivities that differ significantly from the average of the determinations for the area being surveyed, additional soil samples should be taken to represent these locations — particularly those with resistivities significantly below the average.

For example, if the soil resistivities throughout the surveyed area are all at or near an average value of 20 ohm meter, three samples will be enough. If any of the locations tested have resistivities markedly below this average, for example 8 ohm meter, then such “hot spots” should definitely be represented by additional samples. Scattered locations of higher resistivity, for example 30 ohm meter or more, do not require additional samples.

Judgment must be exercised both in the field testing and sampling, and in evaluating the laboratory tests. In all cases, take a minimum of three samples per project. Samples should be about 1 lb. (.45 kg) each and should be identified as to material type and location.

3. METHOD OF DETERMINING pH OF WATER

a. Scope

This method is suitable for use in the field or laboratory for determining the pH of water samples.

b. Apparatus and Materials

(1) 5 oz. (148 ml) or larger nonmetallic wide-mouth container, e.g., glass jar, beaker, or wax-coated paper cup.

(2) pH meter.

(3) pH standard solution of pH 7.

c. Recording Data

Record test data in a field record book and report the results to the Project Engineer and in the Regional Soils Report.
d. Method of Sampling
   (1) To avoid contamination from container, dip the wide-mouth container into the water to be tested, swirl to rinse and pour out contents.
   (2) Dip the container into the water again to obtain a sample.
   (3) Pour off any film which is on the surface of the sample before testing.

e. Standardization of pH Meter
   Follow the instructions provided with the pH meter.

f. Use of pH Meter to Determine pH of Water
   Follow the instructions provided with the pH meter.

g. Precautions
   Follow the manufacturer’s instructions for use of the meter and observe the usual precautions for making chemical tests.

   **Note:** Field pH readings may be taken at any period other than flood flow. For water which has a pH of less than 6, take a 1 L (minimum) sample for laboratory analysis.

4. METHOD OF DETERMINING pH OF SOIL

a. Apparatus and Materials
   (1) Paper cups, 5 oz. (148 ml) wax coated type.
   (2) Teaspoon or small metal scoop.
   (3) Wash bottle containing distilled water.
   (4) pH meter.
   (5) pH standard solution of pH 7.

b. Recording Data
   Record data in a field record book and report the results to the Project Engineer and in the Regional Soils Report.

c. Preparation of Test Specimens
   (1) Place approximately five to six rounded teaspoonfuls (25 g) of the soil to be tested into a 5 oz. (148 ml) paper cup.
   (2) Add enough distilled water to make a slurry in the cup.
   (3) Disperse soil in water by stirring. Wait 5 minutes. The specimen is now ready for testing.

d. Standardization of pH Meter
   Follow the instructions provided with the pH meter.

f. Use of pH Meter to determine pH of Soil
   Follow the instructions provided with the pH meter.

f. Precautions
   If the pH reading is unstable when the electrode is immersed in the soil slurry, leave the electrode immersed until the pH reading has stabilized. In some cases, this waiting period for the stabilization of pH reading may be as long as five minutes.
5. LABORATORY METHOD OF DETERMINING MINIMUM RESISTIVITY

a. Scope

This method covers the procedure for determining the minimum resistivity of soil or water samples selected as indicated in Section 2.

b. Apparatus and Materials

(1) Resistivity meter suitable for laboratory testing.
(2) Soil box calibrated for use with resistivity meter.
(3) U.S. No. 8 (2.36 mm) sieve.
(4) Round tin pans, 12 in. (304.8 mm) diameter and 2 in. (50.8 mm) deep.
(5) Non-absorbent pans, bowls or other containers of sufficient size to eliminate spilling during mixing, moisture conditioning, and sample handling.
(6) 200°F (93°C) oven.
(7) Oven capable of maintaining a temperature of 140°F (60°C) around sample.
(8) Balance or scale, minimum 500 g capacity, accurate to 1 g.
(9) Balance, with sufficient capacity and readable to 0.1% of the sample mass, or better, and conform to the requirements of AASHTO M 231.
(10) Distilled or deionized water.
(11) Spoon or spatula.
(12) 10 ml graduated cylinder or graduated plastic cup.
(13) Graduated cylinder or other suitable device of sufficient size to accurately add quantities of moisture to sample.
(14) Mixing bowl.
(15) Straightedge

Preparation of Soil Samples

After thorough mixing of the sample, screen it on a U.S. No. 8 (2.36 mm) sieve. If the sample is too moist to be sieved, it should first be dried in a 140°F (60°C) oven and then crushed. Do not crush rocks. Only natural material passing the U.S. No. 8 (2.36 mm) sieve is to be used for the test.

Dry the sample as received from the field to a constant mass at a temperature not to exceed 140°F (60°C). (Air drying is also acceptable.) Split or quarter a sufficient amount of the dried material to yield a suitable sample after the material has been pulverized, taking care not to crush rock particles, and screened over a U.S. No. 8 (2.36 mm) sieve. Discard any material retained on the U.S. No. 8 sieve. Only natural material passing the U.S. No. 8 sieve is to be used for the test.
d. Measuring the Resistivity of Soil Sample

(1) Quarter or split out about 80 g of the material passing the U.S. No. 8 (2.36 mm) sieve.

(1) Split or quarter an amount of prepared soil that will fill approximately 4 times the volume of the soil box being utilized to determine resistivity.

(2) If the sample has been dried, add about 10 ml of distilled water to the 80 g of soil and thoroughly mix.

(2) Add approximately 10% by weight of distilled water to the sample and mix thoroughly. Allow the sample to stand in a moisture proof container for a minimum of 12 hours.

(3) After the soil sample is thoroughly mixed, place and compact it (moderate compaction with the fingers is sufficient) in the soil box.

(3) Re-mix the sample and immediately compact it (moderate compaction with the fingers is sufficient) slightly over the top of the soil box that has been cleaned with distilled water prior to use. Strike the material level to the top of the soil box with a straightedge.

(4) Measure the resistivity of the soil in accordance with the instructions furnished with the meter and record the value.

(4) Rinse the soil box with distilled water then immediately place the soil in the soil box and compact as described in step 3.

(5) Remove the soil from the soil box and add an additional 5 ml % by weight of distilled water and again thoroughly mix.

(5) Again place and compact the soil in the soil box and measure its resistivity.

(6) Rinse the soil box with distilled water then immediately place the soil in the soil box and compact as described in step 3.

(6) Repeat this procedure (Steps 5 and 6) one more time.

(7) Measure the resistivity of the soil in accordance with the instructions furnished with the meter and record the value.

(7) Repeat steps 5 through 7 until a minimum value can be determined.

(8) If the resistivity of the soil has not followed a trend of high resistivity, low resistivity, and then an increase in resistivity for the preceding additions of distilled water, continue to add water to the soil in about 3 ml increments, mixing, placing, compacting, and measuring resistivity for each increment, until a minimum resistivity is obtained. The amount of water added is not a rigid or exact figure and may vary depending on the soil type. The idea behind the increments is to gradually increase the moisture content until the minimum resistivity value is found.

(8) Repeat steps 5 through 7 until a minimum value can be determined.

(9) If the sample has not been dried, begin the test procedure by adding 3 ml of distilled water in lieu of the 10 ml specified in Step 2. Continue to add 3 ml increments of distilled water followed by mixing, placing, compacting, and measuring until a minimum value of resistivity is measured.

(10) Record the lowest value measured during the repeated measurements in the soil box. The multiplication factor for the soil box is one, (do not assume this as this value should be verified or reconciled with the manufacturer’s recommendations provided with the soil box) so a direct reading of the meter is the value used. The value thus obtained combined with the pH of the soil and water is used to determine what type of culvert is applicable to this location.
(11) *It is not necessary to continue determining resistivity values once the soil sample becomes liquid.* Deleted by DWB – this statement is pretty vague and could easily be misunderstood and test terminated early.

e. Measuring the Resistivity of a Water Sample
   
   (1) Thoroughly clean the soil box of all soil particles and rinse the soil box a minimum of three times with distilled water.
   
   (2) Fill the soil box with distilled water and measure its resistivity.
   
   (3) If the distilled water in the soil box measures infinite resistivity, empty the soil box of distilled water, fill with the test water, measure its resistivity, and record the measured value.
   
   (4) If the distilled water in the soil box measures less than infinite resistivity, continue to rinse with distilled water until the box is absolutely clean. This condition is indicated by an infinite resistivity measurement when the box is filled with distilled water.

f. Recording Data
   
   Record data in a field record book and report the results to the Project Engineer and in the Regional Soils Report.

g. Precautions
   
   The soil box must be completely filled and level with the top.

6. MINIMUM REQUIREMENTS

   a. Metal pipe may be used at locations where the pH and soil resistivity are within the limits specified in the *Hydraulics Manual* (M 23-01) for Aluminum (Aluminum Coated) Steel Pipe, Aluminum Pipe, and Galvanized (Zinc Coated) Steel Pipe.
Performance Exam Checklist
Method T 417 Checklist
Determining Minimum Resistivity and PH of Soil and Water

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Determining ph of soil</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. 5 to 6 rounded teaspoons full of soil placed in a 5 oz. (148 ml) paper cup?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Enough distilled water added to make a slurry in the cup?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Soil Dispersed in water by stirring. Let stand for 5 minutes?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. PH meter is standardized?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. PH of soil determined by meter?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>If ph reading unstable allow electrode to sit until stabilized. This may take 5 minutes.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Record results</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td><strong>Laboratory method of Determining Minimum Resistivity</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Sample thoroughly mixed and screened through # 8 sieve?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Note: If too moist to screen, dry @ 140 F and screen.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Quarter or split out 80 grams of passing #8 material</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. If sample was dried add 10 ml of distilled water to the 80 gram and thoroughly mix?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Place and compact sample in soil box. Must be level with top of soil box.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Resistivity measured with the instrument?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Remove soil from box, add 5 ml of distilled water and thoroughly mix?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Place soil in soil box and measure resistivity?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Repeat procedure by increasing moisture content until minimum resistivity can be established?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. If sample was not dried, begin with 3 ml of distilled water added and mixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. Continue with 3 ml of moisture added per resistivity reading until minimum reading can be established?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>11. Record the lowest value during the repeated measurements?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>12. Report the resistivity reading and the ph value?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>
First attempt: Pass ☐ Fail ☐  Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

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WSDOT Test Method T 418

Method of test for Corrosion of Deicing Materials

1. SCOPE

The weight loss of steel coupons subjected to corrosion by deicing salts for a given period of time is determined. This weight loss is converted to mils penetration per year of the steel surface by the deicing salt. The ratio of the deicing salt sample to a sodium chloride control sample is reported to provide a means to evaluate and compare the corrosion rates of different deicing salts. (This procedure is a modification of NACE Standard TM-01-69 (1976 Revision)).

2. APPARATUS

a. Timer, adjustable, that can be set for a 50 minute (up position) and a 10 minute (down position) cycle every hour, and able to cycle for 72 hours.
b. Immersion testing device capable of automatically subjecting the coupons to an immersion/no immersion cycle. A device with a bar, configured like a crank to which a line from the coupon holder is attached, with an electric motor, governed by the timer, that cycles the crank up and down, is one type of tester.
c. 500 Erlenmeyer flasks fitted with one-hole rubber stoppers, one flask for each sample plus one for a distilled water control and one for a sodium chloride standard.
d. Non-corrosive (e.g., polyethylene) coupon holders attached to the immersion testing device with non-corrosive line (e.g., polypropylene fish line). Each coupon holder to hold three coupons.
e. Coupons, non-galvanized 1/2 in. (13 mm) flat steel washers having the approximate dimensions of 1.38 in. (35 mm) outside diameter by 0.56 in. (14 mm) inside diameter by 0.12 in. (3 mm) thick. Coupons must meet ASTM F436, Type 1, with a Rockwell Hardness of C 38-45. Three coupons are needed for each sample.
f. Polypropylene bottlers, for sample dissolution, one for each sample.
g. Balance, accurate to 0.0001 g.
h. Metal stamp numbering set.
i. Dial Caliper to measure coupons, accurate to 0.01 mm.

3. REAGENTS

a. Hydrochloric Acid cleaning solution. Make by dissolving 50 g SnCl₂ (stannous chloride) and 20 g SbCl₃ (antimony trichloride) in 1000 ml of concentrated Hydrochloric Acid.
b. Distilled water conforming to ASTM D 1193 Type II.
c. Chloroform, technical grade.
d. Acetone, technical grade.
e. Sodium Chloride standard. Make a 3 percent by weight solution of reagent grade Sodium Chloride in distilled water.
f. Hydrochloric Acid, 1 + 1.
4. PREPARATION OF THE COUPONS
   a. Wipe each coupon with a suitable solvent to remove grease and oil.
   b. Examine each coupon for metallurgical abnormalities and reject those that are suspect to flaws.
   c. Acceptable coupons are stamped for identification.
   d. Coupons are acid etched with 1+1 HCl for approximately 2-3 minutes.
   e. The coupons are quickly rinsed with tap water, then distilled water, wiped dry and placed in chloroform.
   f. When the coupons are removed from the chloroform for use, they are placed on a paper-lined tray (not touching each other) and allowed to air dry in a ventilated hood for a minimum of 15 minutes.
   g. Coupons are measured as specified in Section 5. (Note: If latex gloves are not worn during measuring, the coupons should be rinsed again and dried as prescribed above prior to weighing. This will remove any oils that may be transferred to the coupons.)
   h. Each coupon shall be weighed to a constant weight. The constant weight is obtained when two consecutive weighings of each coupon are within a minimum of 0.5 milligrams of each other. Removal of incidental flash rusting prior to weighing is not necessary.
   i. Three coupons are used in each chemical product solution and for the distilled water and Sodium Chloride controls.

5. MEASURING THE COUPONS
   The outside diameter, inside diameter, and the thickness of each coupon is measured twice at 90 degrees from each initial reading and the averages calculated for each measurement. The averages are than used to calculate the surface area of each coupon with the following formula:
   \[ A = \frac{3.1416}{2} (D^2 - d^2) + 3.1416(t)(D) + 3.1416(t)(d) \]
   Where
   \[ D = \text{average outside diameter} \]
   \[ d = \text{average inside diameter} \]
   \[ t = \text{average thickness} \]

6. SAMPLE PREPARATION
   Make a 3 percent by weight solution of each deicing salt by weighing 30.00 g (as received) of the sample and dissolving in 970.00 g of distilled water. Allow the solutions to sit a minimum of 12 hours in polyethylene bottles to insure maximum solubility and to allow for any reactivity (i.e., heat of hydration and heat of solution).

7. PROCEDURE
   Approximately 300 milliliters of the deicing solution as mixed with distilled water is placed into a 500 ml Erlenmeyer flask. Each flask is equipped with a rubber stopper that has been drilled to allow a line to run through it. One end of the line is attached to a rotating bar and the other end of the line is attached to a plastic frame made to hold the three coupons inside the flask. The rotating bar is controlled by an electric timer that lowers the coupon holding apparatus into the solution for 10 minutes then raises the coupon holding apparatus out of the solution for 50 minutes but still keeps the coupons inside of the flask for the entire duration of the test. The corrosion test is run for 72 hours. No agitation of the solution is made during the corrosion test.
Corrosion tests are conducted at normal room temperature. The room temperature is to be recorded daily during the operation of the test. The room temperature shall be taken with a calibrated thermometer located next to the corrosion-testing instrument.

8. CLEANING OF THE COUPONS

The coupons are removed from the solution after 72 hours. They are placed in glass beakers containing the Hydrochloric Acid cleaning solution. (Note: The fumes given off by the acid during cleaning contain gases formed from the antimony and are extremely hazardous, this portion of the cleaning must be conducted under a fume hood).

After 15 minutes of cleaning the coupons are removed from the cleaning acid, rinsed with tap water, then distilled water, and wiped with a cloth to clean any deposits from the coupons. They are then returned to the cleaning acid and the procedure is repeated. After cleaning, the coupons are rinsed in chloroform, air-dried, and weighed.

Each coupon shall be weighed to a constant weight. Constant weight is obtained when two consecutive weighings of each coupon are within a minimum of 0.5 milligrams of each other.

9. EVALUATION OF CORROSION

The weight loss of each coupon is determined by subtracting the final weight from the original weight. The corrosion rate for each coupon is expressed as mils penetration per year (MPY) by the following formula:

\[
\text{MPY} = \frac{(\text{weight loss (milligrams)})(534)}{(\text{area})(\text{time})(\text{metal density}^*))}
\]

* metal density for steel is 7.85 g/cc

The final MPY value for each solution is determined by calculating an average of the three individual coupons. (Note: Wide variation of MPY of individual coupons inside the same flask typically indicates contamination of a coupon. If variation of individual MPY is too great to determine consistent data the test should be repeated. Typically coupon variation may run plus or minus 3 MPY).

10. REPORTING RESULTS

Results shall be reported in Percent Effectiveness (also referred to as “Percent as Corrosive as NaCl”). Results equal to or less than 30% are passing.

The MPY is determined as above for the deicing salt samples, the distilled water blank and for the sodium chloride standard.

The MPY determined for the distilled water blank is subtracted from the MPY value obtained for the deicing samples to arrive at an adjusted deicing sample MPY. Likewise, the distilled water blank is subtracted from the MPY value of the sodium chloride standard to obtain the adjusted NaCl standard MPY.

The adjusted deicing sample MPY is divided by the adjusted NaCl standard MPY and multiplied by 100 to obtain the corrosion of the sample as a percent of the sodium chloride corrosion result.

\[
\text{% Effectiveness} = \frac{\text{adjusted deicing sample MPY}}{\text{adjusted NaCl standard MPY}} \times 100
\]
WSDOT Test Method T 419

Test Method for Cold Temperature Impact Resistance of the Plastic Coating on Reinforcing Bar Chair Feet

1. SCOPE

This test method covers the resistance to cracking, splitting, or breaking of the plastic coating on rebar chair feet under specified conditions of impact by means of a falling weight.

2. REFERENCE DOCUMENTS

ASTM D 4495 IMPACT RESISTANCE OF PVC by means of a falling weight is used as a guide to the order and method of testing, with the changes as below.

3. TERMINOLOGY

Failure is any breach of integrity of the plastic feet coating such that moisture could be admitted to the steel rebar chair.

4. SIGNIFICANCE AND USE

The plastic feet coating must protect the steel rebar chair from exposure to moisture that could be present under the concrete pad, to keep corrosion from traveling up the chair and corroding the rebar mat in the concrete. The impact represents the stresses placed on the feet as the rebar is dropped on the chair.

5. APPARATUS

a. Impact Tester — Consists of a tube 17 in. (43.18 cm) high with a 1½ in. (4.76 cm) inside diameter, and a 1 in. (2.54 cm) wide by 2 in. (5.08 cm) high slot cut out of the bottom end for access in positioning the sample.

b. Falling Weight — Shall be a solid cylinder made from steel, 1½ in. (3.81 cm.) (nominal) in diameter, and approximately 8 ½ in. (21.6 cm) long, with a flat-bottom surface that strikes the test specimen. The mass of the falling weight shall be 5±0.5 lbs (2.26±0.22 kg). The weight shall have a line inscribed on it approximately 5 in. (12.7 cm) from the bottom striking surface such that when the line is even with the top edge of the impact tube, the bottom end of the weight is 12 in. (30.48 cm) above the sample. This will give an impact of 5 ft./lbs. (6.78 J) upon the sample.

c. The impact tester is positioned on a metal plate placed upon a flat, stable surface, i.e., a concrete floor.

6. TEST SPECIMENS

a. Three plastic coated feet constitutes one test.

7. CONDITIONING

a. Condition the test specimens at 0±3.5°F (−18±2°C) in a freezer for a minimum of six hours.
8. PROCEDURE
   a. Cut the feet off the chair about \( \frac{1}{2} \) in. (1.27 cm) above the plastic coating. One test comprises three plastic coated feet randomly chosen from the chair. Put specimens into the freezer as described in number 7.
   b. Remove the specimens, one at a time, from the freezer, and place on the metal plate. Position the impact tube over it, line the weight up in the tube so that the inscribed line is even with the top edge of the impact tester tube, then release the weight (Note 1). Remove the weight and tube and examine the specimen. Any cracks, splits, or other breaks are recorded as a failure. Test the remaining two specimens and record results.

9. RESULTS
   A satisfactory test is when two out of the three specimens do not crack, split, or reak.

   Note 1: Care must be taken to ensure that the weight falls freely, it must not “chatter” down the tube.
WSDOT Test Method T 420

Test Method for Determining the Maturity of Compost (Solvita Test)

1. SCOPE
   The Solvita test is used for evaluating compost conditions.

2. REFERENCE DOCUMENTS
   AASHTO T-2

3. TERMINOLOGY
   3.1 Definitions
   3.1.1 Compost shall be stable, mature, decomposed organic solid waste that is the result of the accelerated, aerobic biodegradation and stabilization under controlled conditions. The result is a uniform dark, soil-like appearance.
   3.1.2 Maturity of any compost sample may be judged using both color test results from paddle A and C. Paddle A is a styrene paddle with a gel component that measures the ammonia content of the compost. Paddle C is a styrene paddle with a gel component that measures the carbon dioxide emitted by the compost sample.

4. SUMMARY OF TEST METHOD
   There are three easy steps involved in using the Solvita test kit to evaluate compost.
   4.1 Obtain and prepare the sample.
   4.2 Perform the test by placing both Solvita gel-paddles in the jar.
   4.2 Determine compost maturity using the color keys provided in the kit.

5. SIGNIFICANT AND USE
   This test is used to determine the maturity of compost materials delivered in the field for use. This test measures the amount of ammonia and carbon dioxide in the compost.

6. APPARATUS
   6.1 Solvita Kit containing the following:
      a testing jar with lid
      a carbon-dioxide paddle (marked with “C”) is purple
      an ammonia paddle (marked with “A”) is yellow
      color determination charts
   6.2 Shovel
   6.3 Small trowel or spoon
   6.4 A clean container large enough to combine the sample (approximately 5 gallons)
   6.5 A clean surface for mixing the sample such as a tarp or plywood
7. **SAMPLE PREPARATION**

7.1 A composite sample (approximately 1 cubic foot) representing the lot to be tested should be sampled in accordance with AASHTO T-2 “Sampling from Stockpiles” or “Sampling from Transport Units”.

7.2 Place the sample on a hard, clean, level surface where there will be neither loss of material nor the accidental addition of foreign material.

7.3 Particles such wood chips which are too large for the jar (over 1/2 inch) should be removed or screened from the compost sample.

7.4 Checking for optimal moisture is absolutely necessary for accurate maturity testing. Samples which are either too wet or too dry are not likely to produce accurate results. The moisture level should be judged by the squeeze test before proceeding. Perform the Squeeze test by squeezing a small handful of compost. When squeezed tightly the compost should feel wet without producing any free water. Compost that is too dry is dusty and will not clump with hard squeezing.

7.5 Mix the material thoroughly by turning the entire sample over three times. With the last turning, the entire sample shall be placed into a conical pile.

7.6 Using a small trowel, or other device, remove a portion from the center of the pile.

7.7 Fill the jar to the fill line and obtain proper density by sharply tapping the bottom of the jar on a counter. Fluffy or coarse composts should be compacted by pressing firmly into the jar.

7.8 If compost to be tested is in an optimal state, allow to air out for one hour.

7.9 If compost to be tested is not in an optimal state, then the following should be performed:

1. If the sample is hot, it should be covered and allowed to cool to room temperature before testing.

2. If the sample is too wet, it should be dried until it passes the squeeze test.

3. If the sample is too dry, add clean water until it passes the squeeze test. This sample shall be covered and allowed to stand at room temperature for 24 hours before performing the test.

8. **PROCEDURE**

8.1 Open each package by tearing along the top strip and carefully remove the paddle by grasping the handle. *Do not touch the special gel surface, and don’t allow compost to touch it.* Once the gel pack is opened, the test should be started within 30-minutes. The gel is not harmful to touch, but should be kept out of the mouth and eyes.

8.2 Insert the paddles into the sample at right angles to each other so that they can be seen through the viewing side. The edges of the paddles can be touching in the middle. Position the two paddles as indicated by the color squares on the jar label. Push the paddle tips into the compost to the bottom of the jar. Be careful not to jostle or tip the jar. Do not use a paddle if the gel is dried out or if the color is not the “Control Color” indicated on the respective color charts.
8.3 Screw the lid on tight, and keep the jar at room temperature 68-77º F (20-25º C) out of direct sunlight for 4 hours ± 10 minutes.

9. EVALUATING THE RESULTS

9.1 Read the Solvita paddle colors 4 hours after the test is started. To read the colors, observe the paddles through the viewing side of the jar with the lid in place and illuminated from the front. Color rendition is best in moderate-intensity, fluorescent room light. Compare to the color charts provided with the kit, and record the color numbers that most closely match. Since the Solvita colors may continue to change after 4-hours, the proper interpretation for this test is based on a 4-hour ± 10 minute reading.

10. REPORTS
Performance Exam Checklist

_Determining the Maturity of Compost (Solvita Test)_

_WSDOT Test Method T 420_

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

**Sample Preparation**

1. Representative sample obtained per AASHTO T-2? | ☐   | ☐  |
2. Sample placed on clean hard surface? | ☐   | ☐  |
3. Check for optimal moisture? | ☐   | ☐  |
4. Sampled mixed thoroughly? | ☐   | ☐  |
5. Small sample taken from the center of the pile? | ☐   | ☐  |
6. Sample filled in jar to the proper line and compacted? | ☐   | ☐  |
7. Sample allowed to air out for 1 hour or equilibrate for 24 hours | ☐   | ☐  |

**Procedure**

1. Open the gel packs with out touching the gel sticks? | ☐   | ☐  |
2. Is the test started within 30 minutes of opening the gel pack? | ☐   | ☐  |
3. Are the paddles inserted in the compost at right angles to each other? | ☐   | ☐  |
4. Are the paddles positioned to be seen through the viewing window? | ☐   | ☐  |
5. Are the paddles pushed to the bottom of the jar? | ☐   | ☐  |
6. Is the lid screwed on tight? | ☐   | ☐  |
7. Is the jar at room temperature 68-77 F? | ☐   | ☐  |
8. Is the test run for 4 hours ± 10 minutes? | ☐   | ☐  |
9. Maturity determined per Manufacturers instructions? | ☐   | ☐  |

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner ________________________________
WSDOT Test Method T 421

Test Method for Traffic Controller Inspection and Test Procedure

1. SCOPE

The purpose of this procedure is to provide a documented method for the steps involved with inspection and testing of the completed traffic controller cabinets.

2. REFERENCE DOCUMENTS

WSDOT Standard Specifications 9-29.13
NEMA Publication TS-1
FHWA Publication IP-78-16
Caltrans: Traffic Signal Controller Equipment Specification

3. SAFETY

Use proper equipment to reduce the risk of electrical shock.

4. APPARATUS

Equipment as defined to perform WSDOT Test Methods T 422, T 423, T 424, T 425, and T 427.
Resistor load bank to simulate each traffic signal light (150 Ohm 100W wire wound resistors).

5. PROCEDURE

The traffic controller cabinet shall be inspected to ensure that it is in compliance with the contract documents. Ensure that all of the required equipment is installed and the cabinet and meets the requirements of the contract documents. The results of successful completion of this procedure will be acceptance for testing.

As a minimum, the following items shall be inspected:

1) Mylar and four cabinet prints
2) Manuals
3) Labeling
4) Air filter
5) All wire laced and clamped
6) Field wire terminal blocks
7) Door locks
8) Police keys
9) Police panel switches
10) Circuit breakers
11) Transient voltage suppresser
12) Modem(s)
13) Phone jack for modem
14) Radio interference suppresser
15) Door light switch(s)
16) Pedestrian switches
17) Cabinet light
18) 120 VAC receptacle outlet
19) Ground fault interrupter
20) Equipment/Cabinet clearance
21) Load switches
22) Intersection display panel
23) Isolated 120VAC bus bar (neutral)
24) Phase Selectors
25) Flash transfer relays
26) Supplemental resistor loads
27) Two position door stop  
28) Emergency indicator lights  
29) Railroad preemption  
30) Cabinet Construction  
31) Detector Panel  
32) Detector Panel shorting plug (NEMA)  
33) Plastic document envelope (NEMA)  
34) DB9 socket and C20 plug (170)  
35) C-2 plug and cable (170)  
36) Rack mounted document drawer (170)  
37) Verify circuit breakers capacities  
38) Absence of red assembly (170)  
39) Controller  
40) Software  
41) AC isolator (170)  
42) DC isolator (170)  
43) Aux. file (170)  
44) External logic (NEMA)  
45) CMU door interlock switch  
46) Stop Time switch  
47) Cabinet Ground Bus bar  
48) Conflict Monitor  
49) Inside auto/flash switch  
50) Loop Amplifiers  

a. The traffic controller cabinet shall be connected to the resistor load bank.  
b. WSDOT Test Method T 425, Environmental Chamber Test shall be performed. Any deficiencies shall be documented on the test report.

6. REPORT  
Record any response found to be in disagreement with the published standards, report pass or fail and any corrective actions taken on the test report.
Performance Exam Checklist
Method T 421 Checklist
Test Method for Traffic Controller Inspection and Test Procedure

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Cabinet inspected for damage during shipping.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Letter to project office sent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Traffic controller assessed for compliance with contract provisions.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Simulated load connected to the Controller.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Perform Environmental test WSDOT Method T 425</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner __________________________________________

Comments:
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WSDOT Test Method T 422

Test Method for Traffic Controller Transient Voltage Test (Spike Test) Procedure

1. SCOPE

This test method is intended to evaluate traffic signal controllers for the transient voltage test defined in NEMA TS-1 2.2.3.2, as modified. This test is to be performed with the surge protector or line filter properly connected in the cabinet.

2. REFERENCE DOCUMENTS

NEMA Publication TS-1

3. SAFETY

Use proper equipment to reduce the risk of electrical shock.

4. APPARATUS

Transient voltage generator capable of placing a 300 V spike on the power supplied to the controller cabinet.

5. PROCEDURE

a. Program the controller unit to cycle on minimum recall.

b. Connect the controller cabinet to the transient voltage generator as outlined by the manufacturer.

c. With the spike generator set to minimum, apply power to the system. Ensure the controller unit is operating normally.

d. Adjust the generator output so that a 300 V ± 5% positive or negative pulse with a 1 microsecond rise and a 10 microsecond width pulse(spike) is placed on the input power, for 10 minutes.

e. Reverse the polarity from step 5d, reapply the pulse for 10 minutes.

f. Reduce the spike to minimum, then disconnect the power.

g. Restore normal power supply to the cabinet.

h. During the preceding transient test the controller unit must continue its programmed functions. During phase cycling, the controller unit shall not skip intervals; shall not place false calls or produce false indications while in dwell; shall not disrupt normal sequences in any manner; or shall not change timings.

6. REPORT

Record any response found to be in disagreement with the published standards, report pass or fail and any corrective actions taken on the test report.
## Performance Exam Checklist

### Method T 422 Checklist

**Test Method for Traffic Controller Transient Voltage Test (Spike Test) Procedure**

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Program controller for minimum recall</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Connect line noise generator to controller per manufacturer's recommendations.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. With the noise generator set to minimum verify controller is operating normally.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Adjust noise generator for 300V spike.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Reverse polarity and repeat 4</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Reduce spike to minimum and restore normal power.</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner ________________________________

Comments:

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WSDOT Test Method T 423

Test Method for Traffic Controller Conflict Monitor Testing

1. SCOPE

This test method is intended to evaluate traffic signal controller conflict monitors for various simulated conflicting signal indications to ensure that the conflict monitor is effective in sensing conflicting signal conditions.

2. SIGNIFICANCE OF USE

This test method describes procedures to be used for simulating conflicting signals to the conflict monitor, and ensuring the proper response. As an alternative to this procedure, manufactured test equipment designed to tests all parameters can be used.

3. REFERENCE DOCUMENTS

NEMA Publication TS-1
FHWA Publication IP-78-16
Caltrans: Traffic Signal Controller Equipment Specification

4. SAFETY

Use proper equipment to reduce the risk of electrical shock.

5. APPARATUS

A conflict monitor tester capable of testing to NEMA standards (see NEMA Publication TS-1 section 2.2.9) and/or Caltrans standards (see Caltrans Traffic Signal Control Equipment Specifications article 4) as required, such as ATSI DCMT-700 or ATSI PCMT-2000 Conflict Monitor.

6. PROCEDURE

Perform testing as shown in NEMA TS1 section 2.2.9.3 or follow manufacture's recommendations.

7. REPORT

Record any response found to be in disagreement with the published standards. Report pass or fail and any corrective actions taken on the test report.
Performance Exam Checklist
WSDOT Test Method T 423
Test Method for Traffic Controller Conflict Monitor Testing

Participant Name ___________________________________________ Exam Date _____________

Procedure Element Yes No
1. Remove Conflict Monitor from controller cabinet. □ □
2. Configure CMU tester for CMU to be tested. □ □
3. Connect CMU to tester. □ □
4. Perform CMU test as defined in appropriate manufactures test procedures. □ □
5. File raw data as required. □ □

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner ___________________________________________

Comments:

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T 423 January 2004 T 423 Page 3 of 4
WSDOT Test Method T 424

Test Method for Traffic Controller Power Interruption Test Procedure

1. SCOPE

   This test method is intended to evaluate traffic signal controllers for the Power Interruption test defined in NEMA TS-1 2.2.7, as modified, and is not required for the Type 2070 controllers. This test is to be performed at normal voltage and room temperature.

2. REFERENCE DOCUMENTS

   NEMA Publication TS-1

3. SAFETY

   Use proper equipment and training to reduce the risk of electrical shock

4. APPARATUS

   A power interrupting device capable of interrupting the controller power for supplied to the controller cabinet of predetermined time intervals as defined in NEMA TS1.

5. PROCEDURE

   a. Program the controller unit to cycle on minimum recall.

   b. Connect the controller cabinet to the power interrupting device as outlined by the manufacture's instructions.

   c. Apply power to the system with the power line modifier connected as required.

   d. **NEMA** Remove the input voltage for a period of 450 milliseconds. Upon restoration of the input voltage, check to insure that the controller unit continues normal operation as though no power interruption had occurred. Repeat this test three times.

   e. **Type 170** Remove the input voltage for a period of 1500 milliseconds. Upon restoration of the input voltage, check to insure that the controller unit continues normal operation as though no power interruption had occurred. Repeat this test three times.

   f. **NEMA** Remove the input voltage for a period of > 500 milliseconds. Upon restoration of the input voltage, check to insure that the controller unit reverts to its start up sequence. Repeat this test three times.

   g. **Type 170** Remove the input voltage for a period of >2000 milliseconds. Upon restoration of the input voltage, check to insure that the controller unit reverts to its start up sequence. Repeat this test three times.

   h. Restore normal supply to the cabinet.

6. REPORT

   Record any response found to be in disagreement with the published standards. Report pass or fail and any corrective actions taken on the test report.
Performance Exam Checklist

WSDOT Test Method T 424

Test Method for Traffic Controller Power Interruption Test Procedure

Participant Name _______________________________ Exam Date ____________

Procedure Element

1) Program traffic controller for minimum recall
   Yes ☐ No ☐

2) Connect traffic controller to power interrupter per the manufacturer’s recommendations.
   Yes ☐ No ☐

3) Verify traffic controller operates normally for the prescribed interruption.
   Yes ☐ No ☐

4) Verify traffic controller reverted to the start-up sequence.
   Yes ☐ No ☐

5) Restore normal power to the cabinet.
   Yes ☐ No ☐

6) Document test results on test report.
   Yes ☐ No ☐

First attempt: Pass ☐ Fail ☐
Second attempt: Pass ☐ Fail ☐

Signature of Examiner ________________________________

Comments:

________________________________________________________________________
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________________________________________________________________________
WSDOT Test Method T 425

**Test Method for Traffic Controller NEMA and 170 Type Environmental Chamber Test**

1. **SCOPE**

   This test method is intended to evaluate the traffic signal controller to the temperature and environmental extremes as defined in the NEMA TS-1 Standard. This procedure will cover temperatures from minus 30º F (-34º C) to 165º F (74º C) and voltages from 95 VAC to 135 VAC with a power interruption as defined in NEMA TS-1.

2. **REFERENCE DOCUMENTS**

   NEMA Publication TS-1
   FHWA Publication IP-78-16
   Caltrans: Traffic Signal Controller Equipment Specification

3. **SAFETY**

   The environmental chamber provides extreme temperatures. Caution should be exercised to avoid injury.

4. **APPARATUS**

   A suitable chamber in which the traffic controller can be subjected to the specified temperatures (-30º F and 165º F) and provide safe access to the equipment under test. A temperature recording device shall record the temperature in the chamber during the test with an accuracy of ±3º F. The air inside the chamber shall be circulated so that no more than a 3º F difference will occur. The chamber control shall maintain constant absolute humidity from 109º F to 165º F.

   Variable voltage transformer capable of delivering the power required at the voltages defined in NEMA TS-1 (20 amps at 0 to 150 VAC)

   Volt-Ohm-Milliamp meter (VOM)

   Resistance load device to simulate each traffic signal light (150 ohm 100W wire wound resistors)

5. **PROCEDURE**

   5.1 Low-Temperature Low-Voltage Test:

      5.1.1 Test conditions:

      a. Environmental chamber door closed

      b. Temperature: minus 30º F

      c. Low Voltage: 95 VAC

      d. Equipment cabinet door open

      e. Humidity control off
5.1.2 Procedure:

5.1.2.1 While at room temperature, adjust the input voltage to 95 VAC NEMA & Type 170 & 2070 + modified 2010ECL; 102VAC 2070 + standard 2010ECL and verify that the test unit is operational.

5.1.2.2 With the equipment under test cycling on minimum recall, lower the test chamber to –30º F at a rate not to exceed 30º F per hour. With the humidity control off, allow the controller assembly under test to cycle on minimum recall during the time it takes to cool down the chamber.

5.1.2.3 Then operate the test switches listed in TABLE 1 to ensure their proper operation.

5.1.2.4 NEMA only remove power from the controller assembly for a period of 3 hours.

5.1.2.5 Upon restoration of power, the controller assembly shall resume cycling at minimum recall.

5.1.2.6 Upon satisfactory completion of this test proceed to the Low-Temperature High-Voltage Test.

5.2 Low-Temperature High-Voltage Test

5.2.1 Test conditions:
   a. Environmental chamber door closed
   b. Temperature: minus 30º F
   c. High Voltage: 135 VAC
   d. Equipment cabinet door open
   e. Humidity control off

5.2.2 Procedure:

5.2.2.1 While at –30º F and the humidity control off, adjust the input voltage to 135 VAC and allow the controller assembly to cycle on minimum recall.

5.2.2.2 Then operate the test switches listed in TABLE 1 to ensure their proper operation.

5.2.2.3 With the input voltage at 135 VAC bring the chamber and test controller assembly to room temperature at a rate no greater than 30º F per hour.

5.2.2.4 Upon satisfactory completion of this test proceed to the High-Temperature High/Low Voltage Test.

5.3 High-Temperature High/Low Voltage

5.3.1 Test Conditions
   a. Environmental chamber door closed
   b. High temperature 165 F
   c. High voltage 135 VAC
   d. Equipment door open
   e. Humidity control as given in TABLE 2
5.3.2 Procedure:

5.3.2.1 With the controller assembly cycling on minimum recall, raise the test chamber temperature to 165° F at a rate not to exceed 30° F per hour. Check to see that the input voltage is set at 135 VAC.

5.3.2.2 Set the humidity controls to not exceed 95% relative humidity over the temperature range of 40° F to 110° F. When the temperature reaches 109° F readjust the humidity control to maintain constant absolute humidity.

5.3.2.3 Verify that the controller assembly continues to cycle satisfactorily during the period of temperature increase and established levels of relative humidity.

5.3.2.4 Allow the test unit to cycle on minimum recall upon reaching 165° F at 18% relative humidity. Then operate the test switches listed in TABLE 1 to ensure their proper operation.

5.3.2.5 Allow test unit to cycle for a minimum of 2 hours at 165° F and 18% relative humidity and 135 VAC.

5.3.2.6 With the test unit at 165° F and 18% relative humidity, again operate the test switches listed in TABLE 1 to ensure their proper operation.

5.3.2.7 Lower the voltage to 95 VAC NEMA & Type 170 & 2070 + modified 2010ECL; 102VAC 2070 + standard 2010ECL. Bring the chamber and controller assembly back to room temperature at a rate no greater than 30° F per hour.
6. REPORT

Record any response found to be in disagreement with the published standards. Report pass or fail and any corrective actions taken on the test report.

**Table 1**

<table>
<thead>
<tr>
<th>Switches</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Verify function of intersection display panel switches</td>
</tr>
<tr>
<td>2 Verify function of police panel switches (on/off, auto/flash)</td>
</tr>
<tr>
<td>3 Verify stop time switch function (inside)</td>
</tr>
<tr>
<td>4 Verify auto/flash switch function (inside)</td>
</tr>
<tr>
<td>5 Reserve for future use</td>
</tr>
<tr>
<td>6 Verify function of external logic (NEMA)</td>
</tr>
<tr>
<td>7 Verify operation of loop detection panel</td>
</tr>
<tr>
<td>8 Verify function of preemption push button on door (NEMA)</td>
</tr>
<tr>
<td>9 Verify function of preemption switches on phase selectors</td>
</tr>
<tr>
<td>10 Verify operation of emergency indicator light</td>
</tr>
<tr>
<td>11 Verify CMU functioning properly</td>
</tr>
</tbody>
</table>

**Table 2**

<p>| Wet-Bulb Dry-Bulb Relative Humidity at Barometric Pressure of 29.92 in. hg. |
|-----------------|-----------------|-----------------|-----------------|</p>
<table>
<thead>
<tr>
<th>Dry Bulb °F</th>
<th>°C</th>
<th>Relative Humidity, Percent*</th>
<th>Wet Bulb °F</th>
<th>°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>40</td>
<td>4.4</td>
<td>75</td>
<td>37</td>
<td>2.8</td>
</tr>
<tr>
<td>50</td>
<td>10.0</td>
<td>80</td>
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<td>60</td>
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<td>70</td>
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<tr>
<td>80</td>
<td>26.7</td>
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<td>90</td>
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<td>100</td>
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<td>38</td>
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<td>150</td>
<td>65.6</td>
<td>28</td>
<td>109</td>
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<tr>
<td>160</td>
<td>71.1</td>
<td>21</td>
<td>109</td>
<td>42.8</td>
</tr>
<tr>
<td>165</td>
<td>73.9</td>
<td>18</td>
<td>109</td>
<td>42.8</td>
</tr>
</tbody>
</table>

*For dynamic testing.
Performance Exam Checklist  
WSDOT Test Method T 425  
Test Method for Traffic Controller NEMA and 170 Type Environmental Chamber Test

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Place Traffic Controller into the Environmental chamber.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Perform Low-Temperature Low Voltage test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Perform Low-Temperature High Voltage Test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Perform High-Temperature Low voltage test.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Perform High-Temperature High voltage test.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner ________________________________

Comments:

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WSDOT Test Method T-426

Pull-Off Test for Hot Melt Traffic Button Adhesive

1. SCOPE
   This method describes a procedure for determining the force required to pull a Type 1 raised pavement marker, adhered with hot melt button adhesive, from a portland cement mortar surface. This method is also suitable for asphalt concrete surface.

2. APPARATUS AND MATERIALS
   a. Portland Cement mortar block (approximately 8” x 18” by 2”) – commercially available block with a compressive strength of 3000 to 5000 psi, conditioned for 24 hours at standard laboratory conditions prior to test.
   b. Asphalt Concrete puck – 6 inch diameter gyratory compacted asphalt concrete puck, conditioned for 24 hours at standard laboratory conditions prior to test.
   c. Raised pavement marker – WSDOT Type 1 plastic or thermoplastic, drilled in the center to accept a threaded steel rod. The type 1 markers shall be from at least two different manufacturers with different patterns on the bottom of the markers.
   d. Laboratory melter – As described in ASTM D5167.
   e. Drill press with bit sized for the threaded rods described below.
   f. Threaded steel eye bolt for attaching to the raised pavement marker.
   g. Tensile testing apparatus – As described in AASHTO T 237 Section 13, fitted with a threaded steel rod with a 2” hook.

3. PROCEDURE
   a. Pull-off tests shall be run in triplicate.
   b. Hot melt traffic button adhesive shall be heated in the laboratory melter to the manufacturer’s recommended application temperature.
   c. A quantity of adhesive sufficient to squeeze out a small bead around the entire periphery of a 4” button shall be poured onto a portland cement mortar block (or asphalt concrete puck) and a predrilled raised pavement marker shall be seated on the adhesive and allowed to cure for at least 4 hours.
   d. A threaded steel eye bolt shall be inserted into the predrilled hole in the button.
   e. The block (puck) and button shall be placed in the tensile testing apparatus and the threaded hook shall be inserted in the eye bolt.
   f. Load shall be applied slowly until failure and the maximum load shall be recorded.
4. **CALCULATION**
   The pull-off strength shall be calculated as follows:
   
   Pull-off Strength, psi = \( L/A \)
   
   \( L = \) Maximum load, pounds
   
   \( A = \) Surface area of Pavement marker (in\(^2\))

5. **REPORT**
   
   The pull-off strength reported shall be the average of the three determinations on a portland cement mortar block unless otherwise specified.
WSDOT Test Method T 427

Test Method for Loop Amplifier Testing Procedure

1. SCOPE
   This test method is intended to evaluate operation of individual loop amplifiers that are included with the traffic controller cabinet.

2. REFERENCE DOCUMENTS
   WSDOT Standard Specification 9-29.13
   NEMA Publication TS-1

3. SAFETY
   Use proper equipment and training to reduce the risk of electrical shock

4. APPARATUS
   Loop Detector Sensitivity Tester.

5. PROCEDURE
   Perform the following tests per manufacturer’s instructions.
   a. Loop Amplifier Tests:
   b. Maximum Sensitivity Check
   c. Sustained Presence Check
   d. Sustained Presence Recovery Check
   e. Pulse Check
   f. Second Vehicle Check
   g. Delay Check
   h. Extension Check

6. REPORT
   Record any response found to be in disagreement with the published standards. Report pass or fail and any corrective actions taken on the test report.
Performance Exam Checklist
WSDOT Test Method T 427
Test Method for Loop Amplifier Testing Procedure

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Maximum Sensitivity Check</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2) Sustained Presence Check</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3) Sustained Presence Recovery Check</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4) Pulse Check</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5) Second Vehicle Check</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6) Delay Check</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7) Extension Check</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐  Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:
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WSDOT Test Method T 428
Test Method for Traffic Controller Compliance Inspection and Test Procedure

1. SCOPE
The purpose of this procedure is to provide a documented method for the steps involved with inspection and testing of the completed traffic controller cabinets.

2. REFERENCE DOCUMENTS
WSDOT Test Method T 422, Transient Voltage Test Procedure (optional)
WSDOT Test Method T 423, Conflict Monitor Testing
WSDOT Test Method T 424, Power Interruption Test Procedure
WSDOT Test Method T 425, Environmental Chamber Test
WSDOT Test Method T 427, Loop Amplifier Test Procedure

3. SAFETY
Use proper equipment to reduce the risk of electrical shock.

4. APPARATUS
Equipment as defined to perform WSDOT Test Methods T 422, T 423, T 424, T 425, and T 427.
Resistor load bank to simulate each traffic signal light (150 Ohm 100W wire wound resistors).

5. PROCEDURE
a. The traffic controller cabinet shall be inspected to ensure that it is in compliance with the contract documents. Ensure that all of the required equipment is installed and the cabinet meets the requirements of the contract documents. Any deficiencies shall be documented on the test report.

b. Perform the following tests:
   WSDOT Test Method T 422, Transient Voltage Test (Spike Test) Procedure
   WSDOT Test Method T 423, Conflict Monitor Testing
   WSDOT Test Method T 424, Power Interruption Test Procedure
   WSDOT Test Method T 427, Loop Amplifier Test Procedure

c. After performing the Environmental Chamber Test, at a minimum, verify the operation of the following functions. The NEMA or 170 that are in the ( ) are for those type cabinets only. In addition verify the controller assembly will function as required for the intended intersection.
   1) Verify function of test button on the GFI is operational
   2) Verify vent fan functional
   3) Verify operation of cabinet light door switches
   4) Verify the correct operation of controller and of master if supplied
   5) Use computer to verify the DB9 to C20 plug communication (170)
   6) Verify communication using modem if supplied
   7) Verify operation of pedestrian call switches
   8) Verify the pedestrian field terminal
   9) Verify loop amplifiers operational
  10) Use dummy loop and test loop amplifier field terminals (NEMA)
11) Verify function of detection panel shorting plug is operational (NEMA)
12) Verify operation of preemption field terminal with detector and strobe
13) Verify railroad preemption as required
14) Verify internal wiring
15) Verify the operation of all switches on the intersection display panel
     and loop detector panel
16) Verify operation of inside “auto/flash switch”
17) Verify operation of “stop-time” switch
18) Running test: Set up cabinet with load resistors connected to output field
     terminals and run a performance test for a period of 72 hours.
19) Verify 2’ extension of ped. Yellow from the CMU edge connector.

6. REPORT

   Record any response found to be in disagreement with the published standards, report pass or fail
   and any corrective actions taken on the test report.
Performance Exam Checklist
Method T 428 Checklist

Test Method for Traffic Controller Inspection and Test Procedure

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Cabinet inspected for damage during shipping.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Letter to project office sent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Traffic controller assessed for compliance with contract provisions.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Simulated load connected to the Controller.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Perform Transient Voltage Test WSDOT Method T 422 (optional)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Perform Conflict Monitor Test WSDOT Method T 423</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Perform Power Interruption Test WSDOT Method T 424</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Verify Traffic controller function (Section g of T 421)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐
Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

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WSDOT SOP 429

Methods for Determining the Acceptance of Traffic Signal Controller Assembly

1. SCOPE:
   The purpose of this procedure is to provide a description of the steps involved with traffic signal controller assembly testing.

2. REFERENCE DOCUMENTS
   WSDOT Test Method T421 Receiving Inspection and Test Procedure
   WSDOT Test Method T422 Transient Voltage (Spike Test) Procedure
   WSDOT Test Method T423 Conflict Monitor Testing
   WSDOT Test Method T424 Power interruption Test
   WSDOT Test Method T425 Environmental Chamber Test
   WSDOT Test Method T427 Loop Amplifier Test
   WSDOT Test Method T428 Compliance Inspection and Test Procedure

3. PROCESS:
   WSDOT Test Method T421 Receiving Inspection and Test Procedure
   When the traffic controller assembly arrives for testing, the supplier should have arranged an appointment. Within 5 days of arrival the supplier shall assemble and demonstrate the controller assembly, Test Method T421 is to be completed in the presence of the supplier. After acceptance for testing a letter or e-mail is to be sent to the Project Engineer and/or the local agency identifying the assembly as ready of testing.

   WSDOT Test Method T425 Environmental Chamber Test
   After completion of the environmental chamber test the controller assembly is to be sent to the Region Signal Shop to complete the test regimen. The Region Signal Shop will be informed when a traffic signal controller assembly has passed T425. The Region can choose to have the assembly picked up or it can be shipped commercially. If the controller assembly is not to be sent to the Region Signal Shop the Materials Lab will complete the test regimen beginning with Method T428.

   WSDOT Test Method T428 Compliance Inspection and Test Procedure
   After receiving the controller assembly for testing a letter or e-mail is to be sent to the Project Engineer and/or the local agency identifying the assembly as ready for testing. At this point order is no longer important.

   WSDOT Test Method T422 Transient Voltage (Spike Test) Procedure
   Test Method T422 is to be done only randomly and is not to be done on every assembly.

   WSDOT Test Method T423 Conflict Monitor Testing
   WSDOT Test Method T424 Power interruption Test
   Test Method T424 is not to be performed on type 2070 controllers

   WSDOT Test Method T427 Loop Amplifier Test
Upon completion of all testing send the satisfactory test report to the Project Engineer with a copy to the Region Administrator, the State Signal Operation Engineer and the State Material Laboratory Electrical Engineer.

Note: All of the testing may not be performed at the same facility; there may be more than one satisfactory test report to document all of the required tests.
WSDOT Test Method T 601

Method of Test For Sieve Analysis of Soils — Coarse Sieving

1. SCOPE
   a. This test method describes the procedure for quantitative determinations of the distribution of particles larger than the 4.75 mm (U.S. No. 4) sieve.

2. EQUIPMENT
   a. Drying oven to maintain a temperature of approximately 50°C (122°F).
   b. Drying pans.
   c. A sieve shaker with sieves of the following sizes which conform to the Specifications for Sieves for Testing Purposes (Appendix A), 1 in. (25.4 mm), 3/4 in. (19.0 mm), 1/2 in. (12.5 mm), 1/8 in. (9.5 mm), and U.S. No. 4 (4.75 mm).
   d. A platform scale which has a capacity of 100 lbs. (45 kg) and graduations of 0.1 lb. (0.1 kg).
   e. A metal scoop for removing the sample fraction from the sieves. A heavy-duty household dustpan is satisfactory.
   f. Metal tub or pan for retaining the fine fraction of the screened sample.
   g. Paper sacks, size No. 6.
   h. 5 gal. (20 liter).
   i. Riffle splitter.
   j. Box sieves are to be used for determining the size of particles which are larger than 1 1/2 in. (37.5 mm).
   k. Rotary screen.

3. PREPARATIONS
   a. Empty the sample, unless thoroughly dry, into a large drying pan, spread evenly, and break up large lumps. With some materials, especially those containing clay, it will be necessary to break up aggregations of particles. This may be done by using rodding-type blows with a hammer or by using the rotary screen as a maller (see note).

   Note: Some materials are resistant to shock and abrasion, but are sensitive to moisture. For samples of this nature, a representative portion is soaked in water to observe possible slaking or decomposition characteristics. If slaking or decomposition occurs place the entire sample in a large, rectangular, shallow pan, completely soak with water until decomposition ceases, then thoroughly dry the sample for sieve analysis.

   b. Place a sample identification tag marked with the laboratory and storage container number with the sample.
c. Dry the sample in the oven for approximately 24 hours. Although most materials will dry sufficiently in 24 hours, some clays require a longer time.

   (1) An alternate method of drying a large sample is by the “Air-drying Method” below. Use the following procedure:
   
   Spread the sample, in a uniform layer, on a clean, dry, flat surface (e.g., concrete floor), and allow it to air-dry.

4. PROCEDURE

   a. Insert the sieves in the sieve shaker and secure them.
   b. Put the sample in the top sieve and start the machine.
   c. When the sample contains particles larger than 1 1/2 in. (38 mm), place these particles in the box sieves to determine their maximum size. Particles larger than 2 1/2 in. (63.5 mm) must be measured. Weigh the particles and record the weight (mass) on the test record form.
   d. When the sieving is complete, remove the particles retained on each sieve and weigh to the nearest 0.1 lb. (0.056 g). Record the weight (mass) retained on each sieve.
   e. Any sample that contains individual fractured particles must be checked to determine the percent of fracture by WAQTC FOP for TM 1. Record the 1 percent of fracture.
   f. Unless otherwise instructed under “Remarks,” on DOT Form 351-007, the portions of the sample retained on the 1 1/2 in. (38 mm), 1 in. (25 mm), and 3/4 in. (19 mm) sieves are discarded in the waste bin. The 3/4 in. (19 mm) to 3/8 in. (9.5 mm) and 3/8 in. (9.5 mm) to U.S. No. 4 (4.75 mm) fractions are to be labeled and placed in separate paper sacks.
   g. Weigh and record the weight (mass) of the portion passing the U.S. No. 4 (4.75 mm) sieve.
   h. Separate the fines using the riffle splitter until a 3 to 4 lb. (1.4 to 1.8 kg) representative sample is obtained. This portion is placed in a paper sack, identified by laboratory number, and forwarded to the Fine Grading Laboratory for grain-size analysis, AASHTO T 38.
   i. Place the remainder of the fines U.S. No. 4 (4.75 mm) passing sieve in numbered storage container along with the sacks containing the U.S. No. 4 (4.75 mm) plus fractions (from g. above).
   j. Store the storage container pending further soil tests.

5. PRECAUTIONS

   a. Do not wear loose clothing or apron that will catch on the rotary screen.

6. REPORTS

   a. Test results will be reported on a computer generated test report.
WSDOT Test Method T 606

Method of Test for Compaction Control of Granular Materials

1. SCOPE
   a. This test method consists of three separate tests which present a method for establishing the proper maximum density values to be used for controlling the compaction of granular materials. These methods account for variations of maximum obtainable density of a given material for a given compactive effort, due to fluctuations in gradation.
   b. By splitting the material on the U.S. No. 4 (4.75 mm) sieve and determining the specific gravity, the compacted density, and the loose density of each of the two fractions, a curve of maximum density versus percent passing the U.S. No. 4 (4.75 mm) sieve can be plotted. These curve values will correlate closely with the densities obtained in the field; using modern compaction equipment.

<table>
<thead>
<tr>
<th>Test Method Selection Table</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 % by weight passing #4 sieve</td>
</tr>
<tr>
<td>Soil Type</td>
</tr>
<tr>
<td>Sandy, Non-plastic, permeable</td>
</tr>
<tr>
<td>Silt, some plasticity, low permeability</td>
</tr>
<tr>
<td>Sandy/silt, some plasticity, permeable</td>
</tr>
<tr>
<td>Material has &gt;30% by weight retained on #4 sieve and &lt; 100% by weight retained on #4</td>
</tr>
<tr>
<td>Low permeable soils w/ oversize</td>
</tr>
<tr>
<td>100% by weight retained on #4 sieve</td>
</tr>
<tr>
<td>15% or less of the total sample (coarse material) coarse material exceeds ¾”</td>
</tr>
<tr>
<td>15% or more of the total sample (coarse material) coarse material exceeds ¾”</td>
</tr>
</tbody>
</table>
   c. The test methods are applicable either to specifications requiring compacting to a given percentage of maximum density, or to specifications requiring compaction to a given compaction ratio.
   d. Use of these test methods eliminates the danger of applying the wrong “Standard” to compaction control of gravelly soils.
   e. Native soils within the contract limits to be used for embankment construction and/or backfill material do not require the sampling by a qualified tester. For material that requires gradation testing such as but not limited to manufactured aggregates and Gravel Borrow, a qualified testers shall be required for sampling.
Test No. 1
(Fine Fraction-100 Percent Passing U.S. No. 4 (4.75 mm) Sieve)

1.1 SCOPE
   a. This test was developed for the sandy, non-plastic, highly permeable soils which normally occur as the fine fraction of granular base course and surfacing materials.
   b. When the fine fraction is primarily a soil having some plasticity and low permeability, AASHTO T 99 (Standard Proctor Test) may be used. With borderline soils, both tests should be applied and the one yielding the highest density value should be used.

1.2 EQUIPMENT
   a. Vibratory, Spring Load Compactor (Figures 1 and 2) — Specifications for vibratory spring load compactor can be obtained from the State Materials Lab.
   b. Mold — Molds of the above dimensions can be fabricated from standard cold drawn-seamless piles or tubes. The small button at the center of the small mold follower is a measuring point. The height of this button should be adjusted so the machine follower does not bear on it during compaction. See Figure 3 for mold dimensions.
   c. Mold Piston — A piston which has a diameter of 5 ¾ in. (150 mm).
   d. Height-Measuring Device — A scale with an accuracy of 0.01 in. (0.25 mm).
   e. Tamping Hammer — As specified in AASHTO T 99, Section 2.21.
   f. Sieve — U.S. No. 4 (4.75 mm) sieve.
   g. Oven — Capable of maintaining a temperature of 230° ± 5°F (110 ± 5° C) for drying moisture samples.
   h. Balance — A balance having a capacity of 100 lbs. (45 kg) and an accuracy of 0.1 lbs. (50 g).
   i. Tamping Rod — ¾ in. (16 mm) spherical end.

1.3 PROCEDURE
   a. Oven-dry the total original sample at a temperature not to exceed 140°F (60°C).
   b. Obtain tare weight of mold and bottom plate, record weight (mass) to the nearest 0.01 lb. (5 g).
   c. Separate the sample, by screening, into two fractions divided on the U.S. No. 4 (4.75 mm) sieve. The fraction passing the sieve is used in this test and the fraction retained will be used in Test No. 2.
   d. From the fine fraction (U.S. No. 4 (4.75 mm) minus) split or otherwise obtain a representative sample of approximately 13 lbs. (6 kg). (This mass can be adjusted after the first compaction run to yield a final compacted sample approximately 6 in. (150 mm) high.)
   e. Add water to the sample (the sample should be completely and thoroughly mixed) until it is saturated when compacted. Note that the moisture content should be adjusted so that free water will show at the base of the mold between 500 lbs. (227 kg) and 2,000 lbs. (908 kg). (See Section 1.3h.) Most materials will yield the highest density at that moisture content. Some materials may continue to gain density on increasing the moisture above that specified; however, severe washing-out of the fines will occur, which will alter the character of the sample and void the test results.

Note: “Free water” is classified as a drop or two of water.
f. Place the sample in the mold in three layers. Rod each layer 25 times and tamp with 25 blows of the tamping hammer. The blows of the hammer should produce a 12 in. (305 mm) free fall provided severe displacement of the sample does not occur. In such cases, adjust the blow strength to produce maximum compaction. The surface of the top layer should be finished as level as possible.

![Figure 1](image1)

![Figure 2](image2)

Figure 1

Figure 2


g. Place the piston on top of the sample in the mold, and mount the mold on the jack in the compactor. Elevate mold with the jack until the load-spring retainer seats on top of the piston. Apply initial seating load of about 100 lbs. (45 kg) on the sample.

h. Start the compactor hammers and, at the same time, gradually increase the spring load on the sample to 2,000 lbs. (908 kg) by elevating the jack. The rate of load application is as follows:

<table>
<thead>
<tr>
<th>Load in lbs (kg)</th>
<th>Time in Minutes</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 to 500 lbs.</td>
<td>1</td>
</tr>
<tr>
<td>500 lbs. to 1,000 lbs.</td>
<td>½</td>
</tr>
<tr>
<td>1,000 lbs. to 2,000 lbs.</td>
<td>½</td>
</tr>
</tbody>
</table>

i. After reaching 2,000 lbs. (908 kg), stop the hammers, release the jack, and return to zero pressure.

j. Repeat step h. four additional times. After the last run, remove the mold from the compactor.

k. Measure and record the height of the compacted sample to the nearest 0.01 in. (.25 mm) and calculate the volume (see Section 1.4).

l. Remove the sample from the mold, weigh it, and record its mass (weight) to the nearest 0.01 lbs. (5 g), and calculate the wet density.

m. Vertically slice through the center of the sample, take a representative sample (at least 1.1 lbs. (500 g)) of the materials from one of the cut faces (using the entire sample is acceptable), weigh immediately, and dry in accordance with AASHTO T 255 to determine the moisture content, and record the results. Calculate and record the dry density.

n. Repeat steps d. through m. at higher or lower moisture contents, on fresh samples if needed, to obtain the maximum density value for the material, three tests are usually sufficient.
1.4 CALCULATIONS
a. The formula for calculating the volume and dry and wet densities are as follows:
\[
V = \frac{(H_1 - H_2)B}{1728}
\]
\[
H_1 = \text{Inside height of the mold, mm.}
\]
\[
H_2 = \text{Height from top of the sample to the top of the mold, mm.}
\]
\[
B = \text{Inside bottom area of the mold, mm}^2
\]
\[
\text{Wet Density} = \frac{\text{Wet Mass (Weight)}}{\text{Volume in m}^3 (\text{ft}^3)}
\]
\[
\text{Dry Density} = \frac{\text{Wet Density}}{1 + \text{Moisture Content}}
\]

*Note: See AASHTO T 255-92 “Total Moisture Content of Aggregate by Drying,” for moisture content calculations.

Test No. 2
(Coarse Fraction-100 Percent Retained on the U.S. No. 4 (4.75 mm) Sieve)

2.1 SCOPE
a. This test is used when there is 100 percent retained on the U.S. No. 4 (4.75 mm) sieve. There are two separate procedures based on the maximum size of the aggregate being tested. Procedure 1 is used when no more than 15% of the total sample mass of the coarse aggregate exceeds ¾ in. (19 mm). Procedure 2 is used when 15% or more of the total sample mass of the coarse aggregate is greater than ¾ in. (19 mm), but does not exceed 3 in. (76 mm). If there is any aggregate greater than 3 in. (76 mm), it has to be removed before proceeding with the test.

Procedure 1
(Aggregate Size: No. 4 to ¾ in. (19 mm))

2.2 EQUIPMENT
a. The equipment for this test is the same as that used in Test No. 1

2.3 PROCEDURE
a. From the coarse fraction obtained in Test No. 1, Section 1.3(C), separate a representative sample of 10 to 11 lbs. (4.5 to 5 kg) and weigh to 0.01 lbs. (5 g).

b. Dampen the sample to 2 ½% moisture and place it in a 0.1 ft.\(^3\) (0.0028 m\(^3\)) mold, in three lifts. Tamp each lift lightly to consolidate the material to achieve a level surface. Omit rodding. Avoid loss of the material during placement.

c. Place the piston on top of the simple, in the mold, and follow the procedure described in Test No. 1, Sections 1.3g. through 1.3k.

d. Using the original dry weight value, calculate the dry density in kg/m\(^3\). Use the formula for dry density described in Test No. 1, Section 1.4.

Procedure 2
(Aggregate Size: No. 4 to 3 in. (76 mm))
2.4 EQUIPMENT
   a. ½ ft.$^3$ (0.014 m$^3$) standard aggregate measure.
   b. A metal piston having a diameter ⅛ in. (3 mm) less than the inside diameter of the ½ ft.$^3$
      (0.014 m$^3$) measure.

2.5 PROCEDURE
   a. From the coarse fraction in Test No. 1, Section 1.3c., separate a representative sample of 45 lb.
      (20 kg) and weigh to 0.1 lb. (50 g).
   b. Split the sample into five representative and approximately equal parts.
   c. Place the sample in the mold in five separate lifts after each lift is placed in the mold, position
      the piston on the sample, mount the mold in the compactor, and compact as described in Test
      No. 1, Section 1.3h. Spacers between the load spring and piston must be used to adjust the
      elevation of the mold to the height of the lift being compacted.
   d. After the final lift is compacted, remove the mold from the compactor, determine the height
      of the compacted sample, and calculate the volume (see Test No. 1, Section 1.4(A)).
   e. Calculate the dry density in lbs./ft.$^3$ (kg/m$^3$) (see Test No. 1, Section 1.4(A)).

Test No. 3

Specific Gravity Determination for Maximum Density Test

3.1 EQUIPMENT
   a. Pycnometer calibrated at the test temperature having a capacity of at least 1 quart (100 ml).
   b. One vacuum pump or aspirator (pressure not to exceed 00 mm mercury).
   c. One balance accurate to 0.1 g.

3.2 MATERIAL
   a. Fine fraction U.S. No. 4 (4.75 mm) minus 1.1 lbs. (500 g) minimum.
   b. Coarse fraction U.S. No. 4 (4.75 mm) plus 2.2 lbs. (1,000 g) minimum.

3.3 PROCEDURE
   a. Place dry material, either fine or coarse fraction, in pycnometer, add water. Put pycnometer jar
      top in place and connect to vacuum apparatus. Apply vacuum for at a minimum of 20 minutes
      until air is removed from sample. Slight agitation of the jar every 2 to 5 minutes will aid the
      de-airing process. If the material boils too vigorously, reduce the vacuum. Remove vacuum
      apparatus, fill pycnometer with water, dry outside of jar carefully and weigh. Water
      temperature during test should be maintained as close to 68° ± 1°F (20° ± 0.5°C) as possible.

   Calculate Specific Gravity as follows:
   \[
   \text{Sp. Gr.} = \frac{a}{a + b - c}
   \]

   Where:
   \[
   a = \text{Weight of dry material, grams}
   \]
   \[
   b = \text{Weight of pycnometer + water, grams}
   \]
   \[
   c = \text{Weight of pycnometer + material + water, grams}
   \]
3.4 REPORTS

a. All test results are recorded on the maximum density work sheet. A copy of the maximum density work sheet is shown in Figure 4.

b. The four separate test values determined above are all the data necessary to determine the coordinates required for construction of a maximum density curve, see Figure 5. The end points of the curve are the densities determined from tests number 1 and 2. The four intermediate points are determined through the aid of an appropriate computer program.
Figure 3: Mold Dimensions (All Dimensions ± 0.051 mm (0.002 in.))
MAX. DENSITY CURVE WORKSHEET

Lab. No. __________________________
Material Type: ______________________
Compted by: _________________________
Date: ______________________________

SPECIFIC GRAVITY - COARSE
A. Wt Sample (g) ____________________
B. Wt Pycn + H₂O (g) ________________
C. Wt Pycn w H₂O + A (g) ____________
D. Wt Pycn w H₂O w Sample (g) _______
E. Displacement (ml) ________________

SpG Coarse = \( \frac{A}{E} \)

SPECIFIC GRAVITY - FINE
K. Wt Sample (g) ____________________
L. Wt Pycn + H₂O (g) ________________
M. Wt Pycn w H₂O + K (g) ___________
N. Wt Pycn w H₂O + Sample (g) _______
O. Displacement (ml) ________________

SpG Coarse = \( \frac{K}{O} \)

DENSITY - COARSE
F. Wt Sample (g) ____________________
G. Mold Volume (mm³) ________________
H. Height Constant ___________________
I. Volume Correction _________________
J. Corrected Volume: _________________
   \( G - (H \times I) \)

Density = \( \frac{F}{J} \)

DENSITY - FINE
P. Wt Sample (g) ____________________
Q. Mold Volume (mm³) ________________
R. Height Constant ___________________
S. Volume Correction _________________
T. Corrected Volume: _________________
   \( Q - (R \times S) \)

Density = \( \frac{P}{T} \)

LABOR CODES
T43J _______ T633 _______

Figure 4: Maximum Density Worksheet
Figure 5: Maximum Density Curve
Performance Exam Checklist

*Method of Test for Compaction Control of Granular Materials*
*WSDOT Test Method T 606*

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

**FINE FRACTION — 100% PASSING THE US NO. 4 (4.75 MM) SIEVE**

**Sample Preparation**

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Has the sample been oven-dried?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Has the sample been separated on the US No. 4 (4.75 mm) sieve?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Is the sample weight approximately 13 lbs?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

**Procedure**

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Is sample saturated when compacted?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Has sample been placed in three layers, rodded 25 and tamped 25 times, each layer?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Is the hammer blow approximately a 12 inch free fall to prevent severe displacement of the sample?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. The sample is as level as possible?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Has piston been placed on top of the sample?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Has the mold been mounted on the jack in the compactor?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Has the mold been elevated until the load-spring retainer sits on top of the piston?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Has the initial load been set at 100 pounds?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. Is the loading rate applied as specified in the test procedure?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. Has the hammer been stopped, jack released, and pressure returned to zero when 2,000 pounds pressure was reached?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>11. Steps 7 through 10 repeated four additional times?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>12. Is free water present at the base of the mold between 500 and 2,000 pounds pressure?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>13. The mold removed from the compactor?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>14. Has the height of the sample been determined?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>15. Has sample been weighed?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>
### Procedure Element

16. Has sample been removed from mold and a representative portion immediately weighted and the moisture percentage determined?  
17. Moisture content, dry density determined and entered on the testing sheet?  
18. Has maximum density determined by testing fresh samples, as necessary, at different moisture contents and entered on the testing sheets?

### Sample Preparation

#### Aggregate Size: No. 4 to 3/4 in. (19 MM)

1. Has the sample been oven-dried?  
2. Has the sample been separated on the US No. 4 (4.75 mm) sieve?  
3. Does more than 85% of the entire sample pass the 3/4 in. (19 mm) sieve?  
4. Is the sample weight approximately 10 to 11 lbs?

#### Procedure

1. Weight and record sample weight?  
2. Has the sample been dampened to 2 1/2 percent and placed in a 0.1 ft³ mold and placed in three lifts?  
3. The sample is tamped lightly to archive a level surface?  
4. Piston placed on top of sample and mold mounted on jack in compactor?  
5. Mold elevated until the load-spring retainer sits on top of the piston?  
6. Initial load of 100 lbs set prior to starting machine?  
7. Is the load rate applied as specified in the test procedure?  
8. Hammers stopped, jack released, and pressure returned to 100 pounds when 2000 pound load has been reached?  
9. Steps 5 and 6 repeated four additional times?  
10. The mold removed from the compactor and the height measured?  
11. Dry density calculated and entered on the testing sheets?

#### Aggregate Size: No. 4 to 3 in.

#### Sample Preparation

1. Has the sample been oven-dried?  
2. Has the sample been separated on the US No. 4 (4.75 mm) sieve?  
3. Is the sample weight approximately 45 lbs?  
4. Does the entire sample contain 15% or more 3/4 + material?  
5. Has material greater than 3 in. (76 mm) been removed?  
6. Sample separated into 5 approximately equal parts?
**Procedure Element**

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Sample place in the mold in five separate lifts?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. The sample is as level as possible?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. After each lift, mold placed in compactor and compacted according to test procedure?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. After compacting final lift, sample removed from compactor and volume determined?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Dry density determined calculated and entered onto testing sheet?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

**SPECIFIC GRAVITY DETERMINATION FOR MAXIMUM DENSITY TEST**

**Sample Preparation**

1. Has the sample been oven-dried? | ☐ | ☐ |
2. Has the sample been separated on the US No. 4 (4.75 mm) sieve? | ☐ | ☐ |
3. Weight of fine fraction approximately 500g? | ☐ | ☐ |
4. Weight of coarse fraction approximately 1000g? | ☐ | ☐ |

**Procedure**

1. Mass of each fraction determined? | ☐ | ☐ |
2. Material placed in pycnometer and water at 68°F added? | ☐ | ☐ |
3. Vacuum applied for at least 20 minutes? | ☐ | ☐ |
4. Container and contents agitated manually by shaking at intervals of about 2 to 5 minutes? | ☐ | ☐ |
5. Pycnometer filled with water at 68°F? | ☐ | ☐ |
6. Pycnometer dried, weighted, and recorded on testing sheet? | ☐ | ☐ |
7. Specific Gravity calculated and entered onto testing sheet? | ☐ | ☐ |

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

**Comments:**

________________________________________________________________
________________________________________________________________
________________________________________________________________
WSDOT Test Method T 610

Method of Test for the Capillary Rise of Soil

1. Scope

   a. This test method describes the capillary rise test which determines the susceptibility of soil to damage by frost. The soil sample, closing the top of a glass tube, is subjected to the pull of an increasing head of water which compresses the soil and reduces its moisture content until air is forced through the soil by a maximum head which is termed the capillary rise.

2. Equipment

   a. Capillary Apparatus — The capillary apparatus shall conform to the details shown in Figure 1 and shall consist of the following components:

      (1) Glass Filter Tube — A glass filter tube, as detailed in Figure 1, fitted with a cork disk shaped to fit the shoulder of the tube and having a 1.57 in. (40 mm) diameter hole in the center. A disk of No. 200 (0.075 mm) sieve is placed above the cork disk to retain the soil particles.

      (2) Glass Tube — A glass extension tube, 3.3 ft. (1000 mm) in length, and the same diameter as the lower part of the filter tube, connected to the filter tube by means of a short piece of rubber tubing.

      (3) Glass Cylinder — A glass cylinder, 2 in. (50 mm) in diameter and 4 ft. (1220 mm) long, fitted with a single hole rubber stopper with a short piece of glass tubing, and a rubber hose about 3 ft. (1 m) long with a clamp or other device for controlling the inlet-outlet flow of water.

Figure 1: Capillary Apparatus
3. PROCEDURE
   a. Assemble the equipment as shown in Figure 1.
   b. Select from the material passing the No. 10 (2 mm) sieve a 200-g sample. Select from this
      200-g sample a portion large enough to fill the filler tube, without tamping, to a height
      of 1.57 in. (40 mm).
   c. Admit water into the jacket through the bottom tube until it is filled to a level slightly above
      the top of the soil in the filter tube and allow to stand for five minutes.
   d. After five minutes, lower the water until it is level with the bottom of the cork disk. Allow
      the excess water to drain from the soil
   e. After the excess water has drained from the soil, allow the water level in the jacket to drop
      slowly 2 in. (50 mm) every five minutes until the water in the filter tube breaks.

4. CALCULATIONS
   a. The distance, in inches, between the top of the water in the jacket and the top of the soil when
      the water column in the tube breaks is reported as the capillary rise.

5. REPORTS
   a. All test results will be reported to the Soils Engineer.
WSDOT Test Method T 611

Method of Test for Determination of the Resistance (R-Value) of Untreated Bases, Subbases, and Basement Soils by the Stabilometer

1. SCOPE
   a. This method is a modification of AASHTO Test Method T 190 and covers the procedure for determining the Resistance (R-value) of untreated soils or aggregates for use as base, subbase, or basement soil. This test is divided into the following sections:
      I. Method of Preparation of Materials,
      II. Method of Compaction,
      III. Method of Determination of Exudation Pressure of R-Value Test Specimens,
      IV. Method of Determining the Expansion of Swell Pressures of the R-Value Test Specimens, and
      V. Method of Determining the Stabilometer Resistance R-Value by Means of the HVEEM Stabilometer.

Section I
Method of Preparation of Materials

1. SCOPE
   a. This part of the test procedure describes the methods of weighing and batching materials, the tempering of the specimens to reduce excessive swell pressures and the mixing preparatory to compacting.

2. APPARATUS
   a. Balance, 11 lbs. (5,000 g) capacity.
   b. Water spray metering device calibrated in cubic centimeters.
   c. Mixing pans and spoons.
   d. Airtight plastic containers.
   e. Identification tags.
   f. Drying pans.
   g. Drying oven.

3. PROCEDURE
   a. If the soil sample has an “as received” grading of 95 percent or more passing the U.S. No. 4 (4.75 mm) sieve, six specimens of 2.2 lbs. (1000 g) each of the U.S. No. 4 (4.75 mm) minus material are used.
   b. Should the samples have an “as received” grading with less than 95 percent passing the U.S. No. 4 (4.75 mm) sieve, Appendix C is referred to and the six specimens made up accordingly. These specimens will weigh 2.4 lbs. (1100 g), the percentage and weights being taken from the “as used” column of the Soil Sample Test Data form.
   c. After each specimen is weighed, a tag with the laboratory number is placed in the container with the material.
4. EXCEPTIONS
   a. Some samples such as clays and clay silts, will require tempering. This will be left to the judgment of the operator.
   b. If tempering is required, add 1/2 to 2/3 of the total water required for compaction, place the specimens in airtight plastic bags and allow to stand overnight. Equal amounts of water should be added to each sample.
   c. The amount of water added should be carefully noted for each specimen. This is necessary for accurate results.

Section II
Method of Compaction

1. SCOPE
   a. This method covers procedures for the compaction of soil samples for R-value determinations and related tests.

2. APPARATUS
   a. Mechanical kneading compactor.
   b. Compactor Accessories — 4 in. (102 mm) in diameter × 5 in. (127 mm) high stainless steel molds, mold holder, mold funnel, spatula, rubber discs.
   c. Basket Fabrication Equipment — Paper strips and phosphor bronze perforated discs for basket bottoms. The perforations of the bronze discs are of the same size and spacing as the disc used in determining the exudation pressures and serve this purpose when used with aggregate or granular soil samples.

3. PROCEDURE
   a. Thoroughly mix the specimen with water. Place the steel mold into the mold holder.
   b. Set the air pressure in the compactor to give a tamping foot pressure of 689 kPa (100 psi).
   c. Place approximately 1/2 of the soil into the mold and tamp several times with the metal spoon.
   d. Continue adding soil until enough has been added to produce a 2.52 in. ± 0.1 in. (64 mm ± 2.5 mm) briquette and tamp 40 times with the spoon.
   e. Place the rubber disc on top of the specimen and place mold with mold holder in the compactor.
   f. Star compactor.
   g. For aggregate samples, after 15 blows, increase the air pressure to give a 250 psi (1724 kPa) tamping foot pressure. For all other samples the tamping foot pressure will remain at 100 psi (689 kPa). After 40 blows, raise the compactor foot and remove the mold from the holder. Remove the rubber disc from the mold and smooth the top of the specimen if necessary.

Note 1: The water added to the soil specimen prior to compaction requires judgment and experience on the part of the operator. The amount of water will vary widely from one soil to another but is the amount required to produce a compacted soil briquette that will exude water under a static load ranging between 100 psi (689 kPa) and 600 psi (4137 kPa), ideally bracketing 300 psi (2069 kPa).

Note 2: Always handle mold in an upright position to prevent disturbing the specimen.
4. EXCEPTIONS
   a. Some granular materials are very difficult to handle without damage and may require a paper basket to keep the specimen intact. The baskets prevent the specimen from falling out of the mold and crumbling when the specimen is transferred from the mold to the stabilometer. Compact all such material in baskets. The decision whether to use baskets must be based on experience. They should not be used if not needed. Baskets should be used for all aggregate samples. When using baskets always keep the mold upright.
   
b. If some materials are too soft to compact at the 100 psi (689 kPa), the pressure should be reduced. Use the greatest compaction pressure possible but do not allow the foot to penetrate over $1/4$ in. (6 mm) into the surface of the specimen after all the material is in the mold. Record the new pressure and number of blows at this pressure.
   
c. Soil samples will not be compacted at more than 100 psi (689 kPa) unless it is:
      (1) A borrow material to be used as gravel base.
      (2) From the top 12 in. (305 mm) of in-place material on a resurfacing job.
      (3) Definitely a surfacing material.
      (4) A special research material.

Section III

Method of Determination of Exudation Pressure of R-Value Test Specimens

1. SCOPE
   a. This section covers procedures for determining exudation pressures of specimens of the same sample with varying moisture contents prepared and compacted according to Sections I and II of this test method.

   Note 1: To determine the design R-value or the R-value at an exudation pressure of 300 psi (2069 kPa), four specimens of a sample are compacted at varying moisture contents and constant compactive effort. Ideally, the exudation pressure should bracket 300 psi (2069 kPa). Pressures should be no less than 100 psi (689 kPa) or no greater than 600 psi (4137 kPa).

2. APPARATUS
   b. Perforated phosphor bronze disc, $3\frac{15}{16}$ in. (100 mm) in diameter.
   c. 12,000 lb. (5443 kg) capacity testing machine.
   d. Follower ram $3\frac{1}{32}$ in. (101 mm) outside diameter $\times$ 5 in. (152 mm) high.
   e. 4. in. (102 mm) diameter filter papers.

3. PROCEDURE
   a. Place filter paper and bronze disc on the clear plastic top of the moisture exudation device. Do not invert the mold but place it directly on the saturation indicator disc. Use the lever arm extruder to lower the specimen to the bottom of the mold. Insert the follower and bring the top platen of the testing machine into contact with the follower.
   
b. Use the testing machine to apply a load at the rate of 2,000 lbs. (907 kg) per minute until the filter paper shows water under $5/6$ of the bronze disc. The total load at this point is considered the point of exudation and is recorded in kPa. (Reading on outside scale of testing machine dial.)
**Note 2:** Discard the specimen if the exudation pressure is found to be less than 100 psi (689 kPa) or more than 600 psi (4137 kPa).

4. **EXCEPTIONS**

   a. Occasionally, material from very plastic clay test specimens will extrude under the mold; when this extrusion occurs, call this the point of exudation. There are many cases where high quality materials of a gravely, sandy nature will have exudation pressures that are extremely sensitive to slight changes in moisture content.

   ![Specimen in Mold on Exudation Device in Place in Testing Machine](image)

   **Figure 1: Specimen in Mold on Exudation Device in Place in Testing Machine**

   b. A compacted aggregate specimen, due to large particle sizes, will generally have excessive voids and not produce a good moisture pattern on the exudation filter paper. To these and coarse, granular soil samples, a pressure of 400 psi (2758 kPa) is applied to each specimen. An average of the R-value is then used as the design R-value. Should the specimen show an exudation pattern before 400 psi (2758 kPa) is reached the pressure at this point is recorded. As an example, if the exudation pattern is obtained at 200 psi (1379 kPa), it is recorded as 200/400 (1379/2758).
Section IV
Method of Determining the Expansion or Swell Pressures of the R-Value Test Specimens

1. SCOPE

2. APPARATUS
   a. Swell pressure device and small pans.
   b. 0.0001 in. (.0025 mm) deflection gauge (expansion pressure dial).
   c. Perforated brass discs with stems.
   e. 4.3 in. (11 cm) diameter filter papers to insert between the specimen in the mold and the swell device turntable.

3. PROCEDURE
   a. Insert the perforated brass disc with the stem on top of compacted specimen in mold and seat firmly with finger-pressure. Place the mold in swell pressure device after first placing a filter paper on the turntable.

   Note 1: Allow the specimen to sit for at least one-half hour after completion of exudation test before proceeding with the following steps. This is to allow the specimen to rebound from the exudation pressures applied which otherwise might be recorded falsely as swell pressure.
b. Place the deflection gauge in position on top bar of swell pressure device frame, being very careful to center it.

c. Turn the table of swell pressure device until the brass stem just contacts the calibrated bar. Then turn the table again to apply a surcharge pressure of 0.0010 in. (0.025 mm); **immediately** pour 100 ml to 150 ml of water on top of the specimen.

*Note 2:* Care must be taken that excessive force is not applied when turning up the table of the swell pressure device with a specimen in place. Excessive pressure may cause a temporary set in the calibrated steel bar of the swell pressure device which slowly relieves itself during the soaking period, resulting in erroneous deflection readings.

d. Read and record the height of specimen for density determination.

e. Allow specimen in mold to stand, undisturbed, for 16 to 20 hours.

f. Permeability

   (1) If water is still existing on the specimen after the 16 or 20 hours soaking period but slowly dripping in the pan, it is considered to be very slow draining.

   (2) Should the water drain through the specimen in less than 16 to 20 hours, the drainage is considered slow. If possible, note the natural drainage time and record.

   (3) If the water drains in one to two hours, the specimen is considered fast draining.

   (4) If the water drains in one hour or less, the specimen is considered free draining.

   (5) Nondraining specimens are noted and recorded as such.

g. Read and record the dial deflection at end of soaking period. If water has drained through the specimen into the water pan, pour it back on top of the specimen and allow this water to begin to percolate through the specimen before reading dial deflection and pouring off excess water.

*Note 3:* Do not, under any circumstances, leave a test specimen unconfined by the swell pressure device with a layer of free water existing on the specimen in the mold. This is particularly serious with expansive clays and silts since the condition permits the specimen to expand freely, taking up excess water and disrupting the density. The result will be the unjustifiable reduction of the stabilometer R-value. For the reasons above, as soon as a swell pressure dial reading is taken, the mold and specimen should be removed from the device and drained.

**Section V**

**Method of Determining the Stabilometer Resistance (R-Value) by Means of the Hveem Stabilometer**

1. **SCOPE**
   a. This method covers the procedure for determining the resistance (R-Value) of untreated soils or aggregates for use as base, subbase, or basement soil.

2. **APPARATUS**
   a. HVEEM stabilometer and accessories.
   b. 4 in. (102 mm) inside diameter standard metal calibration mold, 6.50 in. (165 mm) long.
   c. Testing machine, minimum 10,000 lbs. (4540 kg) capacity.
3. CONTROL
   
   a. The correct volumetric adjustment of the air cell in the hydraulic chamber of the stabilometer is necessary in order to establish standardized horizontal pressure and displacement readings. The following is the calibration procedure.

   (1) Adjust the bronze nut on the base of the stabilometer so that an effective height of 2.40 in. (61 mm) of the test specimen is obtained when the stabilometer is in position on the base. The effective height is defined as that depth of the specimen which acts against the liquid phase of the stabilometer. The “ideal” specimen is 2.5 in. (64 mm) high and has an effective height of 2.40 in. (61 mm).

   (2) Place the standard metal calibration mold in the stabilometer. Apply a confining load of 100 lbs. (45 kg) with the testing machine and turn pump to a pressure of exactly 5 psi (34.5 kPa). Adjust the turns indicator dial to zero. Turn the pump handle at the rate of approximately two turns per second until the stabilometer pressure dial reads 100 psi (689 kPa). The turns indicator dial shall read 2.00 + 0.05 turns. If it does not, the air cell must be adjusted. Remove or add air by means of the valve and pump handle, and repeat the displacement measurement after each air change until the proper number of turns is obtained.

4. PROCEDURE
   
   a. After reading and recording the swell pressure deflection and noting drainage characteristics, remove one of the four specimens on the sample from the swell pressure device. Hold the perforated brass disc with stem on the specimen and pour any remaining water from the sample. Remove the disc. Drain for approximately 5 minutes. If the specimen is firm, it may be placed on its side. Granular samples must always be left upright.

   b. Force the specimen which has been previously tested for swell pressure into the stabilometer. Place the follower on top of the specimen and center the stabilometer under the top platen of the testing machine. Raise the testing machine base until 100 lbs. (45 kg) has been applied to the specimen. Release the load to 25 lbs. (11 kg). This ensures seating of specimen in the machine. Adjust stabilometer pump to give a horizontal pressure of exactly 5 psi (34.5 kPa). Begin application of a vertical load at a rate of 25 lbs. (11 kg) per second.

   c. Record the stabilometer pressure gauge readings when the testing machine vertical load pressures are 500 lbs. (227 kg), 1,000 lbs. (454 kg), and 2,000 lbs. (907 kg).

   d. Vertical loading must cease at 2,000 lbs. (907 kg) and the load immediately be reduced to 1,000 lbs. (454 kg). Turn the stabilometer pump so that the horizontal pressure is reduced to exactly 5 psi (34.5 kPa). This will result in a further reduction in the applied testing machine load which is normal and should be ignored.

   **Note 1:** Do not allow the vertical load to exceed 2,000 lbs. (907 kg). On a specimen of a soft, fluid material excess loading can result in damage to the stabilometer pressure gauge.
Figure 3: HVEEM Stabilometer

Set the turn indicator dial to zero. Turn the pump handle at approximately two turns per second until the stabilometer pressure gauge reads 100 psi (14.5 kPa). During this operation, the applied testing machine load will increase and sometimes exceed the initial 1,000 lbs. (454 kg) load. These changes in loadings are normal and should be ignored.

e. Record the number of turns indicated as turns displacement, D, of the specimen. The indicator dial reads in 0.001 in. (0.025 mm) and each 0.1 in. (2.5 mm) is equal to one turn.

f. Remove the specimen from the stabilometer and discard, unless it is needed as a moisture sample.

g. Compute the R-value by the following equation:
   \[ R-value = 1 - \frac{100}{\frac{2.5}{D} \left( \frac{P_v}{P_h} - 1 \right) + 1} \]
   where:
   \( P_v = 160 \) psi (1103 kPa)
   \( D = \) Displacement (no. of turns)
   \( P_h = \) Horizontal pressure at 2,000 lb. (907 kg) vertical load

h. Report steps a. through g. for the remaining three sample specimens.

i. If the exudation pressure causes change in the R-value, plot R-value versus exudation pressure to determine the 300 psi (2069 kPa) R-value, otherwise report the average R-value.
1. SCOPE

   This procedure covers the procedures for determining the in-place density, moisture content, gradation analysis, oversize correction, and determination of maximum density of compacted soils and untreated surfacing materials using a nuclear density device in the direct transmission mode.

2. REFERENCES

   a. WSDOT FOP for AASHTO T 99 for Method of Test for Moisture-Density Relations of Soils
   b. WSDOT FOP for AASHTO T 180 for Method of Test for Moisture-Density Relations of Soils
   c. WSDOT FOP for AASHTO T 224 for Correction for Coarse Particles in Soil Compaction Test
   d. WSDOT FOP for AASHTO T 255 for Total Moisture Content of Aggregate by Drying
   e. WSDOT FOP for AASHTO T 272 for Family of Curves — One Point Method
   f. WSDOT FOP for AASHTO T 310 for In-Place Densities and Moisture Content of Soils and Soil-Aggregate by Nuclear Methods (Shallow Depth)
   g. WSDOT T 606 Method of Test for Compaction Control of Granular Materials

3. DENSITY STANDARDS

   Having the proper soils and using the appropriate density standard for that soil is the key component to getting good compaction.

   Fine-grained soil is defined as soils that contain a significant amount of cohesion and little or no internal friction, density depends on compactive effort and moisture content.

   Coarse grained soil is defined as having little or no cohesion, compactive effort is the primary concern, and moisture content is not as significant an issue because these soils are free-draining and do not retain water.

   Use the following density standard with the appropriate soils:

   AASHTO T99 method A is used when the soils mixture has some plasticity and low permeability. This is defined as having approximately 30% or less material retained on the U.S. No.4 sieve.

   WSDOT T606 Test 1 is for fine sandy, non-plastic, highly permeable soils where approximately 100 percent passes the U.S. No. 4 sieve.

   Note: Soils with low permeability, test the material with AASHTO T99 method A. Use WSDOT T606 test 2, for material retained on the No. 4 sieve.

   AASHTO T180 is used when the soils mixture has coarse and fine aggregate. This is defined as having approximately 30 percent or less retained on the 3/4-in. sieve.

   WSDOT T606 Test 2 is for coarse, granular, free-draining materials when there is approximately 100 percent retained on the U.S. No. 4 sieve.
4. TEST LOCATION

When selecting a test location, the tester shall visually select a site where the least compactive effort has been applied. Select a test location where the gauge will be at least 6 in. (150 mm) away from any vertical mass. If closer than 24 in. (600 mm) to a vertical mass, such as in a trench, follow gauge manufacturer correction procedures.

Note: When retesting is required due to a failing test; retest within a 10 foot radius of the original station and offset.

5. NUCLEAR DENSITY TEST

Determine the dry density and moisture content of soils and untreated surfacing materials using the nuclear moisture-density gauge in accordance with WSDOT FOP for AASHTO T 310, and record on DOT Form 350-074 “Field Density Test”

6. OVERSIZE DETERMINATION

a. A sample weighing a minimum of 9 lbs. will be taken from beneath the gauge. Care shall be taken to select material that is truly representative of where the moisture density gauge determined the dry density and moisture content.

b. There are two methods for determining the amount of oversized aggregate, Method 1, dries the sample to an SSD condition before sieving, and Method 2, a rapid test that washes the No. 4 (4.75 mm) minus material out of the sample before sieving. Method 2 is only recommended for crushed surfacing materials, materials with high clay content, or other granular materials that are at or near the optimum moisture content for compaction.

Method 1:

1. Dry the sample to SSD conditions, (i.e. dried until no visible surface moisture present, material may still appear damp). Allow the sample to cool sufficiently and record mass to the nearest 0.1 percent of the total mass or better.

2. Shake sample by hand over the No. 4 (4.75 mm) sieve. Limit the quantity of material on the sieve so that all particles have the opportunity to reach the sieve openings a number of times during the sieving operation. The mass retained on the No. 4 (4.75 mm) sieve at the completion of the sieving operation shall not exceed 800 grams, 1.8 pounds, for a 12" sieve, or 340 grams, 0.75 pounds; for a 8" sieve.

3. Remove and weigh the material on the No. 4 (4.75 mm) sieve to the nearest 0.1% of the total mass or better and record.

Method 2:

1. Determine the mass of the sample to the nearest 0.1% of the total mass or better and record.

2. Charge the material in a suitable container with water, agitate the material to suspend the fines, then slowly decant and screen the material over a verified No. 4 (4.75 mm) sieve. Repeat as necessary to remove as much of the No. 4 (4.75 mm) minus material as possible. DO NOT overload the sieve.

3. Place the washed sample retained on the No. 4 (4.75 mm) sieve into a tared container. Blot the material to a SSD condition (i.e. no visible surface moisture present, material may still appear damp) during this step.

4. Weigh the mass of the material on the No. 4 (4.75 mm) sieve to the nearest 0.1% of the total mass or better and record.
c. Calculate the percent retained and the percent passing the No. 4 (4.75 mm) sieve to the nearest percent and record on DOT Form 350-074 by the following formula:

\[
\text{% Retained No. 4 (4.75 mm)} = \frac{\text{Mass Retained on the No. 4 (4.75 mm) sieve}}{\text{Initial Mass}}
\]

7. **% COMPACTION DETERMINATION BASED ON WSDOT FOP AASHTO T 99**

a. This process is applicable to nongranular, silty materials with less than 30 percent retained on the No. 4 (4.75 mm) sieve. WSDOT FOP AASHTO Test Method T-99 and WSDOT FOP for AASHTO T-272 are used to determine the maximum density of the material passing the No. 4 (4.75 mm) sieve. Record the maximum density on DOT Form 350-074 line “Maximum Density”

b. The maximum density from WSDOT FOP AASHTO T-99 and WSDOT FOP for T-272 must be corrected for material larger than the No. 4 (4.75 mm) sieve. To correct for the oversize, use WSDOT FOP for AASHTO T224, and enter this value on DOT Form 350-074 line “Corrected Maximum Density”. When less than 5% is retained on the No. 4 (4.75 mm) sieve, no correction is necessary.

c. Percent Compaction is calculated by the following formula and entered on DOT Form 350-074:

**English:**

\[
\% \text{ Compaction (kg/m}^3\text{)} = \frac{\text{Dry Density lbs./ft.}^3\text{ (kg/m}^3\text{)} \times 100}{\text{Corrected Maximum Density lbs/ft}^3\text{ (kg/m}^3\text{)}}
\]

8. **% COMPACTION DETERMINATION BASED ON WSDOT FOP AASHTO T 180**

a. This process is applicable to nongranular, silty materials with less than 30 percent retained on the No. 4 (4.75 mm) sieve and less than 30 percent retained on the ¾ in (19.0mm) sieve. WSDOT FOP AASHTO T 180 is used to determine the maximum density of the material passing the ¾ in (19.0 mm) sieve. Record the maximum density on DOT Form 350-074 line “Maximum Density”

b. The maximum density from WSDOT FOP AASHTO T-180 must be corrected for material larger than the ¾ in (19.0 mm) sieve. The maximum density from WSDOT FOP AASHTO T-180 must be corrected for material larger than the ¾ in (19.0 mm) sieve. To correct for the oversize, use WSDOT FOP for AASHTO T 224, and enter this value on DOT Form 350-074 line “Corrected Maximum Density”. When 5% or less is retained on the ¾ in (19.0 mm) sieve, no correction is necessary.

c. Percent Compaction is calculated by the following formula and entered on DOT Form 350-074.

\[
\% \text{ Compaction} = \frac{\text{Dry Density lbs./ft.}^3\text{ (kg/m}^3\text{)} \times 100}{\text{Corrected Maximum Density lbs/ft}^3\text{ (kg/m}^3\text{)}}
\]
9. % COMPACTION DETERMINATION BASED ON WSDOT TEST METHOD No. 606.
   a. This process is applicable to granular, free-draining materials and to materials with 30 percent or more retained on the No. 4 (4.75 mm) sieve. Test Method 606 requires specialized equipment and is run only by the Region or State Materials Laboratory.
   b. Using the appropriate computer-generated chart, determine the maximum density, based on the percent passing the No. 4 (4.75 mm) sieve. This value should be entered on DOT Form 350-074 on line “Maximum Density”.
   c. Percent of compaction is then calculated by the formula and entered on DOT Form 350-074:

   \[
   \% \text{ Compaction} = \frac{\text{Dry Density lbs./ft.}^3 (\text{kg/m}^3) \times 100}{\text{Maximum Density lbs/ft}^3 (\text{kg/m}^3)}
   \]

10. OPTIMUM MOISTURE DETERMINATION
   a. The optimum moisture content for WSDOT FOP for AASHTO T 180 will have to be corrected with the following formula:

   \[
   \text{Corrected Optimum Moisture} = (\text{Optimum Moisture}) \times (\% \text{ Passing 3/4 (19.0mm)})
   \]
   b. The optimum moisture content for WSDOT test method T 606 and WSDOT FOP for AASHTO T 99 will have to be corrected with the following formula:

   \[
   \text{Corrected Optimum Moisture} = (\text{Optimum Moisture}) \times (\% \text{ Passing No. 4 (4.75mm)})
   \]
   c. Record the Optimum Moisture content from the appropriate density curve on DOT Form 350-074.

11. REPORT
   Report compaction data of DOT Form 350-074, “Field Density Test” and on DOT Form 351-015 “Daily Compaction Test, or other report approved by the State Materials Engineer.
   Report percent compaction to the nearest whole number.
WSDOT Test Method T 702

Method for Preparation of Test Specimens of Hot Mix Asphalt by Means of California Kneading Compactor

1. SCOPE
   a. This method for preparation of test specimens of Hot Mix Asphalt mixtures is accomplished by means of a mechanical compactor.
   b. This method is the WSDOT equivalent to AASHTO T 247.

2. APPARATUS
   a. California Kneading Compactor — Mechanical kneading compactor, as shown in Figure 1, for consolidating test specimens.

   Figure 1: California Kneading Compactor

   b. Compactor Foot — A ram having a face shaped as shown in Figure 2, and having an area of approximately 3.1 in.² (20.059 cm²).
   c. Funnel — as shown in Figure 3.
   d. Molds — Molding cylinders, 4.000 in. ± 0.005 in. (101.60 ± 0.13 mm) in inside diameter by 5 in. (127 mm) in height. A minimum of six such compaction molds is recommended.
   e. Metal Followers — Two metal followers, 3.985 in. ± 0.005 in. (101.2 ± 0.11 mm) in diameter; one 5.5 in. (140 mm) high, the other 1.5 in. (38.1 mm) high.
f. Testing Machine — A compression testing machine having a minimum capacity of 13,000 lbs. (55 kN).

g. Ovens — Electric ovens capable of maintaining temperatures of 140 ± 5°F (60° ± 3°C) and 275± 5°F (135° ± 3°C).

h. Measuring Device- A device for measuring the height of the sample to the nearest 0.01 in.

i. Miscellaneous Apparatus — Thermometers, trowels, spatulas scoops, gloves, and an assortment of metal pans.

3. TEST SPECIMENS

a. Preparation of Mixture — The Hot Mix Asphalt mixture shall be prepared in accordance with WSDOT Test Method 726.

b. Size of Specimens — Test specimens for use with the HVEEM Stabilometer in accordance with WSDOT Test Method No. 703 shall be 4-in. (102-mm) in diameter by 2.5 ± 0.1 in. (64 ± 3 mm) in height. Approximately 1240 g of material is adequate for a specimen of the proper height.
4. PROCEDURE
   a. Heat the Hot Mix Asphalt and compaction molds to 275°F to 300°F (149°C) maximum. Place the compaction mold in position in the mold holder and secure. Insert paper disk, 4 in. (102 mm) diameter, to cover the base plate of the mold holder. Start compactor and place one half of the Hot Mix Asphalt mixture into the mold using a spoon or other suitable devise to transfer the material and apply 10 tamping blows. Add the remaining Hot Mix Asphalt to the the mold and apply 150 tamping blows to complete compaction in the California Kneading Machine.
   b. Remove the molded specimen from the machine. Place the specimen (still in the mold) on the compression testing machine. By the double plunger method, apply a “leveling off” load of 12,560 lbs. (86.6 MPa) for a period of 1 minute. Remove the specimen from the mold and cool to room temperature. Measure the height of the sample to the nearest .01 inch.
   c. Determine the density of the sampler per AASHTO T 166.

5. REPORT
   a. The report shall include the following: height, mass, and density of the specimens.
WSDOT Test Method T 703

Method of Test for Resistance to Deformation of Hot Mix Asphalt by Means of HVEEM Stabilometer

1. SCOPE
   a. These methods of test are intended for determining (1). The resistance to deformation of compacted Hot Mix Asphalt mixtures by measuring the lateral pressure developed from applying a vertical load by means of the HVEEM stabilometer (2). The cohesion of compacted Hot Mix Asphalt mixtures by measuring the forces required to break or bend the sample as a cantilever beam by means of the Hveem cohesiometer.

2. OUTLINE OF METHOD
   a. Use current AASHTO Test Method T 246 except as follows:
      (1) Replace 3.2 with the following: “Test specimens shall be formed and compacted in accordance with WSDOT Test Method No. 702.”
      (2) Delete the first sentence of 5.2.
WSDOT Test Method T 706

Method of Static Immersion Asphalt in Water Preferential Stripping Test

1. SCOPE
   a. This method is used to determine the extent of static stripping, a preference of aggregate for asphalt binder or water. The results are used to determine the suitability of the aggregate for Hot Mix Asphalt.
   b. This test is technically equivalent to AASHTO T 182.

2. EQUIPMENT
   a. Mixing Bowl — Hemispherical.
   b. Mixing spoon.
   c. Distilled water.
   d. Glass Jar — 400 ml.
   e. Oven — 140° ± 5°F (60° ± 3°C).

3. PROCEDURE
   a. Use the liquid asphalt binder product called for in the contract, or if a preliminary sample, use both MC-800 and RC-3000.
   b. The material to be tested shall be ⅜ in. to ¼ in. (9.5 mm to 6.3 mm) aggregate or other sized material without the ¼ in. to 0 in. (6.3 mm to 0).
   c. Heat aggregate and liquid asphalt binder at 140° ± 5°F (60° ± 3°C) for approximately two hours.
   d. In the hemispherical bowl, mix until completely coated 94 g of ⅜ in. to No. 4 (9.5 mm to 4.75 mm) aggregate with 6 ± 0.5 g immeasurable of liquid asphalt binder.
   e. Place the sample in a 140°F (60°C) curing oven for approximately 15 hours.
   f. Remove the mix from the oven, stir to recoat particles if necessary, and transfer the coated aggregate to the 400 ml glass jar. Add distilled water, warmed to room temperature, to cover the sample.
   g. Examine the sample for stripping at the end of 1, 2, and 24 hours without removing the aggregate from the water. The stripping is estimated by comparison with a nonstripping aggregate.

4. REPORTS
   a. Stripping results are recorded as none, slight, moderate, or severe.
WSDOT Test Method T 712

Standard Method of Reducing Hot Mix Asphalt Paving Mixtures

Significance

Samples of bituminous paving mixes taken in accordance with FOP AASHTO T 168 are composites and are large to increase the likelihood that they are representative of the product being tested. Materials sampled in the field need to be reduced to appropriate sizes for testing. It is extremely important that the procedure used to reduce the field sample not modify the material properties.

1. SCOPE

This method covers the procedure for reducing samples of Hot Mixed Asphalt (HMA). The samples are to be acquired in accordance with FOP AASHTO T 168. The sample is to be representative of the average of the HMA being produced.

2. APPARATUS

- Flat-bottom scoop,
- Broom or brush,
- Non-stick splitting surface such as metal, paper, canvas blanket or heat-resistant plastic,
- Large spatulas, trowels, metal straight edge or 12 in. dry wall taping knife, sheet metal quartering splitter,
- Mechanical Splitter—The splitter shall have four equal width chutes, which will discharge the material into four appropriate size containers. The splitter shall be designed with a receiving hopper that will hold the HMA field sample until a handle releases the material to fall through a divider and is distributed into four equal portions. The splitter shall be designed so that the HMA field sample will flow smoothly and freely through the divider without loss of materials (See Figures 1 to 3.).
• Oven — An oven of appropriate size, capable of maintaining a uniform temperature within the allowable tolerance for the grade of asphalt.

• Miscellaneous equipment including trowel(s), spatula(s), hot plate, non-asbestos heat-resistant gloves or mittens, pans, buckets, cans.

3. SAMPLE PREPARATION

The sample must be warm enough to separate. If not, warm in an oven until it is sufficiently soft to mix and separate easily.

4. PROCEDURE

Initial Reduction of Field Sample

A. Place the sample on a hard, clean, non-stick, level surface where there will be neither loss of material nor the accidental addition of foreign material. The surface may be...
3. **SAMPLE PREPARATION**

   The sample must be warm enough to separate. If not, warm in an oven until it is sufficiently soft to mix and separate easily.

4. **PROCEDURE**

   **Initial Reduction of Field Sample**

   A. Place the sample on a hard, clean, non-stick, level surface where there will be neither loss of material nor the accidental addition of foreign material. The surface may be covered with a canvas blanket, heavy paper or other suitable material. Remove the sample from the agency approved containers by dumping into a conical pile.

   ![Figure 4](image)

   B. Divide the sample into four approximately equal quarters with a spatula, trowel, flat metal plate, sheet metal quartering splitter, or mechanical splitter.

   C. With the quartering device in place remove all the material from each quarter. If needed for additional testing the material should be placed in agency approved containers for storage or shipment.

   **Note 1:** When testing lean mixes or mixes with aggregate larger than $\frac{3}{4}$ in. (19 mm), sampling as described in Method B will be used, with no remixing and no removal of a similar amount of material from the opposite quarter, is recommended at this point to obtain samples for each acceptance test.

   D. Pay particular attention that excessive amounts of material is not left on the splitting surface or splitting equipment.

   F. When the further reduction of the HMA is to be done, proceed according to step 2 of methods A, B, or C.

   **Note 2:** Identify the opposite quarter as the “Challenge Sample.”

   **Reducing to Test Size — Method A**

   1. On a hard, clean, non-stick, level surface where there will be neither loss of material nor the accidental addition of foreign material. Remove the sample from the agency approved containers by dumping into a conical pile. The surface shall be covered with either a canvas blanket, heavy paper or other suitable material.

   2. With the material on the canvas or paper, mix the sample thoroughly by turning the entire sample over the minimum amount of times to achieve a uniform distribution. Alternately lift each corner of the canvas or paper and pull it over the sample diagonally toward the opposite corner causing the material to be rolled. With the last turning, lift both opposite corners to form a conical pile.
3. Grasp the canvas or paper, roll the material into a loaf and flatten the top.

![Figure 5](image)

4. Pull the canvas or paper so approximately \( \frac{1}{4} \) of the length of the loaf is off the edge of the counter. Allow this material to drop into a container to be saved. As an alternate using a straight edge slice off approximately \( \frac{1}{4} \) of the length of the loaf and place in a container to be saved.

![Figure 6](image)

5. Pull additional material (loaf) off the edge of the counter and drop the appropriate size sample into a sample pan or container. As an alternate using a straight edge slice off appropriate size sample from the length of the loaf and place in a sample pan or container.

6. Repeat step 5 until the proper size sample has been acquired. Step 5 is to be repeated until all the samples for testing have been obtained.

**Note 3:** When reducing the sample to test size it is advisable to take several small increments determining the mass each time until the proper minimum size is achieved. Unless the sample size is below the minimum or exceeds the maximum test size use, the sample as reduced for the test.

**Reducing to Test Size — Method B**

1. On a hard, clean, non-stick, level surface where there will be neither loss of material nor the accidental addition of foreign material. Remove the sample from the agency approved containers by dumping into a conical pile. The surface shall be covered with either a canvas blanket, heavy paper or other suitable material. (See Note 1.)

2. With the material on the canvas or paper, mix the sample thoroughly by turning the entire sample over the minimum amount of times to achieve a uniform distribution. Alternately lift each corner of the canvas or paper and pull it over the sample diagonally toward the opposite corner causing the material to be rolled. With the last turning, lift both opposite corners to form a conical pile.

3. Quarter the conical pile using a quartering device or straightedge.
4. With the quartering device in place using a suitable straight edge slice through the quarter of the HMA from the apex of the quarter to the outer edge. Pull or drag the material from the quarter holding one edge of the straight edge in contact with the quartering device. Two straight edges may be used in lieu of the quartering device.

5. Slide or scoop the material into a sample pan. Repeat step 4 removing a similar amount of material from the opposite quarter. Step 4 is to be repeated until all the samples for testing have been obtained.

**Note 4:** When reducing the sample to test size it is advisable to take several small increments determining the mass each time until the proper minimum size is achieved. Unless the sample size is below the minimum or exceeds the maximum test size use, the sample as reduced for the test.

**Reducing to Test Size — Method C**

1. On a hard, clean, non-stick, level surface where there will be neither loss of material nor the accidental addition of foreign material. Remove the sample from the agency approved containers by dumping into a conical pile. The surface shall be covered with either a canvas blanket, heavy paper or other suitable material.

2. With the material on the canvas or paper, mix the sample thoroughly by turning the entire sample over the minimum amount of times to achieve a uniform distribution. Alternately lift each corner of the canvas or paper and pull it over the sample diagonally toward the opposite corner causing the material to be rolled. With the last turning, lift both opposite corners to form a conical pile.

3. Quarter the conical pile using a quartering device or straightedge.

4. Remove the opposite quarters saving the material for future use.

5. Repeat step 2 through 4 until the proper size sample has been achieved.

6. When additional test specimens are required, dump the removed material into a conical pile as in step 1 and repeat steps 2 through 5. This process may be repeated until sample have been reduced to testing size for all tests.

7. **SAMPLE IDENTIFICATION**

   (1) Each sample submitted for testing shall be accompanied by a transmittal letter completed in detail. Include the contract number, acceptance and mix design verification numbers, mix ID.

   (2) Samples shall be submitted in standard sample boxes, secured to prevent contamination and spillage.

   (3) Sample boxes shall have the following information inscribed with indelible-type marker: Contract number, acceptance and mix design verification numbers, mix ID.

   (4) The exact disposition of each quarter of the original field sample shall be determined by the agency.
# Performance Exam Checklist

## Reducing Samples of Hot Mix Asphalt to Testing Size

**WSDOT Test Method T 712**

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<th>Exam Date</th>
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### Procedure Element

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#### Method A

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#### Method C

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First attempt:  

First attempt: Pass [ ] Fail [ ]  

Second attempt:  

Second attempt: Pass [ ] Fail [ ]

Signature of Examiner __________________________________________
WSDOT Test Method T 716

Method of Random Sampling for Locations of Testing and Sampling Sites

1. SCOPE
   a. This method outlines the procedure for selecting sampling and testing sites in accordance with accepted random sampling techniques. It is intended that all testing and sampling locations be selected in an unbiased manner based entirely on chance.
   b. Testing and sampling locations and procedures are as important as testing. For test results or measurements to be meaningful, it is necessary that the sampling locations be selected at random, typically by use of a table of random numbers. Other techniques yielding a system of randomly selected locations are also acceptable.
   c. This procedure is divided into several sections:
      • Applications for Hot Mixture Asphalt Density and Challenge Cores, Section 5
      • Applications for Hot Mixture Asphalt (HMA) Sampling, Section 6
      • Applications for Portland Cement Concrete, Section 7
      • Applications for Aggregate and other materials, Section 8

2. Straight Random Sampling vs. Stratified Random Sampling:
   Straight random sampling considers an entire lot as a single unit and determines each sample location based on the entire lot size. Stratified random sampling divides the lot into a specified number of sublots or units and then determines each sample location within a distinct sublot. Both methods result in random distribution of samples to be tested for compliance with the agency’s specification.

3. PROCEDURE
   a. Determine the lot, or sublot size and number of tests per LOT or sublot.
   b. Determine the “X” and/or “Y” random number by using values from the random number table.
   c. Multiply the lot or sublot size by the random number. This will give you the approximate test location within the lot or sublot to do the testing.

4. Stratified Random Sampling
   a. Following determination of the LOT length in Example 1, determine the length increment for individual sublots by dividing by the number of such desired sublots. In the case of Hot Mix Asphalt Pavement this would be five sublots
   b. Determine random location factors “X” and/or “Y” values by random entry to the table.
   c. To determine the location of test No. 1 in sublot No. 1 multiply the sublot increment by the selected “X” or “Y” factor from the Random Number table, then add this amount to the beginning location. Test locations within each of the subsequent sublots are determined by calculating the fractional location within the sublot interval then adding the increment of the preceding sublot.
   d. For irregular lot or sublot sizes at the end of production, determine the location by dividing the final increment into 5 equal parts and define a test location within each.
5. APPLICATIONS FOR HOT MIX ASPHALT DENSITY AND CHALLENGE CORES (ENGLISH UNITS)

Note: For metric projects refer to Appendix A.

a. Determine the LOT size and number of tests per LOT. The Standard specifications set the size of a density test lot for Hot Mix Asphalt Pavement to no greater than a single day’s production or 400 tons, whichever is less, and require five tests per LOT. At the end of a day’s production the final lot may be increased to a maximum of 600 tons.

b. Convert this LOT size to an area segment of the roadway based on the roadway section and depth being constructed for the course being tested. The calculations in Example 1 show how this is performed. Table 1 has been provided to give you recommend lot lengths for standard lane widths at various depths. Lot length needs to be determined to the nearest 100 feet.

Example 1
Sample Computation for Lot Length

Using nominal compacted density of 2.05 tons/cy, and a 400 ton lot:

Tons per linear foot = \( \frac{(1.0 \text{ (foot)} \times \text{width (feet)} \times \text{depth(feet)}) \times 2.05 \text{ Tons/cy}}{27} \)

\( \text{Tons per linear Foot = } \frac{1.0 \text{ ft} \times 12 \text{ ft} \times 0.15 \text{ ft} \times 2.05 \text{ tons}}{27} = 0.137 \text{ Tons per linear Foot.} \)

\( \text{Lot length = } \frac{400 \text{ Tons}}{0.137 \text{ Tons per linear Foot}} = 2900 \text{ linear Feet} \)

Table 1:

<table>
<thead>
<tr>
<th>Lane Width</th>
<th>Compacted Depth</th>
<th>Computed Lot Length</th>
<th>Recommended Lot Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 feet</td>
<td>0.12</td>
<td>3655</td>
<td>3700</td>
</tr>
<tr>
<td></td>
<td>0.15</td>
<td>2924</td>
<td>2900</td>
</tr>
<tr>
<td></td>
<td>0.20</td>
<td>2193</td>
<td>2200</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>1754</td>
<td>1800</td>
</tr>
<tr>
<td>11 feet</td>
<td>0.12</td>
<td>3987</td>
<td>4000</td>
</tr>
<tr>
<td></td>
<td>0.15</td>
<td>3189</td>
<td>3200</td>
</tr>
<tr>
<td></td>
<td>0.20</td>
<td>2392</td>
<td>2400</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>1913</td>
<td>1900</td>
</tr>
</tbody>
</table>

LOT length may also be determined based on Nominal Designated LOT sizes. To utilize this concept, compacted mix volumes equivalent to the designated mix quantity per LOT have been determined using the nominal compacted unit weight of Hot Mix asphalt. These volumes are then converted into Density LOT lengths using the typical lane width and specified compacted depth. The included tables present the values for LOT Lengths based on English units.

c. Determine the locations of the test (or sampling) sites by using values from the random number table (Table 2) to determine the coordinate location on the roadway. In the table, use the “X” values as decimal fractions of the total length of the lot; use the “Y” values as fractions of the width, customarily measured from the right edge of the pavement. The values in the table have been set so that no measurements are taken within 1.5 LF (0.45 m) of the edge of the pavement. Whenever a test location is determined to fall within such an area (i.e., bridge end, track crossing, or night joint) the test location should be moved ahead or back on stationing, as appropriate, by 25 LF (8 m).
Table 2

Random Numbers with X and Y values

<table>
<thead>
<tr>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
<th>X</th>
<th>Y</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.0290</td>
<td>0.33</td>
<td>2</td>
<td>0.0119</td>
<td>0.43</td>
<td>3</td>
<td>0.954</td>
<td>0.15</td>
<td>4</td>
<td>0.784</td>
<td>0.39</td>
<td>5</td>
<td>0.857</td>
<td>0.29</td>
<td>6</td>
<td>0.069</td>
<td>0.14</td>
</tr>
<tr>
<td>0.21</td>
<td>0.712</td>
<td>0.17</td>
<td>0.22</td>
<td>0.976</td>
<td>0.69</td>
<td>0.23</td>
<td>0.997</td>
<td>0.63</td>
<td>0.24</td>
<td>0.024</td>
<td>0.47</td>
<td>0.25</td>
<td>0.352</td>
<td>0.17</td>
<td>0.26</td>
<td>0.626</td>
<td>0.29</td>
</tr>
<tr>
<td>0.41</td>
<td>0.172</td>
<td>0.32</td>
<td>0.42</td>
<td>0.430</td>
<td>0.37</td>
<td>0.43</td>
<td>0.704</td>
<td>0.19</td>
<td>0.44</td>
<td>0.009</td>
<td>0.18</td>
<td>0.45</td>
<td>0.552</td>
<td>0.17</td>
<td>0.46</td>
<td>0.196</td>
<td>0.15</td>
</tr>
<tr>
<td>0.61</td>
<td>0.768</td>
<td>0.32</td>
<td>0.62</td>
<td>0.983</td>
<td>0.37</td>
<td>0.63</td>
<td>0.504</td>
<td>0.66</td>
<td>0.64</td>
<td>0.043</td>
<td>0.31</td>
<td>0.65</td>
<td>0.284</td>
<td>0.39</td>
<td>0.66</td>
<td>0.196</td>
<td>0.15</td>
</tr>
<tr>
<td>0.81</td>
<td>0.477</td>
<td>0.87</td>
<td>0.82</td>
<td>0.267</td>
<td>0.44</td>
<td>0.83</td>
<td>0.933</td>
<td>0.28</td>
<td>0.84</td>
<td>0.974</td>
<td>0.87</td>
<td>0.85</td>
<td>0.600</td>
<td>0.19</td>
<td>0.86</td>
<td>0.591</td>
<td>0.19</td>
</tr>
<tr>
<td>0.88</td>
<td>0.668</td>
<td>0.41</td>
<td>0.89</td>
<td>0.327</td>
<td>0.29</td>
<td>0.90</td>
<td>0.473</td>
<td>0.51</td>
<td>0.91</td>
<td>0.598</td>
<td>0.58</td>
<td>0.92</td>
<td>0.373</td>
<td>0.69</td>
<td>0.93</td>
<td>0.44</td>
<td>0.24</td>
</tr>
<tr>
<td>0.94</td>
<td>0.831</td>
<td>0.14</td>
<td>0.95</td>
<td>0.178</td>
<td>0.45</td>
<td>0.96</td>
<td>0.821</td>
<td>0.46</td>
<td>0.97</td>
<td>0.124</td>
<td>0.62</td>
<td>0.98</td>
<td>0.580</td>
<td>0.57</td>
<td>0.99</td>
<td>0.037</td>
<td>0.24</td>
</tr>
</tbody>
</table>
d. In order to determine which “X” and “Y” values should be used, enter the table on a line chosen by chance. Recommended procedure is selection of a line based on the last two digits from the most recent standard count on the nuclear density gage. Subsequent “X” and “Y” values are then taken from the lines that follow. Based on the specified sampling frequency, 20 lots can be accommodated by one cycle through the table. Start each shift with a set of values determined by chance in order to obtain random selection.

e. Example 2 shows the calculations for determining the testing location for asphalt pavement density. No Figure 1

Example 2
Test Location Within the LOT
for Hot Mix Asphalt Density

For the lot: (12 ft. wide, 0.15 ft. deep, starting at station 168 + 75 with paving progressing ahead on station), Lot length was previously determined as 2,900 LF. Using the last two digits of the standard count, as in the example, 2951, assume “X” and “Y” values from line (51) in table 2: X = 0.762, Y = 0.65.

For the first test:

Beginning station: 168 + 75
Sublot length increment: 580 * 0.762 = 442
Width offset: 12 * 0.65 = 7.8 ft. (from right edge)
Location is: station: (168+75) + 442 = 173 + 17, 7.8 ft. from right edge

For the Second test:

Beginning station: (168 + 75) + (580) = 174 + 55
Sublot length increment: 580 * 0.285 = 165
Width offset: 12 * 0.28 = 3.4 ft. (from right edge)
Location is: station: (174 + 55) + 165 = (176 + 20), 3.4 ft. from right edge

For the Third test:

Beginning station: (168 + 75) + 580 + 580 = 180 + 35
Sublot length increment: 580 * 0.347 = 201
Width offset: 12 * 0.87 = 10.4 ft. (from right edge)
Location is: station: (180 + 35) + 201 = (182 + 36), 10.4 ft. from right edge

6. APPLICATIONS FOR HOT MIX ASPHALT (HMA) PAVEMENT MIXTURE

a. Determine the sublot size. The Standard Specifications define a lot as the total quantity of material or work produced for each job mix formula (JMF). The sublot size for HMA gradation, binder content, and/or volumetrics is a maximum of 800 tons, and shall be determined to the nearest 100 tons. At the end of production, the final sublot may be increased to a maximum of 2 times the sublot quantity calculated.

Sampling of binder shall be every other mixture sample.
b. Determine the locations of the test (or sampling) sites as defined in Section 3 using random numbers from table 3, or from another Random Number Generator. Do not sample from the first or last 25 tons. Once the two-digit number is selected the corresponding four-digit number becomes the factor for determining the selection of the next sample. Random sample tonnage may be adjusted per subplot to accommodate field testing. Adjustments to random sample tonnage should be documented.

<table>
<thead>
<tr>
<th>X</th>
<th>X</th>
<th>X</th>
<th>X</th>
<th>X</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.186</td>
<td>0.256</td>
<td>0.201</td>
<td>0.508</td>
<td>0.431</td>
</tr>
<tr>
<td>0.584</td>
<td>0.753</td>
<td>0.699</td>
<td>0.884</td>
<td>0.509</td>
</tr>
<tr>
<td>0.965</td>
<td>0.108</td>
<td>0.785</td>
<td>0.648</td>
<td>0.962</td>
</tr>
<tr>
<td>0.044</td>
<td>0.626</td>
<td>0.874</td>
<td>0.398</td>
<td>0.315</td>
</tr>
<tr>
<td>0.840</td>
<td>0.885</td>
<td>0.604</td>
<td>0.142</td>
<td>0.721</td>
</tr>
<tr>
<td>0.381</td>
<td>0.418</td>
<td>0.087</td>
<td>0.962</td>
<td>0.637</td>
</tr>
<tr>
<td>0.756</td>
<td>0.320</td>
<td>0.334</td>
<td>0.516</td>
<td>0.056</td>
</tr>
<tr>
<td>0.586</td>
<td>0.098</td>
<td>0.189</td>
<td>0.615</td>
<td>0.905</td>
</tr>
<tr>
<td>0.480</td>
<td>0.791</td>
<td>0.777</td>
<td>0.226</td>
<td>0.195</td>
</tr>
<tr>
<td>0.101</td>
<td>0.717</td>
<td>0.704</td>
<td>0.881</td>
<td>0.981</td>
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<td>0.385</td>
<td>0.266</td>
<td>0.744</td>
<td>0.433</td>
</tr>
<tr>
<td>0.456</td>
<td>0.465</td>
<td>0.791</td>
<td>0.229</td>
<td>0.762</td>
</tr>
<tr>
<td>0.778</td>
<td>0.101</td>
<td>0.711</td>
<td>0.906</td>
<td>0.678</td>
</tr>
<tr>
<td>0.243</td>
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<td>0.122</td>
<td>0.413</td>
<td>0.347</td>
</tr>
<tr>
<td>0.578</td>
<td>0.829</td>
<td>0.895</td>
<td>0.827</td>
<td>0.274</td>
</tr>
<tr>
<td>0.966</td>
<td>0.998</td>
<td>0.371</td>
<td>0.984</td>
<td>0.114</td>
</tr>
<tr>
<td>0.373</td>
<td>0.539</td>
<td>0.221</td>
<td>0.641</td>
<td>0.480</td>
</tr>
<tr>
<td>0.834</td>
<td>0.060</td>
<td>0.011</td>
<td>0.068</td>
<td>0.685</td>
</tr>
</tbody>
</table>

c. In order to determine which random values should be used, enter the table on a line chosen by chance. Recommended procedure is selection of a line based on the last two digits of the ignition furnace calibration.

d. Example 3 shows the calculations for determining the testing location for HMA WSDOT Form DOT 350-160 will calculate the testing location for you.
Example 3
Test Location for a Sublot of HMA

The Ignition Furnace calibration is 0.45%. Use 45 as the starting point to enter the random number table 3. The starting random number is 0.604.

**For the First test point:**
- Beginning tonnage: 0
- Sublot increment: $800 \times 0.604 = 483$
- Testing tonnage is at: 483 tons

**For the Second test point:**
- Beginning tonnage: 800
- Sublot increment: $800 \times 0.087 = 70$
- Testing tonnage is at: $800 + 70 = 870$ tons

**For the Third test point:**
- Beginning Tonnage: $800 + 800 = 1600$
- Sublot increment: $800 \times 0.334 = 267$
- Testing tonnage is at: $1600 + 267 = 1867$ tons

**For the Fourth test point:**
- Beginning Tonnage: $1600 + 800 = 2400$
- Sublot increment: $800 \times 0.189 = 151$
- Testing tonnage is at: $2400 + 151 = 2551$ tons

7. APPLICATIONS FOR PORTLAND CEMENT CONCRETE

a. Determine the sublot size. The Standard Specifications states after two successive tests indicate that the concrete is within specified limits; the sampling and testing frequency may decrease to one for every five truck load. Concrete samples other than initial load samples or samples for questioned acceptance will be taken from each sublot by a random selection. Random selection will be accomplished by using the random number table 3. For each day of concrete delivery and placement a new random number will be selected and the process repeated.

b. Determine the locations of the test (or sampling) sites as defined in Section 3 using random numbers from table 3, or from another Random Number Generator. Do not sample concrete from the first ½ cubic yard of the truck.

c. In order to determine which random values should be used, enter the table on a line chosen by chance. As a suggestion, select a line corresponding to the last two numbers on the first civilian license plate you see or other acceptable random means. Subsequent “X” values for following sublots on the same day are taken from the lines, which follow. Start each day with an “X” value determined by chance in order to obtain a random selection.

d. Example 4 shows the calculations for determining the testing location for Portland Cement Concrete.
Example 4
Test Location for a Sublot of Portland Cement Concrete

For this example the random number selected is “37.” Enter the random number table 3 at (37) and the corresponding four-digit number is 0.829, this is the factor.

Based on the delivery of 10 cubic yard loads to the project. This would be adjusted by the quantity of concrete actually being delivered per load.

Next five trucks loads => 10 CY x 5 = 50 CY

50 CY x 0.829 = 41 CY to be sampled

20 CY (first two trucks) + 41 CY = sample at the 61 CY point

Therefore, the sample will be taken from the truck containing the 61st CY. (This would be samples from the first 1/3 of the truck.) After approximately ½ CY of concrete has been discharged the sample should be taken. This is actually the seventh truckload delivered to the project this day as the first two truckloads were sampled before the random selection process started.

The next sample would be taken at random number “38.” Enter the random number table 3 at (39) and the corresponding four-digit number is 0.998, this is the factor.

Based on the delivery of 10 cubic yard loads to the project. This would be adjusted by the quantity of concrete actually being delivered per load.

Next five trucks loads => 10 CY x 5 = 50 CY

50 CY x 0.998 = 50 CY to be sampled

20 CY (first two trucks) 50 CY (from first random test) + 50 CY = sample at the 120 CY point. (This would be samples from the last 1/3 of the truck)

The next sample would be taken at random number “39.” Enter the random number table 3 at (38) and the corresponding four-digit number is 0.539, this is the factor.

Based on the delivery of 10 cubic yard loads to the project. This would be adjusted by the quantity of concrete actually being delivered per load.

Next five trucks loads => 10 CY x 5 = 50 CY

50 CY x 0.539 = 27 CY to be sampled

20 CY (first two trucks) 50 CY (from first random test) + (50 CY from second random test) + 27 CY = sample at the 147 CY point. (This would be samples from the middle to last 1/3 of the truck)
8. APPLICATIONS FOR AGGREGATE AND OTHER MATERIALS
   
a. Determine the lot or subplot size according to the contract documents. The lot or subplot shall be determined to the nearest 100 tons.

b. Determine the locations of the test (or sampling) sites as defined in Section 3 using random numbers from table 3, or from another Random Number Generator.

c. In order to determine which random values should be used, enter the table on a line chosen by chance. The first two or last two digits of the next automobile license plate you see is one way to select the entry point. Another way is to start a digital stopwatch and stop it several seconds later, using the decimal part of the seconds as your entry point.

**Sampling from a Belt or Flowing Stream:** Example: The specification calls for one sample from every 1000 Tons of aggregate. If the random number is 0.371, the sample would be taken at (0.371) (1000 Tons) = 371 Tons.

**Sampling from Haul Units:** Example: The specification calls for the samples to be based on a number of haul units. Determine the number of hauling units that comprise a lot. Multiply the selected random number(s) by the number of units to determine which unit(s) will be sampled.

If 20 haul units comprise a lot and one sample is needed, using the random number 0.773, the sample would be taken from the (0.773) (20) = 15.46, or 15th haul unit.

**Sampling from a Roadway with Previously Placed Material:** Example: The specification calls for a sample from a location on a job. The process as defined in Section 5, Applications for Asphalt Paving Density should be used where a X and Y measurement is needed to determine the testing location.
Appendix A  APPLICATIONS FOR HOT MIX ASPHALT DENSITY AND CHALLENGE CORES (metric Units)

a. Determine the LOT size and number of tests per LOT. The Standard specifications set the size of a density test lot for Asphalt Pavement to no greater than a single day’s production or approximately 400 tonne, whichever is less, and require five tests per LOT. At the end of a days production and the final lot is greater than 400 tonne, it should be broken up into two lots.

b. Convert this LOT size to an area segment of the roadway based on the roadway section and depth being constructed for the course being tested. The calculations in Example 1 show how this is performed. Table 1 has been provided to give you recommend lot lengths for standard lane widths at various depths. Lot length needs to be determined to the nearest 30 meters.

Example 1

Sample Computation for Lot Length (Metric Units)

Using nominal compacted density of 2 439 kg/m$^3$, compacted depth of 40 mm and paving width of 3.6 m:

Lot Length:

400 tonnes equate to 400 000 kg

Cross-section pavement area: 3.6 m wide, 0.040 m (40 mm) deep = 0.144 m$^2$

Unit weight per meter length = 0.144 m$^2$ * 2439 kg/m$^3$ = 351.2 kg/m

Length = 400 000 kg/351.2 kg/m = 1138.9 m round to 1140 m

Sublot length = 1140 m * 0.2 = 228 m

These typical figures may be revised based on the actual densities achieved or the yield results from the paving involved.

Table 1: Hot Mix Asphalt Density Test Lot Sizes Metric Units

<table>
<thead>
<tr>
<th>Lane Width</th>
<th>Compacted Depth</th>
<th>Computed Lot Length</th>
<th>Recommended Lot Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>40 mm</td>
<td>1139</td>
<td>1140</td>
<td></td>
</tr>
<tr>
<td>3.6 meters</td>
<td>1139</td>
<td>1140</td>
<td></td>
</tr>
<tr>
<td>60 mm</td>
<td>759</td>
<td>760</td>
<td></td>
</tr>
<tr>
<td>75 mm</td>
<td>607</td>
<td>610</td>
<td></td>
</tr>
<tr>
<td>40 mm</td>
<td>1242</td>
<td>1240</td>
<td></td>
</tr>
<tr>
<td>3.3 meters</td>
<td>1104</td>
<td>1100</td>
<td></td>
</tr>
<tr>
<td>60 mm</td>
<td>828</td>
<td>830</td>
<td></td>
</tr>
<tr>
<td>75 mm</td>
<td>663</td>
<td>660</td>
<td></td>
</tr>
</tbody>
</table>

LOT length may also be determined based on Nominal Designated LOT sizes. To utilize this concept, compacted mix volumes equivalent to the designated mix quantity per LOT have been determined using the nominal compacted unit weight of Hot Mix Asphalt pavement. These volumes are then converted into Density LOT lengths using the typical lane width and specified compacted depth. The included tables present the values for LOT Lengths based on English units.
c. Determine the locations of the test (or sampling) sites by using values from the random number table (Table 2) to determine the coordinate location on the roadway. In the table, use the “X” values as decimal fractions of the total length of the lot; use the “Y” values as fractions of the width, customarily measured from the right edge of the pavement. The values in the table have been set so that no measurements are taken within 1.5 LF (0.45 m) of the edge of the pavement. Whenever a test location is determined to fall within such an area (i.e., bridge end, track crossing, or night joint) the test location should be moved ahead or back on stationing, as appropriate, by 25 LF (8 m).

d. In order to determine which “X” and “Y” values should be used, enter the table on a line chosen by chance. Recommended procedure is selection of a line based on the last two digits from the most recent standard count on the nuclear density gage. Subsequent “X” and “Y” values are then taken from the lines that follow. Based on the specified sampling frequency, 20 lots can be accommodated by one cycle through the table. Start each shift with a set of values determined by chance in order to obtain random selection.

e. Example 2 shows the calculations for determining the testing location for asphalt pavement density.

Example 2
Test Location Within the LOT
for Hot Mix Asphalt Pavement Density (Metric Units)

For the lot defined above (3.6 m wide, 1140 m long) starting at station 10 000.00 m

Using the last two digits of the standard count. Determine the “X” and “Y” values from line (51) in the table: X = 0.762, Y = 0.65 (these are illustrative examples only. Table format and generation have been randomized so that each replication of the table will vary).

  Beginning station: 10 000.00
  Sublot length increment: 228 * 0.762 = 173.7 m
  Width offset: 3.6 * 0.65 = 2.3 m (from right edge)
  Location is station: 10 000 + 173.7 = 10 173.7, 2.2 m from right edge
**WSDOT Test Method T 718**

Method of test for Determining Stripping of Hot Mix Asphalt

1. **SCOPE**
   a. This test is used to determine the amount of stripping resulting from the effects of water saturation and accelerated water conditioning, with a freeze-thaw cycle of laboratory – compacted Hot Mix Asphalt.
   b. This test is the WSDOT equivalent to AASHTO T 283.

2. **EQUIPMENT**
   a. Water bath controlled at 140 ± 1.8° F.
   b. Vacuum container capable of holding a vacuum of approximately 26mm Hg and large enough to accommodate test samples and volume of water as described in this procedure.
   c. Perforated platform to hold test samples 2 inches off the bottom of the vacuum container.
   d. Vacuum pump, vacuum system or water aspirator, for vacuum saturation of samples.
   e. Air-bath freezer, maintained at 0 ± 5° F.
   f. Water bath maintained at 55 ± 1° F.
   g. Testing machine meeting requirements of AASHTO T 246 and capable of producing a uniform vertical movement of 0.065 inches per minute.

3. **PREPARATION OF LABORATORY-MIXED, LABORATORY-COMPACTED SAMPLES FOR MIX DESIGNS**
   a. Mix samples per WSDOT Test Method 726, at optimum asphalt binder content with appropriate grade and supplier of asphalt binder per the mix design to achieve approximately 4% air voids.
   b. Mix six samples per asphalt binder supplier, two samples with 0% anti-strip additive and the other samples with varying amounts of anti-strip additive (Note 1).
   
   **Note 1**-It is recommended that liquid anti-strip agents, added directly to the asphalt binder, be added at the level of ¼%, ½%, ¾% and 1% by weight of asphalt binder. Latex anti-strip agents must be added to the aggregate in a Saturated Surface Dry (SSD) condition at a level of 0.08%, 0.17%, 0.33% and 0.50% by weight of dry aggregate.
   c. Compact samples per WSDOT Test Method 702.

**PRECONDITIONING OF TEST SPECIMENS**
   a. Once the set of six samples have been compacted and cooled to room temperature, set one of the samples mixed with 0% anti-strip aside to be stored at room temperature, this will be the referee sample.
   b. Test remaining set of samples per AASHTO T166. Calculate the air void level of the sample using mix design Theoretical Maximum Specific Gravity value.
c. Place the samples in the vacuum container. The container must be filled with potable water at room temperature (77 ± 9°F) so that the samples have at least 1 inch of water above their surface. Apply a vacuum for a short amount of time, suitable to saturate the samples air voids between 60 and 80 percent.

d. Determine the mass of the saturated, surface-dry specimen after partial vacuum saturation by method A of T166.

e. Calculate the volume of absorbed water (J) in cubic centimeters by use of the following equation: 
   \[ J = B - A \]
   
   Where: 
   
   - \( J \) = volume of absorbed water, cubic centimeters.
   - \( B \) = mass of saturated, surface-dry specimen after partial vacuum.
   - \( A \) = mass of dry specimen in air.

f. Determine the degree of saturation (S) by comparing the volume of absorbed water (J) with the volume of air voids (Va) using the following equation.

   \[ S = \frac{100J}{Va} \]

   \( S \) = Degree of saturation, percent.

   \( Va \) = Volume of air voids

   Determine the Volume of air voids using the following equation:

   \[ Va = \frac{Pa \times E}{100} \]

   \( Pa \) = Percent of air voids

   E = Volume of Specimen, cubic centimeters. (SSD wt. – wt. In water)

g. If the degree of saturation is between 60 and 80 percent then proceed. If the degree of saturation is less than 60 percent then repeat the procedure beginning with c above, using more vacuum and/or time. If the degree of saturation is more than 80 percent then the sample has been damaged and must be discarded.

h. After saturation is achieved place each sample in a plastic bag, seal the bag and place sample in a freezer at a temperature of 0 ± 5°F for a minimum of 16 hours.

i. Remove samples from the freezer, remove plastic bags and place them in a water bath maintained at 140 ± 2°F for 24 ± 1 hour. (Note 2)

   Note 2: Some samples become fragile after curing in the hot bath for 24 hours, as a precaution it may be necessary to place samples into suitable transfer dishes prior to placing them into the hot bath, to facilitate the movement of samples for the hot bath to the cold-water bath.

j. After 24 ± 1 hour in the 140 ± 2°F water bath, remove the samples and place them into the cold water bath maintained at 55 ± 1°F. At this time the referee sample shall be placed into the cold water bath with the conditioned samples. Testing must begin within 2 hours ± 10 minutes after samples have been placed into the cold water bath.
TESTING

a. After 2 hours ± 10 minutes in the cold water bath, remove and test one sample at a time in the testing machine on the diametrical vertical plane. Apply the diametrical loading at a vertical deformation rate of 0.065 inches per minute. Record the maximum compressive load of each sample.

b. Continue to load sample until specimen can be easily broken open.

c. Remove sample from machine, break sample in half by hand for visual inspection. Record the visual condition of each sample as to stripping action: none, slight, moderate, or severe.

d. Determine the Tensile Strength Ratio (TSR) of each sample by comparing the load needed to break the testing sample to the load needed to break the referee sample, using the following equation: \[ \text{TSR} = \left( \frac{S1}{S2} \right) \times 100. \]

Where \( S1 \) = tensile strength of the conditioned sample
\( S2 \) = tensile strength of the unconditioned sample

A Tensile Strength Ratio of 80 or greater is required to be considered passing for this test.

VISUAL CONDITIONING DEFINITIONS

NONE: The sample condition is solid with no evidence of asphalt binder withdrawing from aggregate. After the sample has air-dried, the appearance is black.

SLIGHT: The sample condition is solid to slightly soft with evidence of the asphalt binder beginning to withdraw from edges and surfaces of the aggregates. After the sample has air-dried, the appearance remains black.

MODERATE: The sample condition is soft, easily broken in half, with partial to completely exposed aggregates. After the sample has air-dried, the appearance is slightly gray.

SEVERE: The sample condition is soft to falling apart with the majority of coarse aggregate completely exposed and asphalt binder almost nonexistent. After the sample has air-dried, the appearance is gray.

A sample must have a TSR of 80 or greater and exhibit no visual stripping to pass this test.

(Note 2)

Note 2: Asphalt Treated Base requires a TSR of 50 or greater and exhibit no visual stripping to pass this test.

REPORT

a. The report shall include the following: TSR and visual stripping of the specimen.
WSDOT Test Method T 720

Method of Test for Thickness Measurement for Hot Mix Asphalt (HMA) Cores

1. SCOPE
   a. This procedure is used to determine the thickness of the lifts in a Hot Mix Asphalt core.

2. APPARATUS
   a. Protection goggles, safety shield, or safety glasses.
   b. Hatchet.
   c. Striking tool such as a hammer, sledge, or maul suitable for striking the hatchet to separate the lifts.
   d. Tape, rule, calipers, or a measuring device suitable for measuring core lifts to 0.01 LF (3 mm).
   e. Hard stable surface, such as a cement concrete table, on which to place core for striking.
   f. Hard rubber pad.

3. PROCEDURE
   a. Measure the total thickness of the core as received to 0.01 LF (3 mm).
   b. Carefully remove all crushed surfacing top course, old pavement, prelevel, and prime coat from the core with the hatchet and striking tool.
   c. Measure the total thickness of the remaining core to 0.01 LF (3 mm).
   d. Split off the individual pavement lifts by placing core on the hard rubber pad, on the hard stable surface. Place the hatchet on the lift line and striking with the striking tool at several points around the core. Care must be taken in order to get a clean split of the core at the lift line and not damage the core.

   Note: Lift lines are often more visible by rolling the core on a flat surface. Chilling the cores may aid in splitting lifts.

   e. Each lift shall be measured from a plane surface to a plane surface. Two or more measurements shall be taken around the lift and the average shall be reported to 0.01 LF (3 mm) for each lift in the core.

   Note: The top lift is designated as lift number one. Each subsequent lift shall be designated as lifts 2, 3, 4, etc.

4. REPORT

   Report the results of the thickness measurements on DOT Form 350-067 (Thickness Measurements: Pavement & Treated Base Cores Transmittal/Report).
WSDOT Test Method T 724

Method of Preparation of Aggregate for HOT MIX ASPHALT (HMA) Mix Designs

1. SCOPE

This method of test is intended for the processing and preparation of aggregate samples for use in HMA mix designs and Ignition Furnace calibration samples for Hot Mix Asphalt, asphalt treated base, or open graded products.

2. APPARATUS

a. Sieves — shall conform to the specifications of sieves for testing purposes.

b. Mechanical sieve shaker — of sufficient size to separate the material to the specification sieves.

c. Oven(s) — of appropriate size, capable of maintaining a uniform temperature of 325 ± 25°F (163 ± 14°C).

d. Container — pans or containers of suitable size to dry and store the aggregate.

e. Balance — capacity of at least 8 kg sensitive to 0.1 g and meeting the requirements of AASHTO M231.

f. Aggregate washer (optional).

3. PROCEDURE

a. Representative sample(s) of the production aggregates shall be obtained.

b. Dry the aggregate in an oven to a constant mass not to exceed 350º F.

   Note: When developing an Ignition Furnace Calibration Factor, samples from separate stockpiles can be combined in the same percentages as the job mix formula prior to further processing. The combined sample should be at least four times the amount required for a single test (i.e., IFCF determination).

c. Sieve the aggregate over all the specification sieves designated for class of mix being tested. Place the material retained on each sieve in separate containers.

d. Wash the separated aggregate samples, except the portion passing the No. 200 (0.075 mm) sieve, in accordance with WSDOT FOP for WAQTC/AASHTO T 27/11.

e. Dry the washed, aggregate samples to constant mass.

f. Recombine the aggregate samples to match the grading of the job mix formula. The sample size as determined by the specific test procedure performed.
Performance Exam Checklist

Method of Preparation of Aggregate for Hot Mix Asphalt (HMA) Mix Designs

WSDOT Test Method T 724

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
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<tr>
<td>2. All equipment is functioning according to the test procedure,</td>
<td></td>
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<tr>
<td>and if required, has the current calibration/verification tags present?</td>
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<tr>
<td>3. Representative sample(s) of the production aggregates obtained per AASHTO T2?</td>
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<tr>
<td>4. Aggregate dried in an oven to a constant mass?</td>
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<tr>
<td>5. Aggregate sieved over designated sieves for class of mix being tested?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Material retained on each sieve placed in separate containers?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Washed separated aggregates, except the portion passing the No. 200 (0.075mm)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sieve, in accordance with WSDOT FOP for WAQTC/AASHTO T27/T11?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Washed aggregate samples dried in an oven to a constant mass?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Aggregate recombined to match the grading of the job mix formula?</td>
<td></td>
<td></td>
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<tr>
<td>10. Sample size determined by the specific test procedure to be performed?</td>
<td></td>
<td></td>
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</tbody>
</table>

First attempt: Pass [ ] Fail [ ]

Second attempt: Pass [ ] Fail [ ]

Signature of Examiner ________________________________________________

Comments:

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WSDOT Test Method T 726

Mixing Procedure for Hot Mix Asphalt (HMA)

1. SCOPE

This is the mixing procedure for laboratory prepared specimens of asphalt concrete, asphalt treated base, or open graded asphalt products. The aggregates used in this procedure are prepared by means of WSDOT Test Method No. 724.

2. EQUIPMENT

a. Mixing Spoon — A large metal spoon capable of handling hot mix asphalt.
b. Scoop — A metal scoop of ample size, capable of handling hot mix asphalt.
c. Curing Pan — A heat resistant pan of ample size to handle samples of hot mix asphalt.
d. Mixing Bowl — A heat resistant bowl for hand mixing or mechanical mixer of ample size to handle samples of hot mix asphalt.
e. Mechanical Mixer — A mechanical mixer with heat source may be used in lieu of hand mixing.
f. Balance — The balance shall have capacity of 11 kg and sensitive to 0.1 gm.
g. Oven — An oven of appropriate size, capable of maintaining a uniform temperature within the allowable tolerance for the grade of asphalt binder.

3. PROCEDURE

a. Place samples of aggregate in oven preheated to mixing temperature specified from supplier of asphalt binder or as indicated on mix design report for at least 2 hours.
b. Heat asphalt binder and mixing bowl(s) to mix temperature specified from supplier of asphalt binder or as indicated on mix design report.
c. Stir the asphalt binder and verify that the temperature of asphalt binder is within the temperature recommended by the asphalt supplier or as indicated on mix design report.
d. After materials are heated place mixing bowl on balance and tare.
e. Place heated aggregate in mixing bowl.
f. Form a crater in the aggregate and weigh in asphalt binder in accordance with design information, see calculation below.

Note: If mixing bowl is not buttered an additional sample should be prepared, mixed and then discarded to properly coat the mixing bowl with asphalt and fines.
g. Mix aggregate and asphalt binder for approximately 3 minutes or until aggregate is completely coated with asphalt binder. This can be accomplished by hand mixing or by mechanical mixer.
h. Transfer mixed material to the labeled heat resistant pan for curing or other testing as required.
i. Repeat steps A thru H for each sample to be mixed.
Calculation for Mass of Asphalt Binder:

Designated Mass of Asphalt binder = \( \frac{(A) \, D}{(1 - A)} \)

Where:  \( A \) = Designated asphalt binder content (expressed in decimal)
\( D \) = dry aggregate mass (from step 3(c))

Example:
The designated asphalt binder content is 5.3%, and dry aggregate mass is 1567.1 grams.

Designated Mass of Asphalt binder = \( \frac{(0.053) \times 1567.1}{(1 - 0.053)} \) = \( \frac{83.1}{0.947} \) = 87.7g
# Performance Exam Checklist

**Mixing Procedure for Hot Mix Asphalt (HMA)**  
**WSDOT Test Method T 726**

<table>
<thead>
<tr>
<th>Participant Name</th>
<th>Exam Date</th>
</tr>
</thead>
</table>

## Procedure Element

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Test</th>
<th>Retest</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Aggregate samples prepared as per WSDOT Test Method T724?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Mixing bowl(s), aggregate and asphalt binder heated to appropriate mixing temperature?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Asphalt binder stirred and temperature confirmed by thermometer?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Heated mixing bowl and paddle placed on scale and scale then tared?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Heated aggregate sample placed in bowl and scale then tared?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Crater formed into center aggregate, weigh in asphalt binder in accordance with mix design information?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>9. Mix aggregate and asphalt binder for approximately 3 minutes or until aggregate is completely coated?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>10. When mixing is complete carefully scrape off mixing apparatus, tools and bowl is dumped into correctly marked pan?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>11. Repeat steps 4 - 8 for each sample to be mixed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>12. All calculations performed correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐  
Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

________________________________________________________________
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WSDOT Standard Operating Procedure SOP 728

Method for Determining the Ignition Furnace Calibration Factor (IFCF) for Hot Mix Asphalt (HMA)

1. SCOPE
   This method may be affected by the type of aggregate in the mixture. Accordingly, to optimize accuracy, a calibration factor will be established with the testing of a set of HMA calibration samples for each mix type. This procedure must be performed before any acceptance testing is completed. The calibration process should be repeated each time there is a significant change in the mix ingredients or design.

2. APPARATUS
   a. Equipment as described to perform FOP for AASHTO T 308 Method A.

3. SAMPLE PREPARATION
   a. Prepare a minimum of two HMA calibration samples in accordance with WSDOT Test Method No. 724 and No. 726 or use previously prepared HMA calibration samples.
   b. If the HMA calibration samples are not sufficiently soft to separate for testing, carefully heat the samples in an oven until sufficiently soft and no appreciable moisture exists, not to exceed 325 ± 25 °F (63 ± 14 °C). Do not heat the sample basket assemblies.

4. PROCEDURE
   a. Test two HMA calibration samples in accordance with WSDOT FOP for AASHTO T 308.
   b. Determine the measured asphalt binder contents for each sample from the printed tickets.
   c. If the difference between the measured asphalt binder contents of the two samples exceeds 0.15 percent, test two additional HMA calibration samples. From the four tests, discard the high and low results and determine the IFCF from the two remaining results. Calculate the difference between the actual and measured asphalt binder contents for each sample. The IFCF is the average of the differences expressed in percent by mass of the HMA.
WSDOT Standard Operating Procedure SOP 729

In-Place Density of Bituminous Mixes Using the Nuclear Moisture-Density Gauge FOP for WAQTC TM 8

1. Number and Locations of Nuclear Tests
   a. Control lots representing 400 tons (400 metric tones) or less of mix shall be established. Nuclear gauge tests for compaction control during paving construction shall be taken at a minimum of five locations per control lot. The locations will be picked at random by WSDOT Test Method No. 716.

2. Theoretical Maximum Density determination FOR PAVEMENT COMPACtion CONTROL
   a. Theoretical Maximum Density is to be determined daily per WSDOT FOP for AASHTO T 209.
   b. On the initial day of production of a new Job Mix Formula (JMF), two determinations shall be made to establish an initial average value. The samples shall not be from the same truck. Average the two Theoretical Maximum Densities and report the result to the Moisture Density Gauge Operator. The Theoretical Maximum Density value from the Mix Design shall not be included in the average.
   c. If the two Theoretical Maximum Densities determined on the initial day do not agree within 3.0 lb./ft.³ (48 kg/m³), a third determination shall be made. The average density shall be based on the two closest sets of results.
   d. The moving average is defined as the average of the most recent last five determinations for the HMA being placed. All Theoretical Maximum Density determinations performed in a day or shift of paving will be included in the moving average. For Non Volumetric projects, a rice density test shall be taken with the first mix sample each day. For Volumetric projects, a rice density test shall be taken with each mix sample and all tests included in the moving average. Until five Theoretical Maximum Density values have been determined, the average will consist of the number of Theoretical Maximum Densities currently available. When five Theoretical Maximum Density values have been determined, the moving average for each day or shift will include the last four Theoretical Maximum Density determinations performed plus the first Theoretical Maximum Density determined for the current day or shift of paving. This new value will be used for the entire day or shift of paving.
   e. Subsequent Theoretical Maximum Density determinations shall be compared with the previously computed moving average. If a determination deviate from the moving average by more than 3.0 lb./ft.³ (± 48 kg/m³), a second determination shall be made on another portion of the same sample. If the second determination is within 3.0 lb./ft.³ (± 48 kg/m³) of the first determination a new moving average will be initiated, discarding all previous results. The new moving average will be sent to the Moisture Density Gauge operator and will replace the current moving average. If the second determination agrees within 3.0 lb./ft.³ (± 48 kg/m³) of the moving average then the first determination will be discarded and the second determination will be included in the moving average.
f. An average Theoretical Maximum Density (moving average) will be sent to the Moisture Density Gauge operator once per day or shift change, unless two determinations during a day or shift are not within 3.0 lb./ft.\(^3\) (± 48 kg/m\(^3\)), then a new moving average will be calculated in accordance with “e” of this procedure and sent to the Moisture Density Gauge operator as the new moving average for the day or shift. The Moisture Density Gauge Operator will continue to use the previous moving average until a new moving average is available.

3. Acceptance

a. For acceptable compaction, nuclear gauge test results for the control lot shall be determined by WAQTC FOP for TM8, as required by current specifications or contract plans.

b. The percent compaction equals the average of two inplace nuclear gauge wet density readings in accordance with TM8, times the gauge correlation factor divided by the current average Theoretical Maximum Density multiplied by 100.

\[
\text{percent compaction} = \frac{(\text{WD})(\text{CF})}{\text{Avg. Gmm}} \times 100
\]

\[
\text{WD} = \text{average of two inplace nuclear gauge wet density readings in accordance with TM8.}
\]

\[
\text{CF} = \text{gauge correlation factor.}
\]

\[
\text{Average Gmm} = \text{Avg. Theoretical Maximum Density}
\]

Report the percent compaction to tenth (0.1 percent)
WSDOT Standard Operating Procedure SOP 730

Correlation of Nuclear Gauge Densities with Hot Mix Asphalt (HMA) Cores

1. Gauge-core correlation shall be required for statistical evaluation of degree of asphalt compaction.
   a. For each combination of gauge and initial job mix formula.
   b. For direct transmission and for back scatter modes (when used).
   c. For a change in the class of HMA.

2. A new gauge correlation is not required.
   a. For different contracts if JMF and gauge are the same.
   b. For a change in bases (i.e., surfacing to overlay).
   c. When the job mix formula has been adjusted in accordance with Section 9-03.8(6)A of the Standard Specifications.

3. Gauge correlation is based on 10 density determinations and 10 cores taken at corresponding locations. Gauge densities shall be determined in accordance with WSDOT FOP for WAQTC TM 8. Cores should be taken no later than the day following paving and before traffic has been allowed on roadway. The sites for correlation cores do not have to be record density core sites and therefore consideration should be given to selecting sites out of the travel way.

   Note1: If a core becomes damaged, it may be eliminated from the average.

   Note2: Cores may be taken sooner than the day after paving by cooling the pavement to allow for hardening of the HMA to prevent damage to the core when taking the sample. Water, ice, or even dry-ice would be expedient means to cool the pavement. Nitrogen gas or CO2 uses as replacement drilling fluids may also be involved.

4. Obtain a pavement core from each of the test sites in accordance with WSDOT SOP 734. The core shall be taken in between the two nuclear gauge footprints. If direct transmission was used, locate the core at least 1 in. (25 mm) away from the edge of the drive pin hole.

5. Core densities shall be determined in conformance with AASHTO T 166 Bulk Specific Gravity of Compacted Bituminous Mixtures Using Saturated Surface-Dry Specimens.

6. Correlation factor shall be determined to 0.001 using Standard Form 350-112: Correlation Nuclear Gauge to Core Density, or other comparable forms.
WSDOT Standard Operating Procedure SOP 731

Method for Determining Volumetric Properties of Hot Mix Asphalt Class Superpave

1. SCOPE

This procedure covers the determination of volumetric properties of Asphalt Concrete Pavement Class Superpave i.e. Air Voids (Va), Voids in Mineral Aggregate (VMA), Voids Filled with Asphalt (VFA), and Dust to Binder Ratio (P_{200}/P_{be}).

2. REFERENCES

   a. T 329, WSDOT FOP for AASHTO Moisture content of Bituminous Mixtures by Oven
   b. T27/11, WSDOT FOP for WAQTC/AASHTO for Sieve Analysis of Fine and Coarse Aggregates
   c. T 166, WSDOT FOP for AASHTO for Bulk Specific Gravity of Compacted Bituminous Mixtures Using Saturated Surface-Dry Specimens
   d. T 168, WSDOT FOP for WAQTC/AASHTO for Sampling of Hot Mix Asphalt Paving Mixtures
   e. T 209, WSDOT FOP for AASHTO FOP for Maximum Specific Gravity of Hot Mix Asphalt Paving Mixtures “Rice Density”
   f. T 308, WSDOT FOP for AASHTO FOP for Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method
   g. T 312, WSDOT FOP for AASHTO for Preparing and Determining the Density of Hot Mix Asphalt (HMA) Specimens by Means of the Superpave Gyratory Compactor
   h. T 712, WSDOT Test Method for Standard Method of Reducing Hot Mix Asphalt Paving Mixtures

3. CALIBRATION OF COMPACTOR

   a. The gyratory compactor will be calibrated in accordance with WSDOT VP-58 and according to the manufacturer’s established calibration procedure. Anytime the gyratory compactor is moved to a new testing site a new calibration is required in accordance with WSDOT VP-58.

4. TEST SAMPLES

   a. All test samples shall be obtained per WSDOT FOP for WAQTC/AASHTO T 168, and reduced in accordance with WSDOT Test Method No. 712. It is recommended that the gyratory test sample be the first sample acquired in order to minimize heat loss.

   b. The size of the gyratory sample shall be such that it will produce a compacted specimen 115.0 ± 5.0 mm in height. Generally, the reference mix design verification report from the State materials Laboratory initial starting mass is adequate.

   c. Place the gyratory sample in an oven set no more than 25º F above the compaction temperature (Note 1) as soon as possible to reduce sample cooling. The gyratory test is temperature sensitive, so the sample should only be heated until it achieves the compaction temperature.

   Note: The compaction temperature for each mix design can be found on the mix design report. Any change in compaction temperature must be confirmed by the temperature viscosity chart provided by the asphalt supplier, which can be obtained from the paving Contractor.
5. PROCEDURE
   a. Place a compaction mold, base plate, and top plate (if required), in an oven set at no more than 25º F above compaction temperature (Note 2) for a minimum of 60 minutes prior to the estimated beginning of compaction. Subsequent uses of a conditioned mold will require 5 minutes reheating.

   **Note 2:** Never heat any gyratory compactor mold in excess of 350º F.

   b. Place a thermometer into the center of the mix, do not stir the mixture. (Note 3) Compact the sample immediately upon achieving compaction temperature.

   **Note 3:** While the gyratory test sample is heating it is beneficial to prepare and/or run the other tests as times permits.

   c. Perform the sample compaction in accordance with WSDOT FOP for AASHTO T312 section 9.

   d. Determine Rice Density per WSDOT FOP for AASHTO T 209.

   e. Determine asphalt content and gradation per WSDOT FOP for AASHTO T 308 and WSDOT FOP for WAQTC/AASHTO T27/11.

   f. Determine moisture content per WSDOT FOP for AASHTO T 329.

   g. Allow the gyratory compacted specimen to cool at room temperature for 15 to 24 hours. Determine the Bulk Specific Gravity (Gmb) of the specimen in accordance with WSDOT FOP for AASHTO T 166 Method A.

6. VOLUMETRIC CALCULATIONS
   a. Calculate %$G_{mm}$ @ $N_{design}$ as follows:

   \[
   \%G_{mm} @ N_{design} = \frac{G_{mb}}{G_{mm}} \times 100
   \]

   Example:

   \[
   \%G_{mm} @ N_{design} = \frac{2.383}{2.493} \times 100 = 95.6\%
   \]

   Where:

   $\%G_{mm}@N_{design}$ = % Theoretical Maximum Specific Gravity @ $N_{design}$

   $G_{mb}$ = bulk specific gravity of the compacted specimen

   $G_{mm}$ = maximum specific gravity of the paving mixture (Rice)

   $N_{design}$ = Number of design gyrations
b. Calculate $\%G_{\text{mm initial}}$ as follows:

$$\%G_{\text{mm initial}} = 100 \times \left( \frac{G_{\text{mb}} \times h_d}{G_{\text{mm}} \times h_i} \right)$$

Example:

$$\%G_{\text{mm initial}} = 100 \times \left( \frac{2.383 \times 110.0}{2.493 \times 123.1} \right) = 85.4\%$$

Where:

$\%G_{\text{mm initial}} = \%$ Theoretical Maximum Specific Gravity @ $N_{\text{initial}}$

$h_d = \text{height of specimen at design gyration level}$

$h_i = \text{height of specimen at initial design gyration level}$

$N_{\text{initial}} = \# \text{ of initial gyrations}$

c. Calculate Air Voids ($V_a$) as follow:

$$V_a = 100 \times \left( 1 - \frac{G_{\text{mb}}}{G_{\text{mm}}} \right)$$

Example:

$$V_a = 100 \times \left( 1 - \frac{2.383}{2.493} \right) = 4.4\%$$

Where:

$V_a = \%$ percent air voids

d. Calculate Voids in Mineral Aggregate (VMA) as follows:

$$VMA = 100 \times \left( 1 - \frac{G_{\text{mb}} \times P_s}{G_{\text{sb}}} \right)$$

Example:

$$VMA = 100 \times \left( 1 - \frac{2.383 \times 0.948}{2.630} \right) = 14.1\%$$

Where:

$P_s = \%$ percent of aggregate in the mix (use decimal form in calculation)

$P_s = 100 - \%$ asphalt binder

Example: 100% mix – 5.2% asphalt = 94.8% aggregate, use 0.948

$G_{\text{sb}} = \text{bulk specific gravity of the combined aggregate}$

$VMA = \%$ Voids in Mineral Aggregate, percent
e. Calculate Voids Filled with Asphalt (VFA) as follows:

\[
VFA = 100 \times \left( \frac{VMA - V_a}{VMA} \right)
\]

Example:

\[
VFA = 100 \times \left( \frac{14.1 - 4.4}{14.1} \right) = 68.8\%
\]

Where:

VFA = Voids Filled with Asphalt, percent

f. Calculate Gravity Stone Effective (G_{se}) as follows:

\[
G_{se} = \frac{100 - P_b}{\frac{100}{G_{mm}} - \frac{P_b}{G_b}}
\]

Example:

\[
G_{se} = \frac{100 - 5.2}{\frac{100}{2.493} - \frac{5.2}{1.025}} = 2.706
\]

Where:

G_{se} = Gravity Stone Effective (specific gravity of aggregates, excluding voids permeable to asphalt)

P_b = The percent by mass of binder in the total mixture including binder and aggregate

G_b = Gravity Binder

Note 4: G_b is the specific gravity of the asphalt binder. It is imperative that current G_b is used in the volumetric calculations. Any changes in the binder specific gravity must be confirmed by the temperature viscosity curve provided by the asphalt supplier, which can be obtained from the paving Contractor.

g. Calculate Percent Binder Effective (P_{be}) as follows:

\[
P_{be} = -P_s \times G_b \left( \frac{G_{se} - G_{sb}}{G_{se} \times G_{sb}} \right) + P_b
\]

Examples:

\[
P_{be} = -94.8 \times 1.025 \left( \frac{2.706 - 2.630}{2.706 \times 2.630} \right) + 5.2 = 4.2
\]

Where:

P_{be} = percent binder effective, the percent by mass of effective asphalt content minus the quantity of binder lost by absorption into the aggregate particles.

P_s = percent aggregate in the mixture

G_b = Gravity binder

G_{se} = effective specific gravity of the aggregate

G_{sb} = bulk specific gravity of the combined aggregate

P_b = percent binder
h. Calculate dust-to-binder ratio ($P_{200}/P_{be}$) as follows:

$$P_{200}/P_{be} = P_{200} ÷ P_{be}$$

Example: 5.0 ÷ 3.6 = 1.4

Where:

$P_{200}/P_{be}$ = dust-to-binder ratio

$P_{200}$ = percent of aggregate passing the No. 200 sieve

7. REPORT

Report asphalt content, gradation, and moisture content on WSDOT Form 350-560EF, and report volumetric properties on WSDOT Form 350-162 or other report approved by the State Materials Engineer.
WSDOT Standard Operating Procedure 732

Superpave-Volumetric Design for Hot-Mix Asphalt (HMA)

1. SCOPE

1.1. This standard for mix design evaluation uses aggregate and mixture properties to produce a hot-mix asphalt (HMA) job-mix formula. The mix design is based on the volumetric properties of the HMA in terms of the air voids (V_a), voids in the mineral aggregate (VMA), and voids filled with asphalt (VFA).

1.2. This standard may also be used to provide a preliminary selection of mix parameters as a starting point for mix analysis and performance prediction analyses that primarily use T 320 and T 322.

1.3. This standard may involve hazardous materials, operations, and equipment. This standard does not purport to address all of the safety problems associated with its use. It is the responsibility of the user of this procedure to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1. AASHTO Standards:

- M 320, Performance-Graded Asphalt Binder
- M 323, Superpave Volumetric Mix Design
- R 30, Mixture Conditioning of Hot-Mix Asphalt (HMA)
- R 35, Superpave Volumetric Design for Hot-Mix Asphalt (HMA)
- T 2, Sampling of Aggregates
- T 11, Materials Finer Than 75-μm (No. 200) Sieve in Mineral Aggregates by Washing
- T 27, Sieve Analysis of Fine and Coarse Aggregates
- T 84, Specific Gravity and Absorption of Fine Aggregate
- T 85, Specific Gravity and Absorption of Coarse Aggregate
- T 100, Specific Gravity of Soils
- T 166, Bulk Specific Gravity of Compacted Hot Mix Asphalt Using Saturated Surface-Dry Specimens
- T 209, Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt Paving Mixtures
- T 228, Specific Gravity of Semi-Solid Bituminous Materials
- T 248, Reducing Samples of Aggregate to Testing Size
- T 275, Bulk Specific Gravity of Compacted Bituminous Mixtures Using Paraffin-Coated Specimens
- T 283, Resistance of Compacted Asphalt Mixture to Moisture-Induced Damage
- T 304, Uncompacted Void Content of Fine Aggregate

1 This Standard Operating procedure is based on AASHTO T 323-04
• T 312, Preparing and Determining the Density of the Hot-Mix Asphalt (HMA) Specimens by Means of the Superpave Gyratory Compactor

• T 320, Determining the Permanent Deformation and Fatigue Cracking Characteristics of Hot-Mix Asphalt (HMA) Using the Simple Shear Test (SST) Device

• T 322, Determining the Creep Compliance and Strength of Hot-Mix Asphalt (HMA) Using the Indirect Tensile Test Device

2.2. Asphalt Institute:
• MS-2, Mix Design Methods for Asphalt Concrete and Other Hot-Mix Types

2.3. ASTM Standards:
• D4791 Flat Particles, Elongated Particles, or Flat and Elongated Particles in Coarse Aggregate

2.4. WSDOT Standards:
• M 41-01, Construction Manual
• M 41-10, Standard Specifications for Road, Bridge, and Municipal Construction (Std. Specs.)
• M 46-01, Materials Manual
• SOP 731 Method for Determining Volumetric Properties of Hot-Mix Asphalt (HMA)
• TM1, WAQTC FOP for Determining the Percentage of Fracture in Coarse Aggregate
• T2, WSDOT FOP for AASHTO for Standard Practice for Sampling Aggregate
• T27/11, WSDOT FOP for WAQTC/AASHTO for Sieve Analysis of Fine and Coarse Aggregates
• T113, Method of Test for Determination of Degradation Value
• T166, WSDOT FOP for AASHTO for Bulk Specific Gravity of Compacted Hot Mix Asphalt Using Saturated Surface-Dry Specimens
• T176, WSDOT FOP for AASHTO for Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test
• T209, WSDOT FOP for AASHTO for Method of Test for Maximum Specific Gravity of Hot Mix Asphalt Paving Mixtures “Rice Density”
• T248, WSDOT FOP for AASHTO for Reducing Samples of Aggregates to Testing Size
• T304, WSDOT Test Method for AASHTO T 304 Uncompacted Void Content of Fine Aggregate
• T 312, WSDOT FOP for AASHTO for Preparing and Determining the Density of Hot-Mix Asphalt (HMA) Specimens by Means of the Superpave Gyratory Compactor
• T 702, Method of Preparation of Test Specimens of Hot Mix Asphalt by Means of California Kneading Compactor
• T 718, Method of Test for Determining Stripping of Hot Mix Asphalt
• T 724, Method of Preparation of Aggregate for HMA Mix Designs
• T 726 Mixing Procedure for Hot-Mix Asphalt (HMA)
• D4791, WSDOT FOP for ASTM D4791 Standard Test Method for Flat Particles, Elongated Particles, or Flat and Elongated Particles in Coarse Aggregate
3. Terminology

3.1. **HMA – hot-mix asphalt**

3.2. **design ESALs**—design equivalent (80kN) single-axle loads

3.2.1. **Discussion**—design ESALs are the anticipated project traffic level expected on the design lane over a 15 to 20-year period. For pavements designed for more or less than 15 to 20 years, determine the design ESALs for 15 to 20 years when using this standard.

3.3. **air voids** ($V_a$)—the total volume of the small pockets of air between the coated aggregate particles throughout a compacted paving mixture, expressed as a percent of the bulk volume of the compacted paving mixture (Note 1).

**Note 1**—Term defined in Asphalt Institute Manual MS-2, Mix Design Methods for Asphalt Concrete and Other Hot-Mix Types.

3.4. **voids in the mineral aggregate** (VMA)—the volume of the intergranular void space between the aggregate particles of a compacted paving mixture that includes the air voids ($V_a$), and the effective binder content ($P_{be}$), expressed as a percent of the total volume of the specimen (Note 1).

3.5. **absorbed binder volume** ($V_{ab}$)—the volume of binder absorbed into the aggregate (equal to the difference in aggregate volume when calculated with the bulk specific gravity and effective specific gravity).

3.6. **binder content** ($P_b$)—The percent by mass of binder in the total mixture including binder and aggregate.

3.7. **effective binder volume** ($V_{be}$)—The volume of binder which is not absorbed into the aggregate.

3.8. **voids filled with asphalt** (VFA)—The percentage of the voids in the mineral aggregate (VMA) filled with binder (the effective binder volume divided by the VMA).

3.9. **dust to binder ratio** Dust/Asphalt Ratio ($P_{200}/P_{be}$)—By mass, ratio between percent passing the No. 200 (0.075-mm) sieve ($P_{200}$) and the effective binder content ($P_{be}$).

3.10. **nominal maximum aggregate size**—For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.

**WSDOT Note 1**: For an aggregate specification having a generally unrestrictive gradation (i.e. wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.

3.11. **maximum aggregate size**—One size larger than the nominal maximum aggregate size (Note 2).

**Note 2**—The definitions given in sections 3.10 and 3.11 apply to Superpave mixes only and differ from the definitions published in other AASHTO standards.

3.12. **reclaimed asphalt pavement** (RAP)—Removed and/or processed pavement materials containing asphalt binder and aggregate.
3.3. \(N_{\text{initial}}, N_{\text{design}}, N_{\text{maximum}}\)—the number of gyrations defined in WSDOT contract provisions.

3.4. Effective Asphalt Content, \(P_{\text{be}}\)—the total asphalt content of a paving mixture minus the portion of asphalt that is lost by absorption into the aggregate particles (Note 1).

4. Summary of the Practice

4.1. Materials Selection—Binder and aggregate and RAP stockpiles are selected that meet the environmental and traffic requirements applicable to the paving project. The bulk specific gravity of all aggregates proposed for blending and the specific gravity of the binder are determined.

Note 3—If RAP is used, the bulk specific gravity of the RAP aggregate may be estimated by determining the theoretical maximum specific gravity \(G_{\text{mm}}\) of the RAP mixture and using an assumed asphalt absorption for the RAP aggregate to back-calculate the RAP aggregate bulk specific gravity, if the absorption can be estimated with confidence. The RAP aggregate effective specific gravity may be used in lieu of the bulk specific gravity at the discretion of the Agency. The use of the effective specific gravity may introduce an error into the combined aggregate bulk specific gravity and subsequent VMA calculations. The Agency may choose to specify adjustments to the VMA requirements to account for this error based on experience with their local aggregates.

4.2. Design Aggregate Structure—It is recommended at least three trial aggregate blend gradations from selected aggregate stockpiles are blended. For each trial gradation, an initial trial binder content is determined, and at least two specimens are compacted in accordance with WSDOT FOP for AASHTO T 312. A design aggregate structure and an estimated design binder content are selected on the basis of satisfactory conformance of a trial gradation meeting the requirements given in Section 9-03.8(2) of the Standard Specifications for Road, Bridge, and Municipal Construction (Std. Specs.) M 323 for \(V_{\alpha}, \text{VMA, VFA, dust-to-binder ratio } \text{Dust/Asphalt Ratio} \) at \(N_{\text{design}}\), and relative density at \(N_{\text{initial}}\).

Note 4—Previous Superpave mix design experience with specific aggregate blends may eliminate the need for three trial blends.

4.3. Design Binder Content Selection—Replicate specimens are compacted in accordance with WSDOT FOP for AASHTO T 312 at the estimated design binder content and at the estimated design binder content ±0.5 percent and ±1.0 percent. The design binder content is selected on the basis of satisfactory conformance with the requirements of Section 9-03.8(2) of the Std. Specs. M 323 for \(V_{\alpha}, \text{VMA, VFA, and dust-to-binder ratio } \text{Dust/Asphalt Ratio} \) \(P_{\text{be}}/P_{\text{be}}\) at \(N_{\text{des}}\) and the relative density at \(N_{\text{ini}}\) and \(N_{\text{max}}\). For WSDOT projects, see contract provisions the design binder content selection is determined by the Contractor and is verified by the WSDOT.

4.4. Evaluating Moisture Susceptibility—The moisture susceptibility of the design aggregate structure is evaluated at the design binder content: the mixture is conditioned according to the mixture conditioning for the volumetric mixture design procedure in R 30, compacted to approximately 4.0 ± 1.0 percent air voids in accordance with T 342 WSDOT T 702, and evaluated according to T 282 WSDOT T 718. The design shall meet the tensile strength ratio requirement of MP 2 WSDOT T 718. The WSDOT State Materials Laboratory will evaluate the HMA for moisture susceptibility.

5. Significance and Use

5.1. The procedure described in this practice is used to produce HMA which satisfies Superpave HMA volumetric mix design requirements.
6. Preparing Aggregate Trial Blend Gradations

6.1. Select a binder in accordance with the requirements of M 323. For WSDOT projects, see contract provisions. The asphalt binder grade will be indicated in WSDOT Contract Special Provisions.

6.2. Determine the specific gravity of the binder according to T 228.

6.3. Obtain samples of aggregates proposed to be used for the project from the aggregate stockpiles in accordance with WSDOT FOP for AASHTO T 2.

Note 5—Each stockpile usually contains a given size of an aggregate fraction. Most projects employ three to five stockpiles to generate a combined gradation conforming to the job-mix formula and Section 9-03.8(6) of the Std. Specs.M 323.

6.4. Reduce the samples of aggregate fractions according to WSDOT FOP for AASHTO T 248 to samples of the size specified in WAQTC FOP for AASHTO T 27/T 11.

6.5. Wash and grade each aggregate sample according to WAQTC FOP for AASHTO T 27/T 11.

6.6. Determine the bulk and apparent specific gravity for each coarse and fine aggregate fraction in accordance with T 85 and T 84, respectively, and determine the specific gravity of the mineral filler in accordance with T 100. WSDOT requires specific gravity determinations to be reported to an accuracy of 0.001.

6.7. Blend the aggregate fractions using Equation 1:

\[ P = Aa + Bb + Cc, \text{ etc.} \]

where:

- \( P \) = percentage of material passing a given sieve for the combined aggregates \( A, B, C, \text{ etc.} \);
- \( A, B, C, \text{ etc.} \) = percentage of material passing a given sieve for aggregates \( A, B, C, \text{ etc.} \); and
- \( a, b, c, \text{ etc.} \) = proportions of aggregates \( A, B, C, \text{ etc.} \) used in the combination, and where the total = 1.00.

6.8. Prepare a minimum of three trial aggregate blend gradations; plot the gradation of each trial blend on a 0.45-power gradation analysis chart, and confirm that each trial blend meets the Aggregate Gradation Control Points in Section 9-03.8(6) of the Std. Specs. M 323 gradation controls (see Tables 32 of M 323). Gradation control is based on four control sieve sizes: the sieve for the maximum aggregate size, the sieve for the nominal maximum aggregate size, the No. 4 or No. 8 (4.75- or 2.36-mm) sieve, and the No. 200 (0.075-mm) sieve. For WSDOT projects, gradation shall be determined by the following sieves as defined in table W1T. An example of three acceptable trial blends in the form of a gradation plot is given in Figure 1.
Sieves Required for Gradation Determination

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>3/8 In.</th>
<th>1/2 In.</th>
<th>3/4 In.</th>
<th>1 In.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 1/2&quot;</td>
<td></td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>1&quot;</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 4</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 8</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 16</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 30</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 50</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 100</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>U.S. No. 200</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

X = indicates sieve is required for gradation determination

Table W1T

6.9. Obtain a test specimen from each of the trial blends according to WSDOT FOP for AASHTO T 248, and conduct the quality tests specified in Section 9-03.8(2) subsections 1, 2, 3, and 4 of the Std. Specs. M 323 to confirm that the aggregate in the trial blends meets the minimum quality requirements specified in Section 9-03.8(2) of the Std. Specs. M 323.

Note 6—The designer has an option of performing the quality tests on each stockpile instead of the trial aggregate blend. The test results from each stockpile can be used to estimate the results for a given combination of materials.

Figure 1—Evaluation of the Gradations of Three Trial Blends (Example)
7. Determining an Initial Trial Binder Content for Each Trial Aggregate Gradation

7.1. Designers can either use their experience with the materials or the procedure given in Appendix A1 to determine an initial trial binder content for each trial aggregate blend gradation.

*Note 7*—When using RAP, the initial trial asphalt content should be reduced by an amount equal to that provided by the RAP.

8. Compacting Specimens of Each Trial Gradation

8.1. Prepare replicate mixtures (Note 8) at the initial trial binder content for each of the chosen trial aggregate trial blend gradations. From Table 1, determine the number of gyrations based on the design ESALs for the project. On WSDOT projects the ESAL level number of gyrations will be indicated in the Contract Special Provisions.

*Note 8*—At least two replicate specimens are required, but three or more may be prepared if desired. Generally, 4500 to 4700 g of aggregate is sufficient for each compacted specimen with a height of 110 to 120 mm for aggregates with combined bulk specific gravities of 2.550 to 2.700, respectively.

8.2. Condition the mixtures according to R 30, and compact the specimens to $N_{\text{design}}$ gyrations in accordance with WSDOT FOP for AASHTO T 312. Record the specimen height to the nearest 0.1 mm after each revolution.

8.3. Determine the bulk specific gravity ($G_{mb}$) of each of the compacted specimens in accordance with WSDOT FOP for AASHTO T 166 or T 275 as appropriate. The bulk specific gravity results of the replicate specimens samples shall not differ by more than 0.020.

**Table 1**—Superpave Gyratory Compaction Effort

| Design ESALs a (million) | Compaction Parameters | Typical Roadway Application
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$N_{\text{trial}}$</td>
<td>$N_{\text{design}}$</td>
</tr>
<tr>
<td>&lt; 0.3</td>
<td>6</td>
<td>50</td>
</tr>
<tr>
<td>Applications include roadways with very light traffic volumes such as local roads, county roads, and city streets where truck traffic is prohibited or at a very minimal level. Traffic on these roadways would be considered local in nature, not regional, intrastate, or interstate. Special purpose roadways serving recreational sites or areas may also be applicable to this level.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3 to &lt; 3</td>
<td>7</td>
<td>75</td>
</tr>
<tr>
<td>Applications include many collector roads or access streets. Medium-traffic city streets and the majority of county roadways may be applicable to this level.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 to &lt; 30</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
<td>Applications include many two-lane, multilane, divided, and partially or completely controlled access roadways. Among these are medium to highly trafficked city streets, many state routes, U.S. highways, and some rural Interstates.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\geq$ 30</td>
<td>9</td>
<td>125</td>
</tr>
<tr>
<td>Applications include the vast majority of the U.S. Interstate system, both rural and urban in nature. Special applications such as truck-weighing stations or truck-climbing lanes on two-lane roadways may also be applicable to this level.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

* The anticipated project traffic level expected on the design lane over a 15-20-year period. Regardless of the actual design life of the roadway, determine the design ESALs for 15-20 years.
As defined by *A Policy on Geometric Design of Highways and Streets, 2001*, AASHTO.

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**Note 9**—When specified by the Agency and the top of the design layer is ≥100 mm from the pavement surface and the estimated design traffic level is ≥0.3 million ESALs, decrease the estimated design traffic level by one, unless the mixture will be exposed to significant mainline construction traffic prior to being overlaid.

**Note 10**—When the estimated design traffic level is between 3 and <10 million ESALs, the Agency may, at its discretion, specify $N_{\text{min}}$ at 7, $N_{\text{design}}$ at 75, and $N_{\text{max}}$ at 115.

8.4. Determine the theoretical maximum specific gravity ($G_{\text{mm}}$) according to WSDOT FOP for AASHTO T 209 of separate samples representing each of these combinations that have been mixed and conditioned to the same extent as the compacted specimens.

**Note 11**—The maximum specific gravity for each trial mixture shall be based on the average of at least two tests. The maximum specific gravity results of the replicate specimens samples shall not differ by more than 0.011.

9. Evaluating Compacted Trial Mixtures

9.1. Determine the volumetric requirements for the trial mixtures in accordance with Section 9-03.8(2) of the Std. Specs. M 323.

9.2. Calculate $V_a$ and VMA at $N_{\text{design}}$ for each trial mixture using equations 2 and 3:

\[ V_a = 100 \times \left(1 - \frac{G_{\text{mb}}}{G_{\text{mm}}}\right) \]  \hspace{1cm} (2)

\[ \text{VMA} = 100 \times \left(1 - \frac{G_{\text{mb}} P_i}{G_{\text{sb}}}\right) \]  \hspace{1cm} (3)

where:

- $G_{\text{mb}}$ = bulk specific gravity of the extruded specimen;
- $G_{\text{mm}}$ = theoretical maximum specific gravity of the mixture;
- $P_i$ = percent of aggregate in the mix in decimal form; and
- $G_{\text{sb}}$ = bulk specific gravity of the combined aggregate.

**Note 12**—Although the initial trial binder content was estimated for a design air void content of 4.0 percent, the actual air void content of the compacted specimen is unlikely to be exactly 4.0 percent. Therefore, the change in binder content needed to obtain a 4.0 percent air void content, and the change in VMA caused by this change in binder content, is estimated. These calculations permit the evaluation of VMA and VFA of each trial aggregate gradation at the same design air void content, 4.0 percent.
9.3. Estimate the volumetric properties at 4.0 percent air voids for each compacted specimen. On WSDOT projects the $V_\text{a}$, VMA, VFA criteria, and gyration level, will be specified in the Contract Provisions.

9.3.1. Determine the difference in average air void content at $N_{\text{design}}$ ($\Delta V_a$) of each aggregate trial blend from the design level of 4.0 percent using Equation 4:

$$VMA = 100 \times \left(1 - \frac{G_{\text{mb}}P}{G_{\text{ib}}}\right)$$

(4)

where:

$V_a$ = air void content of the aggregate trial blend at $N_{\text{design}}$ gyrations.

9.3.2. Estimate the change in binder content ($\Delta P_b$) needed to change the air void content to 4.0 percent using Equation 5:

$$\Delta P_b = -0.4(\Delta V_a)$$

(5)

9.3.3. Estimate the change in VMA ($\Delta VMA$) caused by the change in the air void content ($\Delta V_a$) determined in Section 9.3.1 for each trial aggregate blend gradation, using Equations 6 or 7.

$$\Delta VMA = 0.2(\Delta V_a) \text{ if } V_a > 4.0$$

(6)

$$\Delta VMA = 0.1(\Delta V_a) \text{ if } V_a < 4.0$$

(7)

Note 13—A change in binder content affects the VMA through a change in the bulk specific gravity of the compacted specimen ($G_{\text{mb}}$).

9.3.4. Calculate the VMA for each aggregate trial blend at $N_{\text{design}}$ gyrations and 4.0 percent air voids using Equation 8:

$$VMA_{\text{design}} = VMA_{\text{trial}} + \Delta VMA$$

(8)

where:

$VMA_{\text{design}}$ = VMA estimated at a design air void content of 4.0 percent; and

$VMA_{\text{trial}}$ = VMA determined at the initial trial binder content.

9.3.5. Using the values of $\Delta V_a$ determined in Section 9.3.1 and Equation 9, estimate the relative density of each specimen at $N_{\text{initial}}$ when the design air void content is adjusted to 4.0 percent at $N_{\text{design}}$:

$$\%G_{\text{mb,initial}} = 100 \times \left(\frac{G_{\text{mb}} h_d}{G_{\text{mb}} h_i}\right) - \Delta V_a$$

(9)

where:

$\%G_{\text{mb,initial}}$ = relative density at $N_{\text{initial}}$ gyrations at the adjusted design binder content;

$h_d$ = height of the specimen after $N_{\text{design}}$ gyrations, from the Superpave gyratory compactor, mm; and

$h_i$ = height of the specimen after $N_{\text{initial}}$ gyrations, from the Superpave gyratory compactor, mm.
9.3.6. Estimate the percent of effective binder (P) and calculate the Dust/Asphalt Ratio \( \left( \frac{P_{200}}{P_{be}} \right) \) for each trial blend using Equations 10 and 11:

\[
P_{be} = \left( P_s \times G_b \right) \left( \frac{G_w - G_{sb}}{G_{se} \times G_{sb}} \right) + P_{wter}
\]

where:

- \( P_{be} \) = estimated effective binder content,
- \( P_s \) = aggregate content, \((100 - P_s)\)
- \( G_b \) = specific gravity of the binder,
- \( G_{se} \) = effective specific gravity of the aggregate,
- \( G_{sb} \) = bulk specific gravity of the combined aggregate, and
- \( P_{wter} \) = estimated binder content.

\[
\text{Dust/Asphalt Ratio} = \frac{P_{200}}{P_{be}}
\]

where:

- \( P_{200} \) = percent passing the U.S. No. 200 (0.075 mm) sieve.

9.3.7. Compare the estimated volumetric properties from each trial aggregate blend gradation at the adjusted design binder content with the criteria specified in Section 9-03.8(2) of the Std. Specs. Choose the trial aggregate blend gradation that best satisfies the volumetric criteria.

**Note 14**—Table 2 presents an example of the selection of a design aggregate structure from three trial aggregate blend gradations.

**Note 15**—Many trial aggregate blend gradations will fail the VMA criterion. Generally, the % criterion will be met if the VMA criterion is satisfied. Section 12.1 gives a procedure for the adjustment of VMA.

**Note 16**—If the trial aggregate gradations have been chosen to cover the entire range of the gradation controls, then the only remaining solution is to make adjustments to the aggregate production or to introduce aggregates from a new source. The aggregates that fail to meet the required criteria will not produce a quality mix and should not be used. One or more of the aggregate stockpiles should be replaced with another material which produces a stronger structure. For example, a quarry stone can replace a crushed gravel, or crushed fines can replace natural fines.
Table 2—Selection of a Design Aggregate Structure (Example)

<table>
<thead>
<tr>
<th>Volumetric Property</th>
<th>Trial Mixture (3/4 Inch 19.0-mm Nominal Maximum Aggregate)</th>
<th>Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>15 20-Year Project Design ESALs = 5 million</td>
<td></td>
</tr>
<tr>
<td>At the Initial Trial Binder Content</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$P_b$ (trial)</td>
<td>4.4</td>
<td>4.4</td>
</tr>
<tr>
<td>$%G_{	ext{mm}}$ (trial)</td>
<td>88.1</td>
<td>87.8</td>
</tr>
<tr>
<td>$%G_{	ext{mm}}$ (trial)</td>
<td>95.9</td>
<td>95.3</td>
</tr>
<tr>
<td>$V_a$ at $N_{\text{design}}$</td>
<td>4.1</td>
<td>4.7</td>
</tr>
<tr>
<td>VMA$_{\text{trial}}$</td>
<td>12.9</td>
<td>13.4</td>
</tr>
<tr>
<td>Adjustments to Reach Design Binder Content ($V_a = 4.0%$ at $N_{\text{design}}$)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\Delta V_a$</td>
<td>−0.1</td>
<td>−0.7</td>
</tr>
<tr>
<td>$\Delta P_b$</td>
<td>0.0</td>
<td>0.3</td>
</tr>
<tr>
<td>$\Delta$VMA</td>
<td>0.0</td>
<td>−0.1</td>
</tr>
<tr>
<td>At the Estimated Design Binder Content ($V_a = 4.0%$ at $N_{\text{design}}$)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Estimated $P_b$ (design)</td>
<td>4.4</td>
<td>4.7</td>
</tr>
<tr>
<td>VMA (design)</td>
<td>12.9</td>
<td>13.3</td>
</tr>
<tr>
<td>$%G_{	ext{mm}}$ (design)</td>
<td>88.2</td>
<td>89.5</td>
</tr>
</tbody>
</table>

Notes:

1. The top portion of this table presents measured densities and volumetric properties for specimens prepared for each aggregate trial blend at the initial trial binder content.
2. None of the specimens had an air void content of exactly 4.0 percent. Therefore, the procedures described in Section 9 must be applied to:
   (1) estimate the design binder content at which $TV_a = 4.0\%$, and (2) obtain adjusted VMA and relative density values at this estimated binder content.
3. The middle portion of this table presents the change in binder content ($\Delta P_b$) and VMA ($\Delta$VMA) that occurs when the target air void content ($TV_a$) is adjusted to 4.0 percent for each trial aggregate blend gradation.
4. A comparison of the VMA and densities at the estimated design binder content to the criteria in the last column shows that trial aggregate blend gradation No. 1 does not have sufficient VMA (12.9 percent versus a requirement of >13.0 percent). Trial blend No. 2 exceeds the criterion for relative density at $N_{\text{initial}}$ gyrations (89.5 percent versus requirement of ≤89.0 percent). Trial No. 3 meets the requirement for relative density and VMA and, in this example, is selected as the design aggregate structure.
10. Selecting the Design Binder Content

10.1. Prepare replicate mixtures (Note 8) containing the selected design aggregate structure at each of the following four binder contents: (1) the estimated design binder content, \( P_{b \text{ (design)}} \); (2) 0.5 percent below \( P_{b \text{ (design)}} \); (3) 0.5 percent above \( P_{b \text{ (design)}} \); and (4) 1.0 percent above \( P_{b \text{ (design)}} \).

10.1.1. Use the number of gyrations previously determined in Section 8.1.

10.2. Condition the mixtures according to R 30, and compact the specimens to \( N_{\text{design}} \) gyrations according to WSDOT FOP for AASHTO T 312. Record the specimen height to the nearest 0.1 mm after each revolution.

10.3. Determine the bulk specific gravity of each of the compacted specimens in accordance with WSDOT FOP for AASHTO T 166 or AASHTO T 275 as appropriate.

10.4. Determine the theoretical maximum specific gravity (\( G_{\text{mm}} \)) according to WSDOT FOP for AASHTO T 209 of each of the four mixtures using companion samples which have been conditioned to the same extent as the compacted specimens (Note 8).

10.5. Determine the design binder content which produces a target air void content of 4.0 percent at \( N_{\text{design}} \) gyrations using the following steps:

10.5.1. Calculate \( V_{a} \), VMA, and VFA at \( N_{\text{design}} \) using Equations 2, 3 and 12: The volumetric properties are determined for each specimen and then averaged for each replicate mixture.

\[
VFA = 100 \times \frac{VMA - V_{a}}{VMA} \tag{12}
\]

10.5.2. Calculate the dust-to-binder ratio Dust/Asphalt Ratio, using Equation 13.

\[
\text{Dust / Asphalt Ratio} = \frac{P_{200}}{P_{\text{be}}} \tag{13}
\]

where:

\( P_{\text{be}} = \) effective binder content.

10.5.3. For each of the four mixtures, determine the average corrected specimen relative densities at \( N_{\text{initial}} \) (%), using Equation 14.

\[
\%G_{\text{mm, initial}} = 100 \times \left( \frac{G_{\text{mm, i}}}{G_{\text{mm, h}}} \right) \tag{14}
\]

10.5.4. Plot the average \( V_{a} \), VMA, VFA, and relative density at \( N_{\text{design}} \) for replicate specimens versus binder content.

**Note 17**—All plots are generated automatically by the Superpave software. Figure 2 presents a sample data set and the associated plots.

10.5.5. By graphical or mathematical interpolation (Figure 2), determine the binder content to the nearest 0.1 percent at which the target \( V_{a} \) is equal to 4.0 percent. This is the design binder content (\( P_{b} \)) at \( N_{\text{design}} \).

10.5.6. By interpolation (Figure 2), verify that the volumetric requirements specified in Section 9-03.8(2) of the Std. Specs. M-323 are met at the design binder content.
10.6. Compare the calculated percent of maximum relative density with the design criteria at $N_{\text{initial}}$ by interpolation, if necessary. This interpolation can be accomplished by the following procedure.

10.6.1 Prepare a densification curve for each mixture by plotting the measured relative density at $x$ gyrations, $\%G_{\text{mm}}$, versus the logarithm of the number of gyrations (see Figure 3).

10.6.2 Examine a plot of air void content versus binder content. Determine the difference in air voids between 4.0 percent and the air void content at the nearest, lower binder content. Determine the air void content at the nearest, lower binder content at its data point, not on the line of best fit. Designate the difference in air void content as $\Delta V_a$.

10.6.3 Using Equation 14, determine the average corrected specimen relative densities at $N_{\text{initial}}$. Confirm that satisfies the design requirements in Section 9-03.8(2) of the Std. Specs. M 323 at the design binder content.

10.7. Prepare replicate (Note 8) specimens composed of the design aggregate structure at the design binder content to confirm that $\%G_{\text{mm}}$ satisfies the design requirements in Section 9-03.8(2) of the Std. Specs. M 323.

10.7.1 Condition the mixtures according to R-30, and compact the specimens according to WSDOT FOP for AASHTO T312 to the maximum number of gyrations, $N_{\text{max}}$, from Section 9-03.8(2) of the Std. Specs. Table I.

10.7.2 Determine the average specimen relative density at $N_{\text{max}}$, $\%G_{\text{mm}}$, by using Equation 15, and confirm that $\%G_{\text{mm}}$ satisfies the volumetric requirement in Section 9-03.8(2) of the Std. Specs. M 323,

$$\%G_{\text{mm}} = 100 \frac{G_{\text{mb}}}{G_{\text{mm}}}$$

where:

$\%G_{\text{mm}}$ relative density at $N_{\text{max}}$ gyrations at the design binder content.

**Figure 2 - Sample Volumetric Design Data at $N_{\text{des}}$**

<table>
<thead>
<tr>
<th>$P_0$(%)</th>
<th>$V_i$(%)</th>
<th>VMA (%)</th>
<th>VFA (%)</th>
<th>Maximum Density at $N_{\text{design}}$ (Gmm)</th>
<th>Density at $N_{\text{design}}$ lbs/ft$^3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.3</td>
<td>9.9</td>
<td>17.0</td>
<td>41.8</td>
<td>2.660</td>
<td>165.6</td>
</tr>
<tr>
<td>4.8</td>
<td>8.2</td>
<td>16.7</td>
<td>50.9</td>
<td>2.636</td>
<td>164.1</td>
</tr>
<tr>
<td>5.3</td>
<td>6.9</td>
<td>16.6</td>
<td>58.5</td>
<td>2.617</td>
<td>162.9</td>
</tr>
<tr>
<td>5.8</td>
<td>5.2</td>
<td>16.5</td>
<td>68.5</td>
<td>2.585</td>
<td>160.9</td>
</tr>
<tr>
<td>6.3</td>
<td>3.9</td>
<td>16.2</td>
<td>76.0</td>
<td>2.574</td>
<td>160.2</td>
</tr>
</tbody>
</table>

In this example, the estimated design binder content is 4.8 percent; the minimum VMA requirement for the design aggregate structure (3/4 inch nominal maximum size) is 13.0 percent, and the VFA requirements is 65 to 78 percent.

Entering the plot of percent air voids versus percent binder content at 4.0% percent air voids, the design binder content is determined as 6.2% percent.

Entering the plots of percent VMA versus percent binder content and percent VFA versus percent binder content at 6.2% percent binder content, the mix meets the VMA and VFA requirement.
Figure 3 - Sample Densification Curve
11. Evaluating Moisture Susceptibility

11.1. Prepare six mixture specimens (nine are needed if freeze-thaw testing is required) composed of the design aggregate structure at the design binder content. Condition the mixtures in accordance with R 30, and compact the specimens to 7 ± 1.0 percent air voids in accordance with T 312. Prepare the specimens according to WSDOT T 726, and compact the specimens to approximate 4.0% air voids in accordance to WSDOT T 702. The WSDOT State Materials Laboratory will evaluate the HMA for moisture susceptibility.

11.2. Test the specimens and calculate the tensile strength ratio in accordance with T 283 WSDOT T 718.

11.3. If the tensile strength ratio is less than 0.80, as required in MP 2, remedial action such as the use of anti-strip agents is required to improve the moisture susceptibility of the mix. When remedial agents are used to modify the binder, retest the mix to assure compliance with the 0.80 minimum requirement.

12. Adjusting the Mixture to Meet Properties

12.1. Adjusting VMA—If a change in the design aggregate skeleton is required to meet the specified VMA, there are three likely options: (1) change the gradation (Note 18); (2) reduce the minus U.S. No. 200 (0.075-mm) fraction (Note 19); or (3) change the surface texture and/or shape of one or more of the aggregate fractions (Note 20).

Note 18—Changing gradation may not be an option if the trial aggregate blend gradation analysis includes the full spectrum of the gradation control area.

Note 19—Reducing the percent passing the U.S. No. 200 (0.075-mm) sieve of the mix will typically increase the VMA. If the percent passing the U.S. No. 200 (0.075-mm) sieve is already low, this is not a viable option.

Note 20—This option will require further processing of existing materials or a change in aggregate sources.
12.2. **Adjusting VFA**—The lower limit of the VFA range should always be met at 4.0 percent air voids if the VMA meets the requirements. If the upper limit of the VFA is exceeded, then the VMA is substantially above the minimum required. If so, redesign the mixture to reduce the VMA. Actions to consider for redesign include: (1) changing to a gradation that is closer to the maximum density line; (2) increasing the minus U.S. No. 200 (0.075-mm) fraction, if room is available within the specification control points; or (3) changing the surface texture and shape of the aggregates by incorporating material with better packing characteristics, e.g., less thin, elongated aggregate particles.

12.3. **Adjusting the Tensile Strength Ratio**—The tensile strength ratio can be increased by: (1) adding chemical anti-strip agents to the binder to promote adhesion in the presence of water; or (2) adding hydrated lime to the mix.

13. **Report**

13.1. The report shall include the identification of the project number, mix class designation, traffic level, and mix design number.

13.2. The report shall include information on the design aggregate structure including the source of aggregate, kind of aggregate, required quality characteristics, and gradation, including the blending ratios.

13.3. The report shall contain information about the design binder including the source of binder and the performance grade.

13.4. The report shall contain information about the HMA including the percent of binder in the mix; the relative density; the number of initial, design, and maximum gyrations; and the VMA, VFA, Vw, Vw, Vav and dust-to-binder ratio Dust/Asphalt Ratio P200/Pbe, Pbe, Gmm, Gmb, Gsb and Gse of the aggregate blend, Gs of the fine aggregate, and Gw.

13.5. The report shall contain the results of the moisture susceptibility testing and the required level of anti-strip additive needed.

14. **Keywords**

14.1. HMA mix design; Superpave; volumetric mix design.
Appendix

A1. Calculating an Initial Trial Binder Content for Each Aggregate Trial Blend

Nonmandatory information

A1.1 Calculate the bulk and apparent specific gravities of the combined aggregate in each trial blend using the specific gravity data for the aggregate fractions obtained in Section 6.6 and Equations 16 and 17:

\[ G_{sb} = \frac{P_1 + P_2 + \cdots + P_n}{G_1 + G_2 + \cdots + G_n} \]  
(16)

\[ G_{sa} = \frac{P_1 + P_2 + \cdots + P_n}{G_1 + G_2 + \cdots + G_n} \]  
(17)

\( G_{sb} \) = bulk specific gravity for the combined aggregate;
\( G_{sa} \) = apparent specific gravity for the combined aggregate;
\( P_1, P_2, P_n \) = percentages by mass of aggregates 1, 2, n; and
\( G_1, G_2, G_n \) = bulk specific gravities (Equation 16) or apparent specific gravities (Equation 17) of aggregates 1, 2, n.

A1.2 Estimate the effective specific gravity of the combined aggregate in the aggregate trial blend using Equation 18:

\[ G_{se} = G_{sb} + 0.8 \left( G_{sa} - G_{sb} \right) \]  
(18)

where:
\( G_{se} \) = effective specific gravity of the combined aggregate;
\( G_{sb} \) = bulk specific gravity of the combined aggregate; and
\( G_{sa} \) = apparent specific gravity of the combined aggregate.

Note 21—The multiplier, 0.8, can be changed at the discretion of the designer. Absorptive aggregates may require values closer to 0.6 or 0.5.

Note 22—The Superpave mix design system includes a mixture conditioning step before the compaction of all specimens; this conditioning generally permits binder absorption to proceed to completion. Therefore, the effective specific gravity of Superpave mixtures will tend to be close to the apparent specific gravity in contrast to other design methods where the effective specific gravity generally will lie near the midpoint between the bulk and apparent specific gravities.
A1.3 Estimate the volume of binder absorbed into the aggregate, $V_{ba}$, using Equations 19 and 20:

$$V_{ba} = W_s \left( \frac{1}{G_{sb}} - \frac{1}{G_{se}} \right)$$  \hspace{1cm} (19)

where:

$$W_s, \text{ the mass of aggregate in } 1 \text{ cm}^3 \text{ of mix, g, is calculated as:}$$

$$W_s = \frac{P_b}{P_s} \frac{1 - V_a}{G_b + \frac{P_s}{G_{se}}}$$  \hspace{1cm} (20)

and where:

$P_s = \text{ mass percent of binder, in decimal equivalent, assumed to be 0.05;}$

$P_s = \text{ mass percent of aggregate, in decimal equivalent, assumed to be 0.95;}$

$G_b = \text{ specific gravity of the binder; and}$

$V_a = \text{ volume of air voids, assumed to be 0.04 cm}^3 \text{ in } 1 \text{ cm}^3 \text{ of mix.}$

**Note 23**—This estimate calculates the volume of binder absorbed into the aggregate, $V_{ba}$, and subsequently, the initial, trial binder content at a target air void content of 4.0 percent.

A1.4 Estimate the volume of effective binder using Equation 21:

$$V_{be} = 0.176 - \left[ 0.0675 \log(S_n) \right]$$  \hspace{1cm} (21)

where:

$V_{be} = \text{ volume of effective binder, cm}^3; \text{ and}$

$S_n = \text{ nominal maximum sieve size of the largest aggregate in the aggregate trial blend, mm.}$

**Note 24**—This regression Equation is derived from an empirical relationship between:

(1) VMA and $V_{be}$ when the air void content, $V_a$, is equal to 4.0 percent: $V_{be} = \text{VMA} - P_a = \text{VMA} - 4.0$; and (2) the relationship between VMA and the nominal maximum sieve size of the aggregate in MP 2. For WSDOT projects, see contract provisions.

A1.5 Calculate the estimated initial trial binder ($P_{bi}$) content for the aggregate trial blend gradation using Equation 22:

$$P_{bi} = 100 \times \left( \frac{G_b(V_{be} + V_{ba})}{(G_b(V_{be} + V_{ba}) + W_s)} \right)$$  \hspace{1cm} (22)

where:

$P_{bi} = \text{ estimated initial trial binder content, percent by weight of total mix.}$
INTRODUCTION
This test method explains how to locate and test for cyclic density. WSDOT field personnel are to systematically measure the locations where the new hot mix asphalt (HMA) pavement density may vary due to “spots, streaks” or visual pavement irregularities that may be related to temperature differentials or aggregate surface segregation. The described test method will identify density variations due to both causes.

1. GENERAL SCOPE
   a. Temperature differentials is defined as any area where the surrounding new HMA pavement temperature is 25°F or greater.
   b. Temperature differentials shall be determined when the new HMA pavement has been on the roadway for less than 1 minute, and no compaction has been applied.
   c. Aggregate segregation “Spots, streaks” or visual pavement irregularities is defined as areas of new HMA pavement that has a significantly different texture than the surrounding material.
   d. A systematic density reading shall be performed on locations where a temperature differential exists or where the HMA pavement shows spots, streaks, or has a significantly different texture after then finished rolling.
   e. Only systematic density readings located within the compaction lot should be marked and tested for density.
   f. Hot Mix Asphalt density measurements are made in accordance with WSDOT FOP for WAQTC TM-8 using a nuclear moisture density gauge in direct transmission mode.
   g. A density measurement shall be the result of a single four minute reading taken at the described location.
   h. Gauge-core correlation shall be in accordance with WSDOT SOP 730 is required for the systematic density testing.
   i. Normal Quality Assurance Testing will be performed throughout the entire job, as described in WSDOT SOP 729, in addition to any systematic density readings.

2. EQUIPMENT
   a. An approved infrared camera OR a handheld noncontact infrared thermometer (features for both should include continuous reading, minimum, maximum, and average readings, laser sighting, and a minimum distance to spot size ratio (D:S) of 30:1.
   b. Nuclear density gauge and standardizing block (reference standard).
   c. Tape measure.
   d. A can of spray paint for marking test locations.
   e. Required report form.
3. **GAUGE CALIBRATION**
   a. Shall be in accordance with WSDOT FOP for WAQTC TM-8. Follow the gauge calibration as outlined in FOP for WAQTC TM 8.

4. **TEMPERATURE CRITERIA**
   a. If the new HMA pavement temperature differentials are 25°F or greater then the surrounding new pavement, then a systematic density test is required.
   b. If the new HMA pavement temperature differentials are less than 25°F, then there is no need to perform testing unless an area shows signs of visual pavement irregularities, surface segregation or a significantly different texture.

5. **USE OF INFRARED CAMERA**
   a. View at least five consecutive truckloads of HMA (as described in steps b, c, d, e, and f) being placed and observe the location and temperature of any cool spots within the compaction lot. These observations should allow the operator to become familiar with the location and extent of the temperature differentials, if any, and if the temperature differentials are occurring in a cyclic manner.
   b. Viewing should occur from the side of the paved lane approximately 15 to 20 feet back from the paver looking toward the paver.
   c. The camera should be focused on the freshly placed HMA pavement prior to rolling. The camera should be adjusted to show the high and low temperatures.
   d. One truckload of HMA begins when the truck starts to dump into the paver or material transfer device and ends when another truck starts to dump.
   e. The “spot” function on the camera should be used to obtain the temperature of the cool area and the surrounding HMA to assess the temperature differential.
   f. Only temperature differentials located within the compaction lot should be marked for density testing.
   g. If the temperature differential is 25°F or more, locate the approximate center of the temperature differential area with the camera. The offset is from the center of the temperature differential area to the edge of the lane. Mark the location to be tested for systematic density by placing a paint mark at the edge of the lane corresponding to the center of the temperature differential. Record the HMA surface temperature, temperature differential, offset, and station as shown in Figure 1.
   h. If the temperature differential is less than 25°F, there is no need to mark the location unless an area within the paved lane has a significantly different texture. If testing is performed because of a significantly different textured area, locate the center of the affected area and mark the location as described in step g and as shown in Figure 1 with an (S) after the temperature differential.
6. **USE OF HANDHELD NONCONTACT INFRARED THERMOMETER**

   a. View at least five consecutive truckloads of HMA (as described in steps b, c, d, e, and f) being placed at varying offsets and observe the location and temperature of any cool spots within the compaction lot. These observations should allow the operator to become familiar with the location and extent of the temperature differentials, if any, and if the temperature differentials are occurring in a cyclic manner.

   b. Begin the longitudinal scan when a truck starts to dump into the paver or material transfer device and continue until the paver stops (discontinuous mix delivery) or until another truck starts to dump (continuous mix delivery).

   c. To perform the longitudinal scan, stand at the edge of the paving lane about 5 to 10 feet back from the paver. Scan the mat with the handheld noncontact thermometer continuously in a longitudinal manner by walking behind the paver in the direction of paving, staying the same distance away from the paver for one truckload of HMA. The offset for the longitudinal profile should be anywhere from 18 inches from the edge to no more than half the width of the paved lane. (The need to vary the longitudinal offset will be necessary to get an accurate representation of the whole mat.) Scanning temperatures for the other half of the paved lane should be performed from the other side.

   Note: Typically, temperature differentials or surface segregation at the beginning or end of a truckload can be captured with the longitudinal scan.

   d. Perform a transverse scan after completion of the longitudinal scan, making sure to scan the entire width of the paved lane excluding the outer 8 inches on each side. It should be performed approximately 5 to 10 feet behind the paver (to check for streaking of the mat).

   Note: Typically, streaking caused by temperature differentials or surface segregation will be captured by the transverse scan.

   e. The temperature scan can be stopped as soon as a temperature differential greater than 25°F has been located.

   f. Only temperature differentials located within the compaction lot should be marked for density testing.

   g. If the temperature differential is 25°F or more, locate the approximate center of the temperature differential area by scanning that specified location. The offset is from the center of the temperature differential area to the edge of the paved lane. Mark the location to be tested for systematic density by placing a paint mark at the edge of the lane corresponding to the center of the temperature differential. Record the HMA surface temperature, temperature differential, offset, and station as shown in Figure 1.

   h. If the temperature differential is less than 25°F, there is no need to mark the location unless an area within the paved lane has visual pavement irregularities, surface segregation or a significantly different texture. If testing is performed because of a significantly different textured area, locate the center of the affected area and mark the location as described in step g and as shown in Figure 1 with an (S) after the temperature differential.
7. SYSTEMATIC DENSITY PROCEDURE
   a. Testing shall be performed after the Contractor has finished compaction of the paved lane.
   b. Locate the mark (Figure 1) and record the information as listed.
   c. The probe of the gauge shall be placed at the offset listed and perform the testing according to WSDOT FOP for WAQTC TM 8 (direct transmission mode).
   d. Record the data on the Hot Mix Asphalt Concrete Pavement Compaction Report for Cyclic Density Form.

8. NUMBER AND LOCATION OF TEMPERATURE PROFILES AND SYSTEMATIC DENSITY TESTS
   a. If any temperature differentials were found in the initial assessment of paving operations (as described in 5a or 6a), the Engineer or Inspector shall take at least one temperature profile for every 5 trucks delivered to the paving operation.
   b. If the operation is not producing temperature differentials greater than 25°F in a cyclic pattern or the Engineer is not able to find 4 or more locations to be tested per compaction lot, the testing frequency can be reduced, but should be checked randomly throughout the day and the results recorded.
   c. If any significant equipment or weather changes occur, temperature profiles should be performed to determine if the new operation is capable of producing uniform HMA temperatures. If the paving machine in use is causing surface segregation, spotting or streaking that creates a finish that has a significantly different texture than the surrounding HMA, density tests should be performed in accordance with section 7 of this SOP.
   d. No temperature profiles shall be performed within the first or last 25 tons of production each day or within 25 feet of any transverse joint.
   e. Systematic density testing shall be performed on any location marked for testing.

Marking location of temperature differential.

Figure 1.
SAMPLING HOT MIX ASPHALT AFTER COMPACTION (OBTAINING CORES)

WSDOT SOP 734

1. Scope
   - This method describes the process for obtaining Hot Mix Asphalt test cores for Laboratory testing after compaction has been completed. Cores may range in size from 2 in. to 12 in.

2. SIGNIFICANCE AND USE
   - Samples obtained in accordance with the procedure given in this practice may be used for measuring pavement thickness, density, and acceptance testing.
   - When cores are used to determine nuclear gauge correlation, refer to WSDOT SOP 730.
   - When cores are used to determine pavement density, the Bulk Specific Gravity ($G_{mb}$) is determined according to WSDOT FOP for AASHTO T 166.

3. Apparatus
   - Core Drill Machine – A Core Drill Machine of sufficient horsepower and depth to minimize distortion of the compacted cores of Hot Mix Asphalt.
   - Core Bit – The cutting edge of the core drill bit shall be of hardened steel or other suitable material with diamond chips embedded in the metal cutting edge or as recommended by the core drill bit manufacturer. Typically the core drill bit should have an inside diameter of $4'' \pm 0.25''$ (100 mm $\pm$ 6 mm) or $6'' \pm 0.25''$ (150 mm $\pm$ 6 mm), these core bit dimensions are agency preferred alternatives. Suitable larger and smaller diameter core bit alternatives shall be employed as required by the agency.
   - Tools – Core layers may be separated using a saw or other suitable device which provides a clean smooth surface and does not damage the core.
   - Retrieval Device (Optional) – The retrieval device used for removing core samples from holes must preserve the integrity of the core. The device may be a steel rod of suitable length and with a diameter that will fit into the space between the core and the pavement material. There may be a 90 degree bend at the top to form a handle and a 90 degree bend at the bottom, approximately 2 in. (50 mm) long, forming a hook to assist in the retrieval of the core or other suitable device.

4. Safety
   - This standard does not purport to address all of the safety concerns, associated with its use. It is the responsibility of the user of this standard operating procedure to establish a pre activity safety plan prior to use.
   a. Test Site Location
      - The quantity of cores to be obtained shall be determined by the test procedure to be performed or agency requirements. Refer to WSDOT SOP 730 when taking correlation cores.
      - Determine the location of the core(s) as required by the agency.
      - For challenge cores, new random locations shall be determined according to WSDOT T 716 Hot Mix Asphalt density section.
b. Procedure

- For freshly placed Hot Mix Asphalt materials, the core shall be taken when the material has had sufficient amount of time to cool to prevent damage to the core.
- Pavement may be cooled to expedite the removal of the core by the following methods; water, ice water, ice, or dry ice or liquid nitrogen.
- Place the coring machine and core bit over the selected location.
- Keep the core bit perpendicular to the Hot Mix Asphalt surface during the coring process. Note: If any portion of the coring machine shifts during the operation, the core may break or distort.
- Constant downward pressure should be applied on the core bit. Failure to apply constant pressure, or too much pressure, may cause the bit to bind or distort the core.
- Continue the coring operation until the desired depth is achieved.
- If necessary, use a retrieval device to remove the core.
- Clearly identify the cores location and offset without causing damage (i.e., lumber crayon or grease pencil).
  
  Note: If the core is damaged to a point that it cannot be used for its intended purpose, a new core shall be obtained within 6 in. of the original location.

6. Filling Core Holes

- When necessary, the hole made from the coring operation shall be filled with a material that will not separate from the surrounding material. If a Hot Mix Asphalt is available and used, it shall be compacted into the hole. A fast set grout product may be used in lieu of a Hot Mix Asphalt. A black dye can be used to color the grout on wearing lifts.

7. Transporting Cores

- Transport cores in a suitable container(s) that prevents damage from jarring, rolling, hitting together, and/or impact with any object.
- Prevent cores from freezing or excessive heat above 130° F (54° C), during transport.
  
  Note #1: In extreme ambient temperature conditions, cores should be placed in water during transport.
- If the core is damaged in transport to a point it can not be utilized for its intended purpose the core will not be used.

8. Separate the Layers

- When necessary, separate the lifts or layers of pavement courses by using a water cooled saw to cut the core on the designated lift line or separate by other suitable methods that will not damage the lifts or layers to be tested.
  
  Note #2: Lift lines are often more visible by rolling the core on a flat surface and/or surface drying the core.

9. Length Determination

Measure the thickness of the designated lift to the nearest 0.01’ or ⅛” according to WSDOT Test Method 720.
10. Report

- Core information shall be reported on standard agency forms and should include the following information.
  - The date the cores were obtained
  - Paving date
  - Contract number
  - Project title
  - Location of test
  - The lift being evaluated
  - Type of material being evaluated
  - Mix Design Lab Number
  - Average thickness of each core (to the nearest 0.01’ or ⅛ “)
  - Average Theoretical Maximum Density
1. GENERAL SCOPE
   a. In addition to the 5 random Quality Assurance (QA) density tests performed per compaction lot, one density test at each longitudinal joint will be performed on a confined or unconfined edge.

2. LONGITUDINAL JOINT TESTING
   a. The companion longitudinal joint test will be taken at the same station as the test for the third sublot.
   b. Use the guide and scraper plate as a template and drill a hole to a depth of at least ¼ in. (7 mm) deeper than the measurement depth required for the gauge six inches from the edge of the longitudinal joint, in this case disregard the following portion of WSDOT FOP for WAQTC TM-8 “Test Site Location, item c., or less than 8 in. (450 mm) from a vertical pavement edge,” making sure the gauge will sit flush with the hot-mix asphalt (HMA). See Figure 1.

3. NUMBER OF LONGITUDINAL JOINT TESTS
   a. Two readings (one at each longitudinal joint) will be taken within each compaction lot at the same station location as the third sublot.

4. CALCULATION OF RESULTS
   a. Calculate the Longitudinal Joint Density as defined in step 3 b in WSDOT SOP 729.

5. REPORT
   a. Report the test results on the ‘Longitudinal Joint Density Record Sheet that is an Appendix in this procedure.

   Note: Lot Number corresponds to the lot where the set of longitudinal joint readings were taken. The station corresponds to the station within the lot (i.e. third sublot) where the set of longitudinal joint readings were taken.

![Location of Longitudinal Joint Density Tests No Gauge Rotation.](image)}
WSDOT Test Method No. 802

Method of Test for Flexural Strength of Concrete (Using Simple Beam With Center-Point Loading)

1. SCOPE
   a. This method is similar to AASHTO T 177 and covers the procedure for determining the flexural strength of concrete by the use of a simple beam with center-point loading.

2. APPARATUS
   a. The center-point loading method shall be used in the laboratory. The testing machine shall conform to the requirements of Sections 15, 16, and 17 of the Methods of Verification of Testing Machines (AASHTO T 67). In the field, a manually operated calibrated jack shall be used in conjunction with the field testing machine supplied by the Regional Materials Engineer. The apparatus shall incorporate the following requirements. The load shall be applied at the center point of the span, normal to the loaded surface of the beam, employing bearing blocks designed to ensure that forces applied to the beam will be vertical only and applied without eccentricity. The direction of the reactions shall be parallel to the direction of the applied load at all times during the test. The load shall be applied at a uniform rate and in such a manner as to avoid shock. The edges of the load-applying block and of the supports shall not depart from a plane by more than .002 in. (0.051 mm).
   b. Caliper — A 12-in. (1300 mm) long caliper accurate to 0.01 in. (0.25 mm).

![Diagram](image)

Figure 1: Diagrammatic View of Apparatus for Flexure Test of Concrete by Center-point Loading Method

3. TEST SPECIMEN
   a. As nearly as practicable, the test specimen, as tested, shall have a span three times its depth. The test specimen shall be formed and stored as prescribed in WSDOT Test Method No. 808.
4. PROCEDURE
   a. Turn the specimen on its side with respect to its position when molded, and center it on the
      supporting bearing blocks. The load-applying block shall be brought in contact with the upper
      surface at the center line between the supports.
   b. Bring load applying block in full contact with the beam surface by applying a 100 lbs. (3.1 N)
      preload. Check to ensure that the beam is in uniform contact with the bearing blocks and the
      load applying block.
   c. If load is applied with a hand pump, load the beam by applying the load at a rate of one full
      pump stroke per second. When the applied load is about 4,000 lbs. (125 N), reduce the full
      pump stroke to about a \( \frac{1}{2} \)-pump stroke and maintain the one second stroke rate.
      
      Rate of load application for screw power machines, with the moving head operating at 0.05 in. 
      (1.3 mm) per minute when the machine is running idle, is acceptable.

5. MEASUREMENT OF SPECIMENS
   a. Determine the beam dimensions, width (b) and depth (d), by averaging two measurements for
      width and two measurements for depth. The measurements shall be taken at the failure plane
      to an accuracy of 0.05 in. (1.3 mm).

6. CALCULATION
   a. The modulus of rupture is calculated as follows:

      \[ R = \frac{3Pl}{2bd^2} \]

      where:

      - \( R \) = Modulus of rupture in psi or MPa
      - \( P \) = Maximum applied load indicated by the testing machine in lb•f or N
      - \( l \) = Span length in inches or mm
      - \( b \) = Average width of specimen in inches or mm
      - \( d \) = Average depth of specimen in inches or mm

7. REPORT
   a. The report shall include the following:
      
      (1) Identification number,
      (2) Average width,
      (3) Average depth,
      (4) Span length in inches or mm,
      (5) Maximum applied load in lb•f or N,
      (6) Modulus of rupture calculated to the nearest 5 psi (0.03 MPa),
      (7) Defects in specimen, and
      (8) Age of specimen.
   b. All test results will be reported on DOT Form 350-042.
Performance Exam Checklist

Method of Test for Flexural Strength of Concrete (Using Simple Beam With Center-Point Loading)
WSDOT Test Method T 802

Participant Name ___________________________ Exam Date ________________

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Copy of current procedure available at test site?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. In the field, Jack properly calibrated?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Beam turned on its side with respect to its position when molded, and centered on the supporting bearing blocks?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Load applying block brought into contact with the beam at the center line between the supports?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. 100 lbs. (3.1 N) preload applied and the beam then checked to ensure uniform contact with the bearing blocks and load applying block?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Load applied to the beam at the proper uniform rate?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Equipment

1. Where required are calibration/verifications tags present on equipment used in this procedure? |     |    |
2. All equipment functions according to the requirements of this procedure?             |     |    |

Signature of Examiner ____________________________________________________________

Comments: First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐
Rebound Hammer Determination of Compressive Strength of Hardened Concrete

1. Scope

1.1 This test method covers the determination of a rebound number of hardened concrete using a spring-driven steel hammer.

1.2 The values stated in SI inch-pound units are to be regarded as the standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 ASTM Standards:

C 125 Terminology Relating to Concrete and Concrete Aggregates

C 670 Practice for Preparing Precision and Bias Statements for Test Methods for Construction Materials

E 18 Test Methods for Rockwell and Rockwell Superficial Hardness of Metallic Materials

3. Terminology

3.1 Definitions:

For definitions of terms used in this test method, refer to Terminology C 125.

4. Summary of Test Method

4.1 A steel hammer impacts, with a predetermined amount of energy, a steel plunger in contact with a surface of concrete, and the distance that the hammer rebounds is measured.

5. Significance and Use

5.1 This test method is applicable to assess the in-place uniformity of concrete, to delineate regions in a structure of poor quality or deteriorated concrete, and to estimate in-place strength development.

5.2 To use this test method to estimate strength requires establishing a relationship between strength and rebound number. The relationship shall be established for a given concrete mixture and given apparatus. The relationship shall be established over the range of concrete strength that is of interest. To estimate strength during construction, establish the relationship by performing rebound number tests on molded specimens and measuring the strength of the same or companion molded specimens. To estimate strength in an existing structure, establish the relationship by correlating rebound numbers measured on the structure with the strengths of cores taken from corresponding locations. See ACI 228.R4 for additional information on developing the relationship and on using the relationship to estimate in-place strength.

5.3 For a given concrete mixture, the rebound number is affected by factors such as moisture content of the test surface, the method used to obtain the test surface (type of form material or type of finishing), and the depth of carbonation. These factors need to be considered in preparing the strength relationship and interpreting test results.
5.4 Different hammers of the same nominal design may give rebound numbers differing from 1 to 3 units. Therefore, tests should be made with the same hammer in order to compare results. If more than one hammer is to be used, perform tests on a range of typical concrete surfaces so as to determine the magnitude of the differences to be expected.

5.5 This test method is not intended as the basis for acceptance or rejection of concrete because of the inherent uncertainty in the estimated strength.

6. Apparatus

6.1 *Rebound Hammer*, consisting of a spring-loaded steel hammer that when released strikes a steel plunger in contact with the concrete surface. The spring-loaded hammer must travel with a consistent and reproducible velocity. The rebound distance of the steel hammer from the steel plunger is measured on a linear scale attached to the frame of the instrument.

**NOTE 1**—Several types and sizes of rebound hammers are commercially available to accommodate testing of various sizes and types of concrete construction.

6.2 *Abrasive Stone*, consisting of medium-grain texture silicon carbide or equivalent material.

6.3 *Test Anvil*, approximately 150-mm (6-in.) diameter by 150-mm (6-in.) high cylinder made of tool steel with an impact area hardened to 66 ± 2 HRC as measured by Test Methods E 18. An instrument guide is provided to center the rebound hammer over the impact area and keep the instrument perpendicular to the surface.

6.4 *Verification*—Rebound hammers shall be serviced and verified annually and whenever there is reason to question their proper operation. Verify the functional operation of a rebound hammer using the test anvil described in 6.3. During verification, support the test anvil on a bare concrete floor or slab. The manufacturer shall report the rebound number to be obtained by a properly operating instrument when tested on an anvil of specified hardness.

**NOTE 2**—Typically, a rebound hammer will result in a rebound number of 80 ± 2 when tested on the anvil described in 6.3. The test anvil needs to be supported on a rigid base to obtain reliable rebound numbers. Verification on the test anvil does not guarantee that the hammer will yield repeatable data at other points on the scale. The hammer can be verified at lower rebound numbers by using blocks of polished stone having uniform hardness. Some users compare several hammers on concrete or stone surfaces encompassing the usual range of rebound numbers encountered in the field.

7. Test Area and Interferences

7.1 *Selection of Test Surface*—Concrete members to be tested shall be at least 100 mm (4 in.) thick and fixed within a structure. Smaller specimens must be rigidly supported. Avoid areas exhibiting honeycombing, scaling, or high porosity. Do not compare test results if the form material against which the concrete was placed is not similar (see Note 3). Troweled surfaces generally exhibit higher rebound numbers than screeded or formed finishes. If possible, test structural slabs from the underside to avoid finished surfaces.

7.2 *Preparation of Test Surface*—A test area shall be at least 150 mm (6 in.) in diameter. Heavily textured, soft, or surfaces with loose mortar shall be ground flat with the abrasive stone described in 6.2. Smooth-formed or troweled surfaces do not have to be ground prior to testing (see Note 3). Do not compare results from ground and unground surfaces.
NOTE 3—Where formed surfaces were ground, increases in rebound number of 2.1 for plywood formed surfaces and 0.4 for high-density plywood formed surfaces have been noted.5 Dry concrete surfaces give higher rebound numbers than wet surfaces. The presence of surface carbonation can also result in higher rebound numbers.6 The effects of drying and surface carbonation can be reduced by thoroughly wetting the surface for 24 h prior to testing. In cases of a thick layer of carbonated concrete, it may be necessary to remove the carbonated layer in the test area, using a power grinder, to obtain rebound numbers that are representative of the interior concrete. Data are not available on the relationship between rebound number and thickness of carbonated concrete. The user must exercise professional judgment when testing carbonated concrete.

7.3 Do not test frozen concrete.

NOTE 4—Moist concrete at 0 °C (32 °F) or less may exhibit high rebound values. Concrete should be tested only after it has thawed. The temperatures of the rebound hammer itself may affect the rebound number. Rebound hammers at -18 °C (0 °F) may exhibit rebound numbers reduced by as much as 2 or 37.

7.4 For readings to be compared, the direction of impact, horizontal, downward, upward, or at another angle, must be the same or established correction factors shall be applied to the readings.

7.5 Do not conduct tests directly over reinforcing bars with cover less than 0.75 in. [20 mm].

NOTE 5—The location of reinforcement may be established using reinforcement locators or metal detectors. Follow the manufacturer’s instructions for proper operation of such devices.

8. Procedure

8.1 Hold the instrument firmly so that the plunger is perpendicular to the test surface. Gradually push the instrument toward the test surface until the hammer impacts. After impact, maintain pressure on the instrument and, if necessary, depress the button on the side of the instrument to lock the plunger in its retracted position. Read the rebound number on the scale to the nearest whole number and record the rebound number. Take ten readings from each test area. No two impact tests shall be closer together than 25 mm (1 in.). Examine the impression made on the surface after impact, and if the impact crushes or breaks through a near-surface air void disregard the reading and take another reading.

9. Calculation

9.1 Discard readings differing from the average of 10 readings by more than 6 units and determine the average of the remaining readings. If more than 2 readings differ from the average by 6 units, discard the entire set of readings and determine rebound numbers at 10 new locations within the test area.
10. Report

10.1 Report the following information for each test area:

10.1.1 Date and time of testing,

10.1.2 Identification of location tested in the concrete construction and the type and size of member tested,

10.1.2.1 Description of the concrete mixture proportions including type of coarse aggregates if known, and

10.1.2.2 Design strength of concrete tested.

10.1.3 Description of the test area including:

10.1.3.1 Surface characteristics (trowelled, screeded) of area,

10.1.3.2 If surface was ground and depth of grinding, 5 Gaynor, R. D., “In-Place Strength of Concrete—A Comparison of

10.1.3.3 Type of form material used for test area,

10.1.3.4 Curing conditions of test area,

10.1.3.5 Type of exposure to the environment,

10.1.4 Hammer identification and serial number,

10.1.4.1 Air temperature at the time of testing,

10.1.4.2 Orientation of hammer during test,

10.1.5 Average rebound number for test area, and

10.1.5.1 Remarks regarding discarded readings of test data or any unusual conditions.

11. Precision and Bias

See ASTM C 805 Precision and Bias

11.1 Precision—The single-specimen, single-operator, machine, day standard deviation is 2.5 units (1σ) as defined in Practice C 670. Therefore, the range of ten readings should not exceed 12.

11.2 Bias—The bias of this test method cannot be evaluated since the rebound number can only be determined in terms of this test method.

12. Keywords

12.1 concrete; in-place strength; nondestructive testing; rebound hammer; rebound number
Performance Exam Checklist

Rebound Hammer Determination of Compressive Strength of Hardened Concrete
FOP For ASTM C 805

Participant Name ________________________________ Exam Date ________________

**Procedure Element**

1. Copy of current procedure available at test site? | Yes | No
2. Hammer properly serviced and calibrated or verified? | Yes | No
3. Test location properly prepared? | Yes | No
4. Test location meets minimum size requirement? | Yes | No
5. Ten acceptable readings taken in each test area? | Yes | No
6. Readings properly spaced in test area? | Yes | No
7. Test readings properly converted to estimated strength? | Yes | No
8. Test information properly recorded? | Yes | No
9. All calculations performed correctly? | Yes | No

**Equipment**

10. Where required are calibration/verifications tags present on equipment used in this procedure? | Yes | No
11. All equipment functions according to the requirements of this procedure? | Yes | No

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

**Comments:**

_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
_________________________________________________________________
WSDOT Test Method T 807

Method of Operation of California Profilograph and Evaluation of Profiles

1. SCOPE
   a. The operation of the California Profilograph, the procedure used for determining the Profile Index from profilograms of pavements made with the Profilograph, and the procedure used to locate individual high points in excess of 0.3 in. (7.62 mm) are described in Parts I, II, and III, respectively, in this test method.

   Part I
   Operation of the California Profilograph

2. PROCEDURE
   a. Equipment
      
      The California Profilograph consists of a frame 25 LF (7.62 m) long supported upon multiple wheels at either end arranged in a staggered pattern such that no two wheels cross the same bump at the same time. The profile is recorded from the vertical movement of a wheel attached to the frame at midpoint and is in reference to the mean elevation of the 12 points of contact with the road surface established by the support wheels (see Figure 1). The profilogram is recorded on a scale of 1 in. = 25 LF (25 mm = 7620 mm) longitudinally and 25 mm = 25 mm (or full scale) vertically. Motive power is provided manually.

   b. Operation
      
      The instructions for assembling the Profilograph are contained in a booklet accompanying each unit. Particular attention should be paid to the listed precautions.

      In operation, the Profilograph should be moved at a speed no greater than a walk so as to eliminate as much bounce as possible. Too high a speed will result in a profilogram that is difficult to evaluate.

      Calibration of the Profilograph should be checked periodically. The horizontal scale can be checked by running a known distance and scaling the result on the profilogram. If the scale is off, the profile wheel should be changed to one of a proper diameter. The vertical scale is checked by putting a board of known thickness under the profile wheel and again scaling the result on the profilogram. If the scale is off, the cause of the incorrect height should be determined and corrected.
Figure 1
3. **PROCEDURE**

a. **Equipment**

To determine the Profile Index, use a plastic scale 1.70 LF (43.18 mm) wide and 1.70 in. (536.45 mm) long representing a pavement length of 528 LF (161 m) at a scale of 25 mm = 7.62 m a plastic scale for the Profilograph may be obtained by the districts from the OSC Materials Laboratory. Near the center of the scale is an opaque band 0.2 in. (5.1 mm) wide extending the entire length of 21.12 in. (536.45 mm). On either side of this band are scribed lines 0.1 in. (2.54 mm) apart, parallel to the opaque band. These lines serve as a convenient scale to measure deviations or excursions of the graph above or below the blanking band. These are called “scallops.”

b. **Method of Counting**

Place the plastic scale over the profile in such a way as to “blank out” as much of the profile as possible. When this is done, scallops above and below the blanking band usually will be approximately balanced. See Figure 2.

The profile trace will move from a generally horizontal position when going around super-elevated curves making it impossible to blank out the central portion of the trace without shifting the scale. When such a condition occurs, the profile should be broken into short sections and the blanking band repositioned on each section while counting as shown in the upper part of Figure 3.

Starting at the right end of the scale, measure and total the height of all the scallops appearing both above and below the blanking band, measuring each scallop to the nearest 0.05 in. (1.27 mm) (half a tenth). Write this total on the profile sheet near the left end of the scale together with a small mark to align the scale when moving to the next section. Short portions of the profile line may be visible outside the blanking band but unless they project 0.03 in. (0.762 mm) or more and extend longitudinally for 2 LF (610 mm) (0.8 in. (2 mm) on the profilogram) or more, they are not included in the count. (See Figure 2 for illustration of these special conditions.)

When scallops occurring in the first 0.1 mile (161 m) are totaled, slide the scale to the left, aligning the right end of the scale with the small mark previously made, and proceed with the counting in the same manner. The last section counted may or may not be an even 0.1 mile (161 m). If not, its length should be scaled to determine its length in kilometers. An example follows:

<table>
<thead>
<tr>
<th>Section length, miles (km)</th>
<th>Counts, tenth of an in.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.10 (0.16)</td>
<td>5.0</td>
</tr>
<tr>
<td>0.10 (0.16)</td>
<td>4.0</td>
</tr>
<tr>
<td>0.10 (0.16)</td>
<td>3.5</td>
</tr>
<tr>
<td>400 ft. = 0.076</td>
<td>2.0</td>
</tr>
<tr>
<td>Total</td>
<td>0.376</td>
</tr>
<tr>
<td></td>
<td>14.5</td>
</tr>
</tbody>
</table>

The Profile Index is determined as “inches per mile in excess of the 0.2 in. (5.1 mm) blanking band” but is simply called the Profile Index. The procedure for converting counts of Profile Index is as follows:
Using the figures from the above example:

Length = 0.376 mi. (0.605 km), total count = 14.5 tenths of an inch

Profile Index = \( \frac{1 \text{ mi. (1 km)}}{\text{length of profiles in miles (km)}} \times \text{a total count in mm} \)

\[
\text{Pr I} = \frac{1 \text{ mi. (1 km)}}{0.376 \text{ mi. (0.605 km)}} \times 1.45 = 3.9
\]

(Note that the formula uses the count in inches rather than tenths of a inch and is obtained by dividing the count by ten.)

The Profile Index is thus determined for the profile of any line called for in the specifications.

To determine the daily profile index to check the Contractors methods and procedures, profile indexes may be averaged for two or more profiles of the same section of road if the profiles are the same length.

Example:

<table>
<thead>
<tr>
<th>Section length, mi. (km)</th>
<th>Left wheel track</th>
<th>Right wheel track</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.26 (0.10)</td>
<td>5.0</td>
<td>4.5</td>
</tr>
<tr>
<td>0.26 (0.10)</td>
<td>4.0</td>
<td>5.0</td>
</tr>
<tr>
<td>0.26 (0.10)</td>
<td>3.5</td>
<td>3.0</td>
</tr>
</tbody>
</table>

\[
\text{400 LF = } \begin{array}{c}
0/0 \text{ mi. (0.076)} \\
\text{Total}
\end{array}
\begin{array}{c}
2.0 \\
14.5
\end{array}
\begin{array}{c}
1.5 \\
14.0
\end{array}
\]

\[
\text{PrI (by formula)} = \frac{39 + 3.7}{2} = 3.9
\]

\[
\text{Averages } = \frac{3.9 + 3.7}{2} = 3.8
\]

The specifications state which profiles to use when computing the average Profile Index for control of construction operations.

c. Limitations of Count in 0.1 mi. (0.1 km) Sections

When the specifications limit the amount of roughness in “any one-tenth mile section,” the scale is moved along the profile and counts made at various locations to find those sections if any, that do not conform to specifications. The limits are then noted on the profile and can be later located on the pavement preparatory to grinding.

d. Limits of Counts — Joints

When counting profiles, a day’s paving is considered to include the last portion of the previous day’s work, which includes the daily joint. The last 15 to 30 LF (4.57 to 9.14 m) of a day’s paving cannot usually be obtained until the following day. In general, the paving contractor is responsible for the smoothness of joints if he places the concrete pavement on both sides of the joint. On the other hand, the contractor is responsible only for the pavement placed by him if the work abuts a bridge or a pavement placed under another contract. Profilograph readings when approaching such joints should be taken in conformance with current specifications.
Figure 2
METHOD OF COUNTING WHEN POSITION OF PROFILE-shifts as it may
WHEN Rounding SHORT RADIUS CURVES WITH SUPERELEVATION

Incorrect position of blanking band

Blanking band shifted to accommodate lowering of profile

METHOD OF PLACING TEMPLATE WHEN LOCATING BUMPS TO BE REDUCED

Figure 3

BUMP TEMPLATE

Baseline approx. 25 feet

Baseline less than 25 feet

Height of peak is less than 0.3"

Baseline more than 25'
Part III

Determination of High Points in Excess of 0.3 in. (7.62 mm)

4. PROCEDURE

a. Equipment

Use a plastic template having a line 1 in. (25.4 mm) long scribed on one face with a small hole or scribed mark at either end, and a slot 0.3 in. (7.62 m) from and parallel to the scribed line. See Figure 3. (The 1 in. (25.4 mm) line corresponds to a horizontal distance of 0.3 in. (7.62 m) on the horizontal scale of the profilogram.) The plastic template may be obtained from the Olympia Service Center Materials Laboratory.

b. Locating High Points in Excess of 0.3 in. (7.62 m).

At each prominent peak or high point on the profile trace, place the template so that the small holes or scribe marks at each end of the scribed line intersect the profile trace to form a chord across the base of the peak or indicated bump. The line on the template need not be horizontal. With a sharp pencil, draw a line using the narrow slot in the template as a guide. Any portion of the trace extending above this line will indicate the approximate length and height of the deviation in excess of 0.3 in. (7.62 m).

There may be instances where the distance between easily recognizable low points is less than 1 in. (25.4 mm) (30 LF (7.62 m)). In such cases, a shorter chord length shall be used in making the scribed line on the template tangent to the trace at the low points. It is the intent, however, of this requirement that the baseline for measuring the height of bumps will be as near 30 LF (7.62 m) (1 in. (25.4 mm)) as possible, but in no case to exceed this value. When the distance between prominent low points is greater than 0.3 in. (7.62 m) (1 in. (25.4 mm)), make the ends of the scribed line intersect the profile trace when the template is in a nearly horizontal position. A few examples of the procedure are shown in the lower portion of Figure 3.
WSDOT Test Method T 808

Method for Making Flexural Test Beams

1. SCOPE
   a. This method covers the procedures for molding and curing Portland cement concrete flexural test beams.

2. EQUIPMENT
   a. Test beam molds, 6 in × 6 in. × 21 ± ½ in. (150 mm × 150 mm × 550 ± 13 mm) or 8 in. × 8 in. × 26 ± ½ in. (200 mm × 200 mm × 670 ± 13 mm).
   b. Vibrator, capable of 7,000 vibrations per minute with a diameter not less than ¾ in. (19.0 mm) or greater than 1 ½ in. (38.1 mm).
   c. Tamping Rod — The tamping rod is a round, straight steel rod ¾ in. (16.0 mm) diameter and approximately 24 in. (610 mm) long, having the tamping end rounded to a ¾ in. (16.0 mm) diameter hemispherical tip.
   d. Mallet — A mallet with a rubber or rawhide head weighing 1.25 ± 0.50 lb. (0.57 ± 0.23 kg).
   e. Assorted tools such as scoops, shovels, etc.

3. PROCEDURE
   a. For laboratory made beam specimens, mix sufficient concrete to make all the required specimens from one batch. Each beam specimen requires approximately .45 yd.³ (0.015 m³) of concrete.
      For field-made beam specimens, the concrete sample is obtained in accordance with WSDOT Test Method No. 803, Method of Sampling Fresh Concrete. Making of the beam specimens shall begin within 15 minutes of remixing the sample.
   b. Mold specimens as near as practicable to the place where they are to be stored during the first 24 hours.
   c. Assemble the molds on a rigid surface free from vibration and other disturbances. Remix the concrete to a uniform appearance. When the method of consolidation is by internal vibrators, the mold is filled in a single layer. Make sure that each shovel or scoop of concrete is representative of the batch. When the method of consolidation is by rodding, the mold is filled in two layers with each layer being rodded one time for each 2 in.² (1290 mm²) of surface area. The rodding should be distributed evenly over the entire surface. On the succeeding layers, the rod should not penetrate the previous layer more than ½ in. (13 mm). After each layer is rodded, tap the outsides of the mold lightly 10 to 15 times with a mallet.
   d. Insert the vibrator at intervals not to exceed 6 in. (150 mm) along the centerline of the long dimension of the beam. For specimens wider than 6 in. (150 mm), use alternating insertions along two lines at least 2 in. (50 mm) away from the sides of the mold. Withdraw the vibrator so that no air voids are left in the concrete. Then tap the mold lightly 10-15 times with mallet.
   e. Finish the surface of the concrete by striking off with a straightedge. Use the minimum amount of manipulation necessary to leave a flat surface that has no depressions or projections larger than ½ in. (3.2 mm) and is level with the sides of the mold.
f. The top surface of the laboratory-made specimen shall be covered with a saturated towel and a plastic sheet to prevent moisture loss from the concrete.

For the field made specimen, the top surface of the beam shall be sprayed with the same curing compound as is used for the pavement and covered with a plastic tarpaulin.

4. STORAGE AND HANDLING

The method of storing and handling the beam specimen depends on the purpose for which the beam is intended. Two methods are provided as follows:

a. Laboratory Method — Beam for determining the acceptability of a contractor-provided paving mix.

Cover the beam to prevent moisture loss and allow beam to remain undisturbed for an initial cure period of 24 ± 4 hours at a temperature of 60° to 80°F (16° to 27°C). After the initial cure period, the beam will be removed from the mold and within 30 minutes stored in saturated limewater at 73.4° ± 3°F (23° ± 2°C) for a minimum of 20 hours prior to testing. Surface drying of the beam between removal from the limewater and completion of testing shall be prevented. Relatively small amounts of drying of the test beam surfaces induces tensile stress in the extreme fibers that will markedly reduce the indicated flexural strength.

b. Field Method — Beam for determining the flexural strength of the inplace pavement.

After applying the curing compound to the top surface, cover the beam specimen with white reflective sheeting and allow beams to remain undisturbed for an initial cure period of 24 ± 4 hours at ambient conditions. After the initial cure period, remove the specimen from the mold and cure the specimen either by:

(1) Burying the specimen in wet sand making sure that the specimen is never allowed to become surface dry. Temperature of the sand should be similar to the concrete pavement temperature, or

(2) Wrap the beam in a saturated towel, place in a plastic bag, and seal the opening. The plastic should be at least 4 mils thick. Leave the specimen on the pavement in the vicinity where it was molded until time to test. Take specimen to the testing location and store in lime water at 73.4° ± 5°F (23° ± 2.8°C) for 24 ± 4 hours immediately before time of testing to ensure uniform moisture condition from specimen to specimen.

Note: The beam specimen must be kept in a surface moist condition or wet environment for the entire time in storage and testing. Even minor amounts of surface drying of the specimen induces extreme fiber stresses which can markedly reduce the flexural strength.

5. TESTING

a. Beam specimens are tested for flexural strength in accordance with WSDOT Test Method No. 802.
Performance Exam Checklist

Method for Making Flexural Test Beams

WSDOT T 808

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Copy of current procedure available at test site?</td>
<td></td>
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</tr>
<tr>
<td>2. Making of test specimens begins within 15 minutes for sampling?</td>
<td></td>
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<tr>
<td>3. Assemble of molds on a rigid surface free from vibration and other disturbances?</td>
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<tr>
<td>4. Concrete remixed to a uniform appearance?</td>
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<tr>
<td>5. When method of concrete consolidation is by rodding:</td>
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<tr>
<td>a. Mold filled in two layers?</td>
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<tr>
<td>b. Each layer rodded one time for each 2 in.² (1290 mm) of mold surface area?</td>
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<tr>
<td>c. Rodding, evenly distributed over the entire surface area?</td>
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<tr>
<td>d. After rodding each layer, mold tapped lightly 10-15 times with mallet?</td>
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<tr>
<td>6. When method of concrete consolidation is by internal vibrators:</td>
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<tr>
<td>a. Mold filled in a single layer?</td>
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<tr>
<td>b. Vibrator inserted at intervals not to exceed 6 in. (150 mm) along the centerline of the long dimension?</td>
<td></td>
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<tr>
<td>c. For molds wider than 6 in. (150 mm), vibrator inserted along two alternating lines at least 2 in. (50 mm) away from sides of mold?</td>
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<tr>
<td>d. Mold tapped lightly 10-15 times with mallet?</td>
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<tr>
<td>7. Top of mold properly finished?</td>
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<td></td>
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<tr>
<td>8. Top of mold properly treated to prevent moisture loss?</td>
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</tbody>
</table>

Equipment

<table>
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<tr>
<th>Equipment Element</th>
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<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Where required are calibration/verifications tags present on equipment used in this procedure?</td>
<td></td>
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<tr>
<td>2. All equipment functions according to the requirements of this procedure?</td>
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</tbody>
</table>

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________
1. SCOPE
   a. This method of test is a modification of AASHTO T 85 and is intended for use in determining the density of Portland cement concrete pavement cores. The object of this test is to determine the inplace density of the concrete as it exists, a minimum of 24 hours after placement. This density is the value desired for comparison to the density of freshly mixed portland cement as determined by AASHTO T 121 when specific gravity values on absolute terms are required, AASHTO T 85 shall be used.

2. EQUIPMENT
   a. Balance — A balance having capacity of 5 kg or more and sensitive to 1 g or less.
   b. Wire Basket — A wire basket of appropriate size, constructed of wire mesh.
   c. Container — A container suitable for immersing the wire basket in water, and an apparatus for suspending the wire basket from the center of the scale pan of the balance. Maintain a constant water level when weighing under water.
   d. Absorbent towels.
   e. Temperature Measuring Device — The temperature measuring device shall be verified and readable to 1°F (0.5°C). Thermometers having a range of 0 to 120°F (-18 to 49°C) are satisfactory. Other thermometers of the required accuracy, including the metal immersion type, conforming to ASTM E 1 are acceptable.

3. PROCEDURE
   a. Density determinations are made as soon as practicable after coring and with a minimum change in moisture content from the condition as taken. Where on-site determination is not practicable within one hour, cores are stored in airtight plastic bags or completely immersed in water until weighed. Core densities shall be determined within 4 hours after coring.

<table>
<thead>
<tr>
<th>Temperature °F</th>
<th>Pounds perCubic Foot</th>
</tr>
</thead>
<tbody>
<tr>
<td>65</td>
<td>62.336</td>
</tr>
<tr>
<td>66</td>
<td>62.329</td>
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<tr>
<td>67</td>
<td>62.322</td>
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<td>75</td>
<td>62.261</td>
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<tr>
<td>80</td>
<td>62.216</td>
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</table>
b. Wash thoroughly to remove dust or other coatings from the surface of the core. Place the sample in the wire basket and determine its mass in water. Determine this and all subsequent weights to the nearest gram. Determine the temperature of the water to the nearest degree.

c. Remove the sample from the water and roll it in a large absorbent cloth until all visible films of water are removed, although the surfaces still appear to be damp. Take care to avoid evaporation from aggregate pores during the operation of surface drying. Obtain the weight of the sample in the surface dry condition.

4. CALCULATION
a. Calculate the density as follows:

\[ \text{Density (surface-dry basis)} = \frac{A}{A - B} \times d_w \]

where:
A = mass in grams of the surface-dry sample in air
B = mass in grams of the sample in water
d, = density of the water at the test temperature (see Table 1)

Calculate the density to the nearest 0.1 lb. per ft.\(^3\) (1 kg per m\(^3\)).

5. REPRODUCIBILITY OF RESULTS
a. Duplicate determinations should check to within 0.1 lb. per ft.\(^3\) (3 kg per m\(^3\)).

6. REPORTS
a. The test results will be reported on the appropriate test data sheet.
Performance Exam Checklist

Method of Test for Determination of the Density of PCC Pavement Cores
(WSDOT TM 810)

<table>
<thead>
<tr>
<th>Procedure Element</th>
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<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
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</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and</td>
<td></td>
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<tr>
<td>if required, has the current calibration/verification tags present?</td>
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<tr>
<td>3. Density determined approximately 24 hours after concrete placement?</td>
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<tr>
<td>4. Core’s moisture content preserved in bags or by immersion?</td>
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<tr>
<td>5. Density determined within 4 hours of coring?</td>
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<tr>
<td>6. Core washed thoroughly?</td>
<td></td>
<td></td>
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<tr>
<td>7. Weight in water determined to nearest gram?</td>
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<td></td>
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<tr>
<td>8. Temperature of water determined to nearest degree</td>
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<td></td>
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<tr>
<td>9. Core rolled on absorbent towel removing visible films of water but</td>
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<tr>
<td>still appearing damp?</td>
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</tr>
<tr>
<td>10. Core not over-dried or allowed to evaporate?</td>
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</tr>
<tr>
<td>11. Weight of surface-dry core determined to nearest gram?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12. All calculations performed correctly?</td>
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<td></td>
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</tbody>
</table>

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:
________________________________________________________________
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________________________________________________________________
________________________________________________________________
WSDOT Test Method T 812
Method of Test for Measuring Length of Drilled Concrete Cores

1. SCOPE
   a. This method is similar to AASHTO T-148 and covers the procedure for determining the length of a core drilled from a concrete structure, and from Portland cement concrete pavement.

2. APPARATUS
   a. The apparatus shall be a calipering device that will measure the length of axial elements of the core. While the details of the mechanical design are not prescribed, the apparatus shall conform to the requirements of 2 (B) to 2 (F).
   b. The apparatus shall be designed so that the specimen will be held with its axis in a vertical position by three symmetrically placed supports bearing against the lower end. These supports shall be short posts or stubs of hardened steel, and the ends that bear against the surface of the specimen shall be rounded to a diameter of not less than \(\frac{1}{4}\) inch more than \(\frac{1}{2}\) inch.
   c. The apparatus shall provide for the accommodation of specimens of different nominal lengths. (A range of at least 9 to 12 inches.)
   d. The calipering apparatus shall be designed so that it will be possible to make a length measurement at the center of the upper end of the specimen and at nine additional points spaced at equal intervals along the circumference of a circle whose center point coincides with that of the end area of the specimen and whose radius is not less than one-half nor more than three-fourths of the radius of the specimen.
   e. The measuring rod or other device that makes contact with the end surface of the specimen for measurement shall be rounded to a radius of \(\frac{1}{8}\) inch. The scale on which the length readings are made shall be marked with clear, definite, accurately-spaced graduations. The spacing of the graduations shall be 0.10 inch or a decimal part thereof.
   f. The apparatus shall be stable and sufficiently rigid to maintain its shape and alignment without a distortion or deflection of more than 0.01 inch during all normal measuring operations.

3. TEST SPECIMENS
   a. Cores used as specimens for length measurement shall be in every way representative of the concrete in the structure from which they are removed. The specimen shall be drilled with the axis normal to the surface of the structure, and the ends shall be free from all conditions not typical of the surfaces of the structure. Cores that show abnormal defects or that have been damaged appreciably in the drilling operation shall not be used.
4. PROCEDURE
   a. Before any measurement of the core length is made, the apparatus is calibrated with suitable gauges so that errors caused by mechanical imperfections in the apparatus are known. When these errors exceed 0.01 inch, suitable corrections are applied to the core length measurements.
   b. The specimen is placed in the measuring apparatus with the smoothest end of the core facing down, to bear against the three hardened-steel supports. The specimen is placed on the supports so that the central measuring position of the measuring apparatus is directly over the mid-point of the upper end of the specimen.
   c. Ten measurements of the length are made on each specimen, one at the central position and one each at nine additional positions spaced at equal intervals along the circumference of a circle of measurement as described in 2(D). Each of these ten measurements is read directly to 0.10 inch and to 0.01 inch either directly or by estimation.
   d. If, in the course of the measuring operation, it is discovered that at one or more of the measuring points the surface of the specimen is not representative of the general plane of the core end because of a small projection or depression, rotate the specimen slightly about its axis, and make a complete set of nine measurements in the new position.

5. REPORT
   a. The individual observations are recorded to the nearest 0.01 inch and the average of the ten measurements, expressed to the nearest 0.01 foot, are reported on DOT Form No. 350-067 under the column “Measured Thickness.”
Performance Exam Checklist
Method T 812 Checklist
Measuring Length of PCC Cores

Participant Name __________________________ Exam Date __________

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Only concrete measured?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Damaged cores not measured?</td>
<td></td>
<td></td>
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<tr>
<td>3. Apparatus calibrated?</td>
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<td></td>
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<tr>
<td>4. Smooth (top) end of core set on pins?</td>
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<tr>
<td>5. Center probe located at center of core?</td>
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<tr>
<td>6. Ten measurements taken?</td>
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<tr>
<td>7. Measurements read to 0.10 in. directly?</td>
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<td></td>
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<tr>
<td>8. Measurements read indirectly to 0.01 in.?</td>
<td></td>
<td></td>
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<tr>
<td>9. Measurements recorded to 0.01 in.?</td>
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<tr>
<td>10. Averaged and reported to 0.01 foot?</td>
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</tbody>
</table>

First attempt: Pass ☐ Fail ☐  Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________

Comments:
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WSDOT Test Method T 813

Field Method of Fabrication of 2-in. (50-mm) Cube Specimens for Compressive Strength Testing of Grouts and Mortars

1. SCOPE

This method covers the fabrication of 2-in. (50-mm) cube specimens for compressive strength testing of grouts and mortars.

2. EQUIPMENT

   a. Specimen Molds

   Specimen molds for the 2-in. (50-mm) cube specimens shall be tight fitting. The molds shall not have more than three cube compartments and shall not be separable into more than two parts. The parts of the molds, when assembled, shall be positively held together. The molds shall be made of hard metal not attacked by the cement mortar. For new molds, the Rockwell hardness number shall not be less than HRB 55. The sides of the molds shall be sufficiently rigid to prevent spreading or warping. The interior faces of the molds shall conform to the tolerances of Table 1.

      | 2 in. Cube Molds | 50-mm Cube Molds |
      |------------------|-------------------|
      | Parameter        | New | In Use | New | In Use |
      | Planeness of Sides | <0.001 in. | <0.002 in. | <0.025 mm | <0.05 mm |
      | Distance Between Opposite Sides | 2 in. + 0.005 in. | 2 in. + 0.02 in. | 50 mm + 0.13 mm | 50 mm + 0.50 mm |
      | Height of Each Compartiment | 2 in. + 0.001 in. to -0.005 in. | 2 in. + 0.01 in. to -0.015 in. | 50 mm + 0.25 mm to -0.013 mm | 50 mm + 0.25 mm to -0.38 mm |
      | Angle Between Adjacent FacesA | 90 + 0.5° | 90 + 0.5° | 90 + 0.5° | 90 + 0.5° |

   b. Base Plates

   Base plates shall be made of a hard metal not attacked by cement mortar. The working surface shall be plane and shall be positively attached to the mold with screws into the side walls of the mold.

   c. Cover Plates

   Cover plates shall be made of a hard metal or glass not attacked by cement mortar. The surface shall be relatively plane.

A Measured at points slightly removed from the intersection. Measured separately for each compartment between all the interior faces and the adjacent face and between interior faces and top and bottom planes of the mold.
d. Tamper

The tamper shall be made of a nonabsorptive, nonabrasive, nonbrittle material such as a rubber compound having a Shore A durometer hardness of 80 ± 10, or seasoned oak wood rendered nonabsorptive by immersion for 15 minutes in paraffin at approximately 392°F (200°C), and shall have a cross-section of ½ in. × 1 in. (13 mm × 25 mm) and a length of about 5 to 6 in. (125 to 150 mm). The tamping face shall be flat and at right angles to the length of the tamper.

e. Trowel

A trowel which has a steel blade 4 to 6 in. (100 to 150 mm) in length, with straightedges.

3. FIELD PROCEDURE

a. Three or more specimens shall be made for each period of test specified.

b. All joints shall be water tight. If not water tight, seal the surfaces where the halves of the mold join by applying a coating of light cup grease. The amount should be sufficient to extrude slightly when the halves are tightened together. Repeat this process for attaching the mold to the base plate. Remove any excess grease.

c. Apply a thin coating of release agent to the interior faces of the mold and base plate. (WD-40 has been found to work well as a release agent) Wipe the mold faces and base plate as necessary to remove any excess release agent and to achieve a thin, even coating on the interior surfaces. Adequate coating is that which is just sufficient to allow a distinct fingerprint to remain following light finger pressure.

d. The grout or mortar shall be mixed according to the manufacturer’s instructions. Begin molding the specimens within an elapsed time of not more than 2 ½ minutes from completion of the mixing.

e. For plastic mixes, place a first layer of mortar about 1 in. (25 mm) deep in all the cube compartments (about one-half the depth of the mold). Tamp the mortar in each cube compartment 32 times in about 10 seconds making four rounds, each round perpendicular to the other and consisting of eight adjoining strokes over the surface of the specimen, as illustrated in Figure 1, below. The tamping pressure should be just sufficient to ensure uniform filling of the molds. The four rounds of tamping (32 strokes) shall be completed in one cube before going on to the next. When the tamping of the first layer is completed, slightly over fill the compartments with the remaining mortar and then tamp as specified for the first layer. During tamping of the second layer, bring in the mortar forced out onto the tops of the molds after each round of tamping, by means of gloved fingers and the tamper, before starting the next round of tamping. On completion of tamping, the tops of all the cubes should extend slightly above the tops of the molds.
f. Bring in the mortar that has been forced out onto the tops of the molds with a trowel and smooth off the cubes by drawing the flat side of the trowel (with the leading edge slightly raised) once across the top of each cube at right angles to the length of the mold. Then, for the purpose of leveling the mortar and making the mortar that protrudes above the top of the mold of more uniform thickness, draw the flat trailing edge of the trowel (with leading edge slightly raised) once lightly along the length of the mold. Cut off the mortar to a plane surface flush with the top of the mold by drawing the straight edge of the trowel (held nearly perpendicular to the mold) with a sawing motion over the length of the mold.

g. When fabricating fluid mixes, steps e. and f. need not be followed. Instead, the cube mold is filled with mortar and cut off to a plane surface with a sawing motion over the length of the mold.

h. Immediately after molding, place cover plate on top of the mold, cover the sample with wet burlap, towels, or rags, seal it in a plastic sack in a level location out of direct sunlight, and record the time. Allow the sample to set undisturbed, away from vibration, for a minimum of four six hours before moving.

i. Deliver the sample to the Regional or State Materials Laboratory in the mold with the cover plate in wet burlap, towels or rags sealed in a plastic bag within 24 hours. Time of molding MUST be recorded on the Transmittal. If delivery within 24 hours is unachievable, contact the Laboratory for instructions on caring for the cubes.

j. Once received in the lab, the molded sample is to be immediately placed in a moist curing room, with the upper surfaces exposed to the moist air but protected from dripping until the sample is a minimum of 20 hours old or has cured sufficiently that removal from the mold will not damage the cube. If the specimens are removed from the mold before they are 24 hours old they are to be kept on the shelves of the moist curing room until they are 24-36 hours old.

k. When the specimens are 24-36 hours old, immerse them in a lime-saturated water storage tank. (Note 1) The specimens are to remain in the storage tank until time of test. (Curing test specimens of material other than hydraulic cement shall be in conformance with the manufacturer’s recommendations.)

Note 1: The storage tank shall be made of noncorroding materials. The water shall be saturated with calcium hydroxide such that excess is present. Stir the lime-saturated water once a month and clean the bath as required by AASHTO M-201.
**Performance Exam Checklist**

*Field Method of Fabrication of 50-mm (2-in.) Cube Specimens for Compressive Strength Testing of Grouts and Mortars*

**WSDOT Test Method T 813**

<table>
<thead>
<tr>
<th>Procedure Element</th>
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<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>has the current calibration/verification tags present?</td>
<td></td>
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<tr>
<td>3. Three cubes made for each time period of test?</td>
<td></td>
<td></td>
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<tr>
<td>4. All joints (mold halves, mold to base plate) shall be water tight?</td>
<td></td>
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</tr>
<tr>
<td>5. Adequate coating of release agent applied to interior surfaces of the mold?</td>
<td></td>
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</tr>
<tr>
<td>6. Grout or mortar mixed according to manufacturer’s instructions?</td>
<td></td>
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</tr>
<tr>
<td>7. Molding began within 2-1/2 minutes from completion of mixing?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Molding performed in two lifts? (not necessary if mix is fluid)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Lifts tamped 32 times, made up of 4 rounds of 8, each perpendicular to the other?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. For second layer, mortar forced out of the mold brought back in before each round?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Mix extends slightly above the mold at the completion of tamping?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12. Mortar smoothed by drawing flat side of trowel across each cube at right angles?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. Mortar leveled by drawing the flat side of trowel lightly along the length of mold?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14. Mortar cut off flush with mold with edge of trowel using sawing motion?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15. Time of molding recorded?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16. Cover plate placed on top of the mold and covered with wet burlap, towel or rag?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17. Covered sample sealed in a plastic sack in a level location out of sunlight?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18. Sample delivered to the laboratory in the mold within 24 hours?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>19. Transmittal includes the time of molding?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

First attempt: Pass ☐  Fail ☐  Second attempt: Pass ☐  Fail ☐

Signature of Examiner __________________________________________


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T 813  January 2006  T 813
Page 5 of 6
WSDOT Test Method T 814

Method of Test for Water Retention Efficiency of Liquid Membrane-Forming Compounds and Impermeable Sheet Materials for Curing Concrete

1. SCOPE

This method is intended for laboratory use in determining the efficiency of liquid membrane-forming compounds, papers, and other impermeable sheet materials for curing concrete, as measured by their ability to prevent moisture loss during the early hardening period.

2. APPARATUS

a. Molds — The molds shall be seamless tin ointment cans 3.4374 in. (87.31 mm) in diameter by 1.1874 in. (30.16 mm) deep.

b. Mortar — Mortar for test specimens shall consist of a portland cement mixture of equal parts of Type II and Type III Portland cements, air entraining agent, sand and water, all well mixed in the proportions by mass of 1 part cement, 0.003 parts neutralized Vinsol resin (NVX), 2.2 parts sand, and 0.4 parts water. The portland cement shall conform to the Standard Specifications for Portland cement, ASTM Designation: C 150. The sand shall conform to the specifications for graded standard sand, ASTM Designation C 778.

   (1) Mixing — Mixing shall preferably be done in a room in which the air temperature is 70°F ± 5°F (21°C ± 2.8°C) and the relative humidity 40 to 60 percent. The temperature of the mortar at the time of molding shall be 70°F ± 5°F (21°C ± 2.8°C). The cement and sand shall be mixed dry in a nonabsorptive vessel and the specified quantity of water, in which the Vinsol resin has been dissolved, shall be added. The mixture shall be allowed to absorb the water for a period of one minute. Testing shall be accomplished by WSDOT Test Method No. 307.

   (2) Molding — The mold shall be filled with mortar and the mortar puddled with the gloved fingers only to the extent required for uniform and thorough consolidation. Excess mortar shall be removed and the surface smoothed with the gloved hands, immediately after which the surface shall be finished with one pass of a straightedge, using pressure and a sawing motion.

   (3) Storage — Immediately after molding, the excess mortar on the outside of the molds shall be removed with a damp cloth. The molds containing the specimens shall be placed in a moist closet in an atmosphere maintained at 73.4°F ± 3.6°F (23°C ± 2°C) and relative humidity of not less than 95 percent.
3. PROCEDURE

a. Application of Compound — Four hours after molding, the specimens shall be removed from the moist closet and the junction of specimen and mold sealed by means of a hot applied rosin/paraffin wax sealing material applied to the edges by means of a small brush or syringe, and shall not extend more than \( \frac{3}{16} \) in. (4.8 mm) onto the surface of the specimen.

A single coat of curing compound shall be applied to each specimen by the use of a brush. The compound shall be applied at the rate of 1.29 ml per test specimen. The proper mass of compound for the prescribed coverage shall be determined from the specific gravity of the material. This mass of compound (±0.05 g) shall be measured by weighing the test specimens before and after application. When testing impermeable sheet materials, the sample is taped over the specimen with a pressure sensitive tape.

b. Determination of Moisture Loss — One-half hour after application of the curing compound, each of the coated specimens shall be place individually in 4-in. (101.6 mm) Scheibler desiccators. A 30 to 50 mm low form glass weighing bottle containing approximately 20 g of potassium hydroxide pellets accurately weighed to the nearest 5 mg, shall be placed on a wire screen supported about \( \frac{1}{4} \) in. (6.3 mm) above the surface of the mortar. The cover of the weighing bottle shall be removed and desiccator closed tightly with its cover. The desiccators shall be kept at a temperature of 70°F ± 2°F (21°C ± 1°C) for 72 hours. At the conclusion of this period, the desiccator shall be opened, the cover shall be placed on the weighing bottle, and the mass (weight) determined to the nearest 5 mg. The increase in mass (weight) of the potassium hydroxide shall be taken as the loss of moisture from the specimen. The results of not less than three test specimens shall be averaged in computing the loss of moisture.

4. REPORTS

a. Test results will be reported on Test Data Form WSDOT 350-034.
1. SCOPE
   a. This method is used to compare the compressive strength of cylinders and/or the flexural strength of beams made from Steilacoom Aggregates (PS-B-1) with those made from other pits. Data may be maintained on work sheets or input into the LIMS Computer Technical System labeled “Steilacoom.”

2. EQUIPMENT
   a. Balance — The balance or scale shall be sensitive to 0.01 lb. and will conform to AASHTO M 231.
   b. Oven — An oven of appropriate size which shall be capable of maintaining a uniform temperature of 230° ± 9°F (110° ± 5°C).
   c. Sieves shall conform to AASHTO M 92.
   d. Sand Separator — Screens of the following sizes for fine aggregates: U.S. No. 4, U.S. No. 8, U.S. No. 16, U.S. No. 30, and US No. 50 (4.75 mm, 2.36 mm, 1.18 mm, 600 µm, and 300 µm), or other commercial sizes which produce the gradation required.
   e. Plastic Buckets with Sealing Lids — The buckets shall be strong enough for repeated use of being filled with aggregate and water.
   f. Concrete Mixer — A power driven drum or revolving paddle mixer shall be capable of thoroughly mixing batches of prescribed sizes at the required slump.
   g. Molds — Molds for cylinders must conform to AASHTO M 205, 6 in. × 12 in. (150 mm × 300 mm) size. Molds for beams must conform to AASHTO T 126, 6 in. × 6 in. × 21 in. (150 mm × 150 mm × 425 mm) size.
   h. Curing Room and Lime Bath — Curing apparatus shall conform to AASHTO M 201.
   j. Slump Cone and Tamping Rod — The cone and tamping rod shall conform to AASHTO T 119.
   k. Aggregate Washer — A rotating drum type concrete mixer with the paddle size reduced to 1 ¾ in. × 12 ½ in. (44.45 mm × 317.50 mm) and have the capability to wash all sizes of aggregate without loss of fines.

3. SAMPLES
   a. Comparison Sample: A representative sample of pit run material shall be forwarded to WSDOT, FOSSC Materials Laboratory, 1655 South Second Avenue, Tumwater, WA 98512. Sufficient material will be required to meet the grading requirements shown in Table 1.
### Table 1

<table>
<thead>
<tr>
<th>Aggregate Size</th>
<th>Structural Concrete</th>
<th>Concrete Paving</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ½ in. - 1 ¼ in. (37.5 mm - 31.5 mm)</td>
<td>0 lbs. (0 kg)</td>
<td>50 lbs. (25 kg)</td>
</tr>
<tr>
<td>1 ½ in. - 1 in. (31.5 mm - 25.0 mm)</td>
<td>0 lbs. (0 kg)</td>
<td>15 lbs. (7 kg)</td>
</tr>
<tr>
<td>1 in. - ½ in. (25.0 mm - 19.0 mm)</td>
<td>10 lbs. (5 kg)</td>
<td>45 lbs. (20 kg)</td>
</tr>
<tr>
<td>½ in. - ¼ in. (19.0 mm - 9.5 mm)</td>
<td>60 lbs. (27 kg)</td>
<td>55 lbs. (25 kg)</td>
</tr>
<tr>
<td>¼ in. - US #4 (9.5 mm - 4.75 mm)</td>
<td>25 lbs. (11 kg)</td>
<td>20 lbs. (10 kg)</td>
</tr>
<tr>
<td>US #4 - US #8 (4.75 mm - 2.36 mm)</td>
<td>15 lbs. (7 kg)</td>
<td>20 lbs. (10 kg)</td>
</tr>
<tr>
<td>US #8 - US #16 (2.36 mm - 1.18 mm)</td>
<td>15 lbs. (7 kg)</td>
<td>20 lbs. (10 kg)</td>
</tr>
<tr>
<td>US #16 - US #30 (1.18 mm - 600 µm)</td>
<td>15 lbs. (7 kg)</td>
<td>20 lbs. (10 kg)</td>
</tr>
<tr>
<td>US #30 - US #50 (600 µm - 300 µm)</td>
<td>15 lbs. (7 kg)</td>
<td>20 lbs. (10 kg)</td>
</tr>
<tr>
<td>US #50 - US #200 (300 µm - 75 µm)</td>
<td>10 lbs. (5 kg)</td>
<td>15 lbs. (7 kg)</td>
</tr>
</tbody>
</table>

b. If the pit source is a commercial pit, the material may be obtained from the stockpiles, however, sufficient material must be acquired to satisfy the grading requirements shown in Table 1.

c. Steilacoom and Candidate Aggregates — Shall be washed, dried, and graded. The coarse aggregate shall be midline graded to concrete aggregate specifications Standard Specification 9-03.1(3)C Gradings No. 5 for structural concrete or Gradings No. 2 for paving concrete. The fine aggregate shall be midline graded to conform to Standard Specifications 9-03.1(2)B Class 1.

d. Batch sizes shall be 1.23 cu. ft. (0.03 cubic meters) for cylinders and 1.64 cu. ft. (0.05 cubic meters) for beams. (This includes 25 percent additional material to make this a more workable batch.)

4. PREPARATION OF AGGREGATES

a. Obtain representative samples of fine and coarse aggregate from the Candidate Pit to test for LA Abrasion (AASHTO T 96), Specific Gravity and Absorption (SSD) of fine and coarse aggregate (AASHTO T 85 and 84), and Organic Content (AASHTO T 21). The test for Organic Content is to be done on washed sand which has been dried back to a constant mass (weight) at a temperature not to exceed 140°F (60°C). If the Candidate Aggregate fails the Organic Content (AASHTO T 21), additional samples shall be tested as per AASHTO T 71. The cube comparison result from AASHTO T 71 shall be minimum of 95 percent.

b. If the material is wet, oven dry to constant mass (weight).

c. Split out approximately 50 lbs. (25 kg) of pit run material. Grade it through all required screens in order to calculate the amount of pit run material needed to supply enough of each size required. If it appears that there is not enough material on hand, request more from the same source.

d. Wash the aggregates thoroughly and dry in an oven at 230 + 9°F (110 ± 5°C).

e. Separate oven dry aggregates into coarse and fine. Use the following screens for separating the coarse aggregate for cylinders: 1 in. (25.0 mm), ½ in. (19.0 mm), ¼ in. (9.5 mm), and U.S. No. 4 (4.75 mm). For separating coarse aggregate to be used for beams use the following screens: 1 ½ in. (37.5 mm), 1 ¼ in. (31.5 mm), 1 in. (25 mm), ½ in. (19 mm), ¼ in. (9.5 mm), and U.S. No. 4 (4.75 mm). For separating fine aggregate use U.S. No. 8 (2.36 mm), U.S. No. 16 (1.18 mm), U.S. No. 30 (600 µm), U.S. No. 50 (300 µm) and U.S. No. 200 (75 µm)
sieves. Obtain representative samples of fine and coarse aggregate for LA test (coarse aggregate only). Specific gravity, SSD absorption and determination of Organic content. For organics a sample of fine aggregate (approximately 300 g) is needed (AASHTO T 21). The test for Organic content is to be done on washed sand which has been dried back to a constant weight (mass) at a temperature not to exceed 140°F (60°C). If the material fails this Organic Content test then additional samples are to be taken for tests as per AASHTO T 71. This cube comparison result shall be a minimum of 95 percent.

f. If the coarse aggregate is crushed, an adjustment must be made according to Construction Manual Section 6-2.1C. Subtract 8 percent from the coarse aggregate batch weights. Calculate the total weight subtracted. Divide and add this weight (mass) proportionately among the grading of the fine aggregate.

g. If the material is 100% fractured the following shall be followed:

**Proportional Breakdown**

1 in. – ¾ in.
(25.0 mm – 19.0 mm) = 8.5 lb. (4.0 kg) of total coarse aggregate × 8 percent = correction of 0.7 lb. (0.32 kg)

¾ in. – ⅞ in.
(19.0 mm – 9.5 mm) = 55.4 lb. (25 kg) of total coarse aggregate × 8 percent = correction of 4.4 lb. (2.00 kg)

⅞ in. – U.S. No. 4
(9.5 mm – 4.75 mm) = 21.3 lb. (10 kg) of total coarse aggregate × 8 percent = correction 1.7 lb. (0.8 kg)

h. If the material is not 100 percent fracture, the percentage of fracture shall be determined and the sample shall be adjusted proportionally. Example: If the midline graded mix sample is 25 percent fractured, the adjustment should be 2 percent. If the midline graded sample is 50 percent fractured, the adjustment should be 4 percent, etc. Fracture shall be determined by WSDOT Test Method No. 103 and rounded to the nearest 10 percent for this adjustment.

i. Record all data using the worksheets or the LIMS Computer Technical System, for cylinders and/or beams, weigh the aggregate, place in tared plastic buckets, fill with water, cover with lid, and soak for approximately 15 hours. At the end of the soaking period and just prior to making concrete, drain as much water as possible from the aggregate. Weigh the aggregate and compute the amount of free water present.

j. Calculate the add water needed and subtract the amount of free water in both the coarse and fine aggregates from the maximum amount of water recommended.

5. **CALCULATION OF BATCH WEIGHTS**

a. Adjust the Candidate batch weights for Specific Gravity (Sp. Gr.) by multiplying each batch weight by the Sp. Gr. of the Candidate pit then divide by the Sp. Gr. of the Steilacoom pit, which is kept on file in at the Materials Laboratory pit file.

Example: size ¾– ⅞ (19.0 mm – 9.5 mm), 55.35 lb. (25.101 kg) × \( \frac{2.71}{2.67} \) = 55.90 lb. (25.351 kg)

**Note:** The coarse Sp. Gr. is often different from the fine Sp. Gr.
6. WEIGHING UP THE BATCHES
   a. Steilacoom Aggregate: The sum of the coarse aggregate will be divided amongst several
      buckets for ease of handling. The fine aggregate will be in several buckets also.
         (1) Place first bucket for Steilacoom coarse aggregate on the scale.
         (2) Zero the scale, lift the bucket, and record the bucket tare weight.
         (3) Weigh out approximately 30 lb. (13 kg) from the first several sizes of aggregate, zeroing
             the scale after each size.
         (4) Zero the scale, remove the bucket of aggregate, and record the weight (mass) of the
             aggregate w/bucket.
         (5) Proceed with the next bucket as in steps 1 through 4, being very careful when breaking
             up the weight (mass) of one size aggregate between two different buckets.
         (6) Weigh out all the Steilacoom coarse aggregate. Then, in separate buckets, proceed with
             the Steilacoom fine aggregate in the same manner.
   b. Repeat the process of steps 1 through 6 for the candidate aggregate. Buckets should all be
      distinctly marked.
   c. Fill each bucket with water to cover aggregate and soak overnight.
   d. Weigh up cement for each batch. (Type II, in stock at the FOSSC Materials Laboratory.)
7. CALCULATIONS FOR THE WATER
   a. Drain each bucket over U.S. No. 200 (75 μm) reinforced sieve. Return material from sieve to
      the bucket. Weigh and record wet weight aggregate with bucket. Cover with tight fitting lid to
      retain moisture.
         Dry aggregate = aggregate w/bucket – bucket tare
   b. Calculate the amount of absorbed water by multiplying the percent absorption (results of
      coarse and fine absorption, AASHTO T 84 and T 85) by the amount of dry aggregate in each
      bucket. The Steilacoom Coarse and Fine Absorption results are on file at the FOSSC Materials
      Laboratory.
         Dry aggregate wt = dry aggregate with bucket – bucket wt (tare)
   c. Calculate the free water in each bucket by subtracting the dry aggregate w/bucket and the
      absorbed water from the weight of the wet aggregate w/bucket. This figure represents the
      water surrounding the aggregate above saturated surface dry.
   d. Calculate the total free water for all the Steilacoom buckets and for all the Candidate buckets.
   e. Based upon the maximum water/cement ratio for structural concrete class 4000 No Air
      (Standard Specifications 6-02.3(2)C), calculate the maximum amount of water per batch.
      Subtract from that the total amount of free water. Record this as the H2O to weigh up for
      the batch. Calculate and weigh out water for the Steilacoom and for the Candidate batches.
      Label each flask.
8. MIXING PROCEDURE
   a. Make five cylinders for structural concrete and three beams for paving concrete from pit PS-B-1 and from the candidate pit. The proper sequence for mixing samples shall be first, the buttering batch; second, the candidate batch; and third, the Steilacoom batch. (The entire buttering batch is to be discarded.)
   b. Lay out all the necessary equipment.
   c. Begin by making up a buttering batch using Steilacoom aggregates.

   **Size Aggregate**
   - U.S. No. 16 - U.S. No. 30 (1.18 mm - 600 µm) .65 lb. (1.45 kg)
   - U.S. No. 30 - U.S. No. 50 (600 - 300 µm) .65 lb. (1.45 kg)
   - U.S. No. 50 - U.S. No. 200 (300 - 75 µm) .65 lb. (1.45 kg)
   - Cement .80 lb. (1.75 kg)
   - Water 250 ml.

   Mix by hand in a container, then butter mixer sides, bottom, and paddles. Remove and discard any excess.
   
   d. Always start with the Candidate batch first. Place the coarse aggregate in mixer. Add part of the weighed up water. Start the mixer, add fine aggregate and cement. Slowly add water from the flask until the mix appears to be at the desired slump, 3 in. (75 mm) for cylinders, 1 in. (25 mm) for beams. Start timer. Mix for three minutes. Stop mixer. Let rest for three minutes, covering the bowl with damp towels to retain moisture. Remove towels and mix two more minutes.
   
   e. Perform a slump test according to AASHTO T 119. Return slump test material to mixer. Record slump to nearest 1/4 in. (6 mm) If slump is too low, continue mixing, adding more water from the flask. **Note:** Do not add more than what is in the flask so as not to exceed the water/cement ratio. Mix 1 more minute. Repeat slump test and record result. Return the slump material to the mixer. Mix for 30 seconds.
   
   f. Pull concrete to side of the bowl, scraping the bottom and sides. Remix with hands to eliminate segregation.
   
   g. Cast five cylinders or three beams according to AASHTO T 126.
   
   h. Scrape out and discard concrete left in the bowl, leaving approximately the same amount as was left by the buttering batch.
   
   i. Weigh and record the amount of water left remaining in the flask as the amount of water not used, subtract this from the maximum amount of water recommended. This will be the actual water in the mix and will be used to calculate the W/C ratio.
   
   j. Immediately proceed with mixing Steilacoom batch in the same manner as steps d.-h. above.
   
   k. Thoroughly wash all the equipment.
   
   l. 16 to 32 hours after molding, strip cylinders and/or beams. With permanent marker, label each with date of test (14 days from date made), Lab I.D. No., either Steilacoom or Candidate batch, and cylinder or beam number. Cylinders shall be stored in the Moist Curing Room; beams shall be immersed in saturated-lime solution at 73.4 ± 3°F (23 ± 1.7°C) for a minimum period of 20 hours prior to testing.
9. FINAL CALCULATIONS
   a. Subtract the amount of water not used from the maximum amount of water recommended to calculate the W/C ratio. This figure represents the actual water in the mix.
   b. Calculate the W/C ratio:
      \[
      \text{W/C Ratio} = \frac{\text{actual water in mix}}{\text{cement batch weight}}
      \]

10. HANDLING, TESTING, AND REPORTING OF SPECIMENS
    a. Cylinders are to be tested per AASHTO T 22 and beams are to be tested per AASHTO T 177 at 14 days.
    b. Record total load for each cylinder, and the total load for each beam. The computer database will do the final calculations.
    c. Final report shall include results of:
        Candidate Pit Number or Numbers
        Organics
        LA Abrasion
        Degradation (if requested)
        Specific Gravity and Absorption (Course and Fine Aggregates)
        Compare the strength of the Candidate as a percent of the Steilacoom strength.
WSDOT Test Method T 816

Method of Test for Parting Compound

1. SCOPE:
   a. This method of test covers the procedure used for determining if parting compound enables concrete to release cleanly from forms without any dusting or staining.

2. APPARATUS:
   b. APA grade High Density Overlaid (HDO) Plyform Class I.
   c. Vibrator or other consolidation device.
   d. Rubber gloves, clean rag, spray apparatus, or other type of applicator as per mfg. recommendations.

3. PROCEDURE:
   a. Apply parting compound to plyform surface as per manufactures recommendation.
   b. Prepare PCC Cl. 4000 concrete by mixing all preweighed materials together in mixing bowl. Visually mix the concrete to a 3.00” slump. Mix 3 minutes, rest 3 minutes with damp towel covering mixing bowl, and then resume mixing for another 2 minutes. Scrape concrete to one side of bowl. Remix by hand to get a representative sample. Take a proportionate amount of concrete for the size of board you have.
   c. Place concrete on prepared surface of the plyform board. Place vibrator on the board and allow vibrations to consolidate the concrete mix or utilize other consolidation method.
   d. Allow to sit uncovered for 24 +/- 2 hours
   e. Lift the specimen vertically without any twisting or sliding movement.
   f. Specimen should separate cleanly from form without any staining or dusting of concrete surface as per Standard Specification 6-02.3(17)J.

4. REPORT
   a. Note any non-specification elements observed. The Physical Testing Supervisor and/or Engineer will determine acceptability of the parting compound.
**Method T 816 Checklist**

*Method of Test for Parting Compound*

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Parting Compound applied to the plyform surface as per Mfg. recommendations?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Class 4000 concrete mixed correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. A portion of the concrete placed on the prepared surface of the plyform?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Concrete consolidated?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Allowed to sit uncovered for 24 +/- 2 hours?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Specimen lifted vertically without any twisting or sliding?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Separation, Staining, and Dusting determinations made correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. If non-conforming elements are observed, is the supervisor notified?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt:  Pass ☐  Fail ☐  
Second attempt: Pass ☐  Fail ☐

Signature of Examiner __________________________________________

Comments:

________________________________________________________________________
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________________________________________________________________________
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(Blank lines available for comments)
WSDOT Test Method T 817

Standard Test Method for Slump Flow of Self-Compacting Hydraulic-Cement Concrete

1. SCOPE

1.1 The slump flow test is used to characterize the fluidity of the concrete both in the laboratory and in the field. The Slump Flow test measures the diameter of the flow, and observe the material for resistance to segregation.

1.2 The values stated in English units are to be regarded as the standard.

1.3 This standard does not purport to address all of the safety problems, 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 AASHTO Standards:
T 141 Sampling Freshly Mixed Concrete

2.2 ASTM Standards:
C 172 Practice for Sampling Freshly Mixed Concrete

3. SUMMARY OF TEST METHOD

3.1 A sample of freshly mixed self compacting concrete is placed in a traditional slump cone from ASTM C 143. The mold is raised, and the concrete is allowed to flow. The diameter of the concrete is measured and reported as the slump flow of the concrete. The concrete is then observed for signs of segregation.

4. SIGNIFICANCE AND USE

4.1 This test method is intended to provide the user with a procedure to determine slump flow of plastic hydraulic-cement concretes.

4.2 This test method is considered applicable to plastic concrete having coarse aggregate up to 1 1/2 in. (37.5 mm) in size.

5. APPARATUS

5.1 Mold — The test specimen shall be formed in a mold made of metal not readily attacked by the cement paste. The metal shall not be thinner than 0.060 in. (1.5 mm) and if formed by the spinning process, there shall be no point on the mold at which the thickness is less than 0.045 in. (1.15 mm). The mold shall be in the form of the lateral surface of the frustum of a cone with the base 8 in. (200 mm) in diameter, the top 4 in. (100 mm) in diameter, and the height 12 in. (300 mm). Individual diameters and heights shall be within ± 1/8 in. (3.2 mm) of the prescribed dimensions. The base and the top shall be open and parallel to each other and at right angles to the axis of the cone. The mold shall be provided with foot pieces and handles similar to those shown in Figure 1. The mold shall be constructed without a seam. The interior of the mold shall be relatively smooth and free from projections. The mold shall be free from dents, deformation or adhered mortar.
5.1.1 If the condition of any individual mold is suspected of being out of tolerance from the as manufactured condition, a single traditional slump comparative test shall be performed. If the test results differ by more than 0.50 in. (15 mm) from that obtained using the metal mold, the mold shall be removed from service.

5.2 Torpedo level

5.3 Base — Flat, rigid surface, ½ to ¾ inch thick, cut 32 inches by 32 inches (800mm X 800mm) with surface sealed to have a non-absorbent face. Draw two centerlines on the board and two concentric circles, the first circle has a diameter of 20 inches (500mm). The second circle has a larger diameter, which may be 26 to 28 inches in diameter (about 650 to 700 mm). The second circle serves as a visual guide.

5.4 Ruler or tape measure 33 inches (800mm) minimum

![Mold for Slump Test](Figure 1)
6. **SAMPLE**

6.1 The sample of concrete from which test specimens are made shall be representative of the entire batch. It shall be obtained in accordance with FOP for WAQTC TM 2.

7. **PROCEDURE**

7.1 Place slump flow base constructed of a rigid, smooth non absorbent material on a stable horizontal surface. Moisten the surface of the board and wipe off excess water. Place the slump cone in the center of the flow board. The slump cone shall be held firmly in place by the operator standing on the two foot pieces.

7.2 Completely fill the slump cone in one lift. Do not rod or vibrate the sample. A funnel is helpful to prevent excess spillage.

7.3 Strike off excess concrete from the top of the slump cone, and remove and concrete that has fallen onto the board around the cone. Lift the cone vertically approximately 12 inches, hold in center position until all concrete has exited the slump cone.

7.4 Measure the final flow across the diameter at two points, 90 degrees apart. The two diameter measurements should not be greater than two inches in difference (if they are it may indicate a sloping board). Complete the entire test from the start of the filling through removal of the mold without interruption and complete it within an elapsed time of $2\frac{1}{2}$ min.

7.5 The concrete should flow out to final slump diameter with no segregation. Check paste bleed on the surface and around the perimeter. The mix should appear uniform, with the coarse aggregate visible and evenly dispersed.

8. **Report**

8.1 Report the slump flow as the average of the two measurements. Report the spread to the nearest $\frac{1}{2}$ inch (12mm).
Performance Exam Checklist

Slump Compacting Hydraulic Cement Concrete
FOP for AASHTO T 817

Participant Name ________________________________ Exam Date ____________

Procedure Element                          Yes No
1. The tester has a copy of the current procedure on hand? ☐ ☐
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? ☐ ☐
3. Cone placed in the center of the base plate? ☐ ☐
4. Cone held firmly against the base by standing on the two foot pieces? Cone not allowed to move in any way during filling? ☐ ☐
5. Representative samples scooped into the cone? ☐ ☐
6. Cone filled in one lift? ☐ ☐
7. Concrete struck off level with top of cone using tamping rod? ☐ ☐
8. Excess concrete removed from around the base of the slump cone? ☐ ☐
9. Cone lifted upward approximately 12 in. (300 mm) in one smooth motion, without twisting the cone? ☐ ☐
10. Diameter measured at 90 degrees to each other and averaged ☐ ☐
11. The concrete seems to be uniform ☐ ☐

First attempt:  Pass ☐ Fail ☐                      Second attempt:  Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
WSDOT Test Method T 818

Air Content of Freshly Mixed Self-Compacting Concrete by the Pressure Method

1. The air test will be performed in accordance with WSDOT FOP for WAQTC T 152 with the following modifications to Procedure:
   a. Delete Note 6
   b. Change item 3 to read; Fill the base completely in one continuous lift.
   c. Change item 4 to read; Do not consolidate the concrete by rodding, vibration or tamping. When the base is filled, lightly tap around the exterior of the base with a rubber mallet to allow entrapped air bubbles to escape.
   d. Delete items 5-11.
Performance Exam Checklist

Air Content of Freshly Mixed Self Compacting Concrete by the Pressure Method

WSDOT T 818

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Container filled in one layers, slightly overfilling?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Sides of the container lightly tapped with the mallet?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Concrete struck off level with top of container using the bar and rim cleaned off?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Inside of cover cleaned and moistened before clamping to base?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Using a Type B Meter

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>10. Both petcocks open?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>11. Air valve closed between air chamber and the bowl?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>12. Water injected through petcock until it flows out the other petcock?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>13. Water injection into the petcock continued while jarring and tapping the meter to insure all air is expelled?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>14. Air pumped up to initial pressure line?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>15. A few seconds allowed for the compressed air to stabilize?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>16. Gauge adjusted to the initial pressure?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>17. Both petcocks closed?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>18. Air valve opened between chamber and bowl?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>19. Sides of bowl tapped with the mallet?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>20. Air percentage read after lightly tapping the gauge to stabilize the hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>21. Air valve closed and then petcocks opened to release pressure before removing the cover?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>22. Air content recorded to 0.1 percent?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>23. All calculations performed correctly?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt:  Pass ☐  Fail ☐  Second attempt:  Pass ☐  Fail ☐

Signature of Examiner __________________________________________

This checklist is derived, in part, from copyrighted material printed in ACI CP-1, published by the American Concrete Institute.
WSDOT Test Method T 819
Making and Curing Self-Compacting Concrete Test Specimens in the Field

1. The cylinders will be made and cured in accordance with WSDOT FOP for AASHTO T 23 with the following modifications to Section 8 Molding Specimens:
   a. Change 8.2 to read; Place the concrete in the mold using a scoop, blunted trowel or shovel. Ensure that the scoopful, trowelful or shovelful is representative of the entire sample. Fill the mold completely in one continuous lift.
   b. Delete 8.2.1 – 8.2.2 including Table 2 and Table 3.
   c. Change 8.3 to read: Consolidation:
      Do not consolidate the concrete by rodding, vibration or tamping. When the cylinder is filled lightly tap around the exterior with an open hand 10 to 15 times to release any large air bubbles that may have been trapped.
   d. Delete 8.3.1 – 8.3.3.2
Performance Exam Checklist

Making and Curing Self-Compacting Concrete Test Specimens in the Field

WSDOT T 819

Participant Name __________________________________________ Exam Date ________________

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Molds placed on a level, rigid, horizontal surface free of vibration?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Making of specimens begun within 15 minutes of sampling?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>4. Concrete placed in the mold, moving a scoop or trowel around the perimeter of the mold to evenly distribute the concrete as discharged?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>5. Mold filled in one lift?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>6. Sides of the mold lightly tapped 10-15 times after rodding each layer?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>a. with mallet for reusable steel molds</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>b. with the open hand for flexible light-gauge molds</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>7. Strike off excess concrete, and finished the surface with a minimum of manipulation?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>8. Specimens covered with non-absorbent, nonreactive cap or plate?</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

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Comments:

_________________________________________________________________________________________
_________________________________________________________________________________________
_________________________________________________________________________________________
_________________________________________________________________________________________
_________________________________________________________________________________________
WSDOT Test Method T 914

Practice for Sampling of Geotextiles for Testing

1. SCOPE
   a. This practice covers a procedure for use in the division of shipments of geotextiles into lots and the sampling of lots for testing.

2. DEFINITIONS
   a. Geotextile — Any permeable textile used with foundation, soil, rock, earth, or any other geotechnical material, as an integral part of a manmade product, structure, or system.
   b. Lot — All geotextile rolls within a consignment (i.e., all rolls sent to the project site) which were manufactured at the same manufacturing plant having the same product name and specifications, style, or physical characteristics of a particular geotextile product.
   c. Lot Sample — Sample(s) from one or more geotextile rolls taken at random to represent an acceptance sampling lot and used as a source of laboratory samples.
   d. Production Unit — As referred to in this practice, it shall be considered to be synonymous with the geotextile roll as shipped by the manufacturer. Two or more geotextile rolls joined together by sewn seams shall be considered as separate rolls.
   e. Minimum Average Roll Value — The test results of any sampled roll in a lot shall meet or exceed the minimum values specified.

3. SUMMARY OF PRACTICE
   a. Instructions are given within this practice for dividing shipments or consignments of geotextiles into lots and for the determination of the number of production units in a lot sample.

4. SIGNIFICANCE AND USE
   a. This sampling procedure will provide a representation of the lot which is adequate to establish minimum average roll values as defined by this practice.

5. PROCEDURE
   a. Division into lots — Divide the shipment or consignment into lots as defined by 2.b. above.
   b. Determination of lot sample size.
      (1) Take geotextile rolls for the lot sample. Consider the geotextile roll to be the primary sampling units.
      (2) Select at random the number of geotextile rolls from each lot for the lot sample corresponding to the total number of units in the lot, as shown in Table 1. If the lot as defined in this practice contains only a portion of a geotextile roll, the lot shall be considered to contain one production unit for the purpose of sampling.
c. Laboratory sample selection.

(1) Obtain a laboratory sample from each geotextile roll in the lot sample. The minimum laboratory sample size shall be a minimum of 6 feet long by the full width of the geotextile roll. The laboratory sample must also contain a minimum area of 6.0 yd.² (5.0 m²) of geotextile.

(2) The laboratory sample should not be taken from the outer wrap of the roll nor the inner wrap of the core (i.e., do not take the sample from the geotextile at the very ends of the roll).

<table>
<thead>
<tr>
<th>Number of Rolls in Lot</th>
<th>Number of Rolls to be Selected for Lot Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 to 24</td>
<td>1</td>
</tr>
<tr>
<td>25 to 49</td>
<td>2</td>
</tr>
<tr>
<td>50 to 99</td>
<td>3</td>
</tr>
<tr>
<td>100 to 125</td>
<td>5</td>
</tr>
<tr>
<td>125 to 216</td>
<td>6</td>
</tr>
<tr>
<td>217 to 343</td>
<td>7</td>
</tr>
<tr>
<td>344 to 512</td>
<td>8</td>
</tr>
<tr>
<td>513 to 729</td>
<td>9</td>
</tr>
<tr>
<td>730 to 1,000</td>
<td>10</td>
</tr>
</tbody>
</table>

6. SAMPLE SUBMITTAL

a. All geotextile samples submitted to the State Material Laboratory are to be prepared and shipped as follows:

Woven Geotextiles — Roll sample around a 4-in diameter minimum, tube such as PCV pipe or cardboard mailing tube and wrap to protect sample from shipping damage and ultraviolet light (UV) exposure.

Nonwoven Geotextiles — Fold sample to a minimum 2 ft x 2 ft (0.6 m x 0.6 m) square, or roll as for woven geotextile. Wrap or box sample for shipment to protect from shipping damage and ultraviolet (UV) light exposure.

b. If sample is for Acceptance of Lots used on project, the following information must be submitted with the sample:

(1) Manufacturer’s name and current address.
(2) Full product name.
(3) Geotextile roll number(s).
(4) Proposed geotextile use(s).
(5) Certified test results.
(6) The Lot Number being submitted for acceptance. In lieu of a manufacturer provided Lot Number, the Bill of Lading Number can be used.

Testing by the State Materials Laboratory will not begin until all of the required information is received.
Performance Exam Checklist

Practice for Sampling Geotextiles for Testing
WSDOT Test Method T 914

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. Sampling</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>a. Shipment or consignment divided into lots.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>b. Determine the number of rolls in the shipment or consignment to be sampled from Table 1.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>c. Rolls to be sampled selected at random.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>d. Samples are a minimum 1.5 yd. (1.37 m) 6 ft. (1.83 m) long by the full width of the roll and a minimum of 6 sy (5 square meters).</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>e. Sample does not include outer wrap or inner wrap of the roll.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Shipment Preparation</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>a. Woven geotextiles must be rolled and shall not be folded.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>b. Non-woven geotextiles should be rolled or folded but not folded to less than a minimum of 2 feet square.</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>c. Wrap the sample (or box if folded) to protect from ultra-violet light exposure.</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

First attempt: Pass ☐ Fail ☐ Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

________________________________________________________________
________________________________________________________________
________________________________________________________________
________________________________________________________________
WSDOT Test Method T 915

Practice for Conditioning of Geotextiles for Testing

1. SCOPE
   a. This practice covers a procedure for conditioning geotextile specimens for testing and establishes atmospheric conditions which are acceptable for testing when the standard atmosphere for testing cannot be obtained due to local laboratory conditions.

2. APPLICABLE DOCUMENTS
   a. ASTM Standards.
      D 123 Terminology Relating to Textiles
      D 1776 Practice for Conditioning Textiles for Testing
      D 4439 Terminology for Geotextiles
      D 4533 Standard Test Method for Trapezoid Tearing Strength of Geotextiles
      D 4595 Standard Test Method for tensile Properties of Geotextiles by the Wide-Width Strip Method
      D 4632 Standard Test Method for Breaking Load and Elongation of Geotextiles (Grab Method)

3. DEFINITIONS
   a. Atmosphere for Testing Geotextiles — Air maintained at a relative humidity of 55 ± 25 percent relative humidity and temperature of 70° ± 4°F (21° ± 2°C).
   b. Geotextile — Any permeable textile used with foundation, soil, rock, earth, or any other geotechnical material, as an integral part of a manmade product, structure, or system.
   c. Specimen — A specific portion of a material or laboratory sample upon which a test is performed or which is taken for that purpose.
   d. Preconditioning — Atmospheric conditioning of a test specimen prior to testing in a specified environment in which the specimen is allowed to come to equilibrium with that specified preconditioning environment.

4. SUMMARY OF PRACTICE
   a. Specimens are preconditioned by soaking them in distilled water for a specified period of time and are tested at ambient laboratory room temperature and humidity conditions without allowing the specimens time to come to equilibrium with the ambient testing atmosphere.

5. USES AND SIGNIFICANCE
   a. The conditioning prescribed in this practice is designed to obtain reproducible test results on geotextiles.
6. APPARATUS
a. Water filled pan for soaking specimens.
b. Equipment for recording the temperature of the air and the water, and the humidity of the air.

7. PROCEDURE
a. Precondition specimens by immersing them in distilled water maintained at a temperature of 70\(^\circ\) ± 4\(^\circ\)F (21 ± 2\(^\circ\)C). The time of immersion must be sufficient to wet-out the specimens thoroughly, but must be a minimum of two hours. To obtain thorough wetting, add not more than 0.05 percent of a nonionic neutral wetting agent to the water.

b. After the specimens have been thoroughly wetted, remove each specimen from the water, and allow excess water contained in the pores of the specimen to drain from the specimen for a period of time less than or equal to one minute. After the specimen has drained during the maximum allowed time period of one minute, the specimen test must begin before nine minutes of time have elapsed from the end of the draining period.

Note 1: If more than a total of nine minutes from the time the specimen is removed from the water is allowed to elapse before the specimen test is actually begun, the specimen should not be considered to be thoroughly wetted. If this occurs, the specimen should be reimmersed for a minimum of two hours before a test is attempted again for that specimen. Thorough wetting is needed to ensure that the specimen is not affected by the ambient humidity conditions during testing if those ambient conditions are not at the standard atmosphere for testing.

c. The atmosphere for testing, geotextiles must be maintained at a temperature of 70\(^\circ\) ± 4\(^\circ\)F (21 ± 2\(^\circ\)C), and a relative humidity of 55 ± 25 percent.

d. If dry testing of the geotextile is required in addition to wet testing, the specimens must be conditioned in the atmosphere for testing as stated in Section 7.3. Specimen conditioning shall be accomplished in this case by allowing the specimens to reach moisture equilibrium in the atmosphere for testing. Equilibrium is considered to have been reached when the change in the mass of the specimen in successive weighings made at intervals of not less than two hours does not exceed 0.1 percent of the mass of the specimen. Specimen immersion requirements do not apply to specimens which are to be tested dry. Specimens tested dry must be tested in the atmosphere for testing as previously defined.
WSDOT Test Method T 923  
*Thickness Measurement of Geotextiles*

1. **SCOPE**
   a. This test method provides procedures for measuring the thickness of geotextiles.

2. **SUMMARY OF METHOD**
   a. Specimens from the geotextile sample submitted for testing are measured for thickness and the average of those measurements is used as the multiplier in converting permittivity to permeability. The measured thickness is reported as information only in case disagreements arise concerning the permeability of the material.

3. **APPARATUS**
   a. The thickness measurement instrument shall be a precision electronic micrometer conforming to ASTM D 1777-64
   b. The specimen cutting template shall meet the requirements of ASTM D 4491 for cutting the specimens to size.

4. **SPECIMENS FOR TESTING**
   a. Specimens shall be cut from the sample submitted for testing in the pattern and number specified in ASTM D 4491.

5. **CONDITIONING**
   a. Thickness testing shall be done prior to conditioning the specimens for permeability testing.

6. **PROCEDURE**
   a. Handle the specimens carefully to avoid altering the natural state of the material.
   b. Turn the machine on and allow it to warm up for approximately five minutes. As the machine cycles, clean presser foot and base by inserting a clean, lint free cloth between presser foot and base while in the raised position. Carefully and gently pull the cloth from between presser foot and base as the foot is in lowered position. Allow the machine to cycle for several readings and if no variation, zero machine with the zero set knob. Repeat cleaning and zeroing process a second time to ensure the base and foot are clean.
   c. As the machine cycles to a raised presser foot position, (with a zero reading) quickly but carefully insert the sample under the presser foot in a centered position.
   d. The machine will cycle presser foot down. Record the reading of the first cycle only. Record the reading to 0.00030 in. (0.01 mm).
   e. When the machine cycles up remove the sample and allow the machine to cycle with no sample. Check for zero reading. If machine does not read zero repeat cleaning procedure b.
f. Continue to measure the remaining specimens using these same procedures.

g. Average the readings and record the results.

h. Report the average reading on the worksheet for permeability testing as the nominal thickness of the geotextile.

7. CALIBRATION

a. The thickness measuring micrometer shall be calibrated in accordance with the manufacturers’ directions once a year, and will be checked accuracy on the first working day of each month. Accuracy shall be within 0.00030 in. (0.01 mm).
WSDOT Standard Practice T 925

Standard Practice for Determination of Long-Term Strength for Geosynthetic Reinforcement

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WSDOT Standard Practice T 925

Standard Practice for Determination of Long-Term Strength for Geosynthetic Reinforcement

Summary and Use of Standard Practice

Through this protocol, the long-term strength of geosynthetic reinforcements can be determined. This protocol contains test and evaluation procedures to determine reduction factors for installation damage, creep, and chemical/biological durability, as well as the method to combine these factors to determine the long-term strength. The long-term strength values determined from this protocol can be compared to the required design strengths provided in the contract for the geosynthetic structure(s) in question to determine if the selected product meets the contract requirements. This protocol can be used for initial product acceptance (e.g., for inclusion in the Qualified Products List), or for quality assurance (QA) to facilitate periodic review of products for which the long-term strength has been previously determined using this Standard Practice.

This protocol has been developed to address polypropylene (PP), polyethylene (PE or HDPE), and polyester (PET) geosynthetics. For other geosynthetic polymers (e.g., polyamide or PVA), the installation damage and creep protocols provided herein are directly applicable. While the chemical and biological durability procedures and criteria provided herein may also be applicable to other polymers (for example, hydrolysis testing as described in Appendix D is likely applicable to polyamide and PVA geosynthetics), additional investigation will be required to establish a detailed protocol and acceptance criteria for these other polymers. These other polymers may be considered for evaluation using this protocol once modifications to the chemical/biological durability aspects of this protocol have been developed and are agreed upon by the approval authority (e.g., WSDOT).

Abbreviations and Symbols

\[\begin{align*}
AASHTO &= \text{American Association of State Highway and Transportation Officials} \\
d_{50} &= \text{The grain size at 50\% passing by weight for the backfill} \\
HDPE &= \text{High Density Polyethylene} \\
MARV &= \text{The minimum average roll value for the geosynthetic, defined as two standard deviations below the mean for the product (i.e., 97.5\% of all test results will meet or exceed the MARV). For practical purposes from the user’s viewpoint, the average for a sample taken from any roll in the lot shipped to the job site should meet or exceed the MARV.} \\
MSE &= \text{Mechanically Stabilized Earth} \\
PET &= \text{Polyester} \\
PP &= \text{Polypropylene} \\
QPL &= \text{Qualified Products List} \\
RF &= \text{Combined reduction factor to account for long-term degradation due to installation damage, creep, and chemical/biological aging} \\
RF_{CR} &= \text{Strength reduction factor to prevent long-term creep rupture of the reinforcement} \\
RF_{DB} &= \text{Strength reduction factor to prevent rupture of the reinforcement due to long-term chemical and biological degradation}
\end{align*}\]
\[ RF_{id} = \text{Strength reduction factor to account for installation damage to the reinforcement} \]

\[ T_{dl} = \text{The long-term tensile strength which will not result in rupture of the reinforcement during the required design life, calculated on a load per unit of reinforcement width basis} \]

\[ T_{ult} = \text{The ultimate tensile strength of the reinforcement determined from wide width tensile tests} \]

\[ UV = \text{Ultraviolet light} \]

\[ WSDOT = \text{Washington State Department of Transportation} \]

**Definitions**

**Apertures**

The open spaces formed between the interconnected network of longitudinal and transverse ribs of a geogrid.

**Class 1 Structure**

Typically includes geosynthetic walls or slopes that support bridge abutments, buildings, critical utilities, or other facilities for which the consequences of poor performance or failure would be severe. In general, geosynthetic walls greater than 6 m (20 ft) in height and reinforced slopes greater than 9.2 m (30 ft) in height will be considered to be Class 1.

**Class 2 Structure**

All geosynthetic walls and slopes not considered to be Class 1.

**Confined Testing**

Geosynthetic testing in which the specimen is surrounded and confined by soil to simulate conditions anticipated for the geosynthetic in use.

**Effective Design Temperature**

The temperature that is halfway between the average yearly air temperature and the normal daily air temperature for the warmest month at the wall site.

**Hydrolysis**

The reaction of water molecules with the polymer material, resulting in polymer chain scission, reduced molecular weight, and strength loss.

**In-isolation Testing**

Geosynthetic testing in which the specimen is surrounded by air or a fluid (not soil).

**Installation Damage**

Damage to the geosynthetic such as cuts, holes (geotextiles only), abrasion, fraying, etc., created during installation of the geosynthetic in the backfill soil.

**Load Level**

For creep or creep rupture testing, the load applied to the test specimen divided by \( T_{lot} \), the short-term ultimate strength of the lot or roll of material used form the creep testing.

**Nonaggressive Environment**

For geosynthetic walls and slopes, soils which have a \( d_{50} \) of 4.75 mm or less, a maximum particle size of 31.5 mm or less, a pH of 4.5 to 9, and an effective design temperature of 30°C or less.

**Oxidation**

The reaction of oxygen with the polymer material, initiated by heat, UV radiation, and possibly other agents, resulting in chain scission and strength loss.

**Post-consumer Recycled Material**

Polymer products sold to consumers which have been returned by the consumer after use of the products for the purpose of recycling.
Product Line  A series of products manufactured using the same polymer in which the polymer for all products in the line comes from the same source, the manufacturing process is the same for all products in the line, and the only difference is in the product weight/unit area or number of fibers contained in each reinforcement element.

Sample  A portion of material which is taken for testing or for record purposes, from which a group of specimens can be obtained to provide information that can be used for making statistical inferences about the population(s) from which the specimens are drawn.

Specimen  A specific portion of a material or laboratory sample upon which a test is performed or which is taken for that purpose.

Survivability  The ability of a geosynthetic to survive a given set of installation conditions with an acceptable level of damage.

Test Methods and Practices Used

The following test methods and practices are used or referenced by Standard Practice T 925:

2.  AASHTO Bridge LRFD Specifications for Highway Bridges, 3rd Edition, 2004 with current interims
3.  ASTM D4354 Standard Practice for Sampling of Geosynthetics for Testing
4.  ASTM D4873 –Standard Guide for Identification, Storage, and Handling of Geosynthetic Rolls and Samples
5.  ASTM D5261 – Standard Test method for Measuring Mass per Unit Area of Geotextiles
8.  ASTM D-1248 – Standard Specification for Polyethylene Plastics Extrusion Materials for Wire and Cable
10.  WSDOT Test Method T 926 – Geogrid Brittleness Test
12.  ASTM D5818 – Standard Practice for Obtaining Samples of Geosynthetics from a Test Section for Assessment of Installation Damage
14.  ASTM D1557 – Standard Test Methods for Laboratory Compaction Characteristics of Soil Using Modified Effort (56,000 ft-lbf/ft³)(2700 kN-m/m³)
15.  ASTM D6992 – Accelerated Tensile Creep and Creep-Rupture of Geosynthetic Materials Based on Time-Temperature Superposition Using the Stepped Isothermal Method
19. ASTM D4355 – Standard Test Method for Deterioration of Geotextiles from Exposure to Ultraviolet Light and Water (Xenon-Arc Type Apparatus)
21. GRI-GG7 – Carboxyl End Group Content of PET Yarns
22. GRI-GG8 – Determination of the Number Average Molecular Weight of PET Yarns Based on a Relative Viscosity Value
24. ASTM D 3417-99 - Enthalpies of Fusion and Crystallization of Polymers by DSC
27. ASTM D 5885 – Standard Test Method for Oxidative Induction Time of Polyolefins by High-Pressure Differential Scanning Calorimetry

Per mutual agreement between the testing laboratory, the geosynthetic manufacturer, and the approval authority, “equivalent ISO standards and practices may be used in lieu of ASTM, AASHTO, or GRI standards and practices where equivalent procedures are available.

**Data Requirements for Initial Product Acceptance**

1. General Product Information (required for all geosynthetic reinforcement products):
   a. Geosynthetic type and structure.
   b. Spacing and dimensions of geogrid elements.
   c. Polymer(s) used for fibers, ribs, etc.
   d. Polymer(s) used for coating, if present.
   e. Roll size (length, width, and area).
   f. Typical lot size.
   g. Polymer source(s) used for product.
   h. For HDPE and PP, primary resin ASTM type, class, grade, and category (for HDPE use ASTM D-1248, and for PP use ASTM D-4101).
   i. For PET, minimum production number average molecular weight (ASTM D4603 and GRI:GG8) and maximum carboxyl end group content (GRI:GG7), with supporting test data. Information regarding the laboratory where the testing was conducted and date of testing shall also be provided.
j. % of post-consumer recycled material by weight.

k. Minimum weight per unit area for product (ASTM D5261).

l. MARV for ultimate wide width tensile strength (ASTM D4595 or ASTM D6637), with supporting test data. Information regarding the laboratory where the testing was conducted and date of testing shall also be provided.

n. UV resistance at 500 hours in weatherometer (ASTM D4355), with supporting test data (as a minimum, provide supporting data for one product in the product line, preferably the lightest weight product submitted in the product line). Information regarding the laboratory where the testing was conducted and date of testing shall also be provided.

o. In addition, to establish a baseline for quality assurance testing, oven aging tests conducted in accordance with ENV ISO 13438:1999, Method A (PP) or B (HDPE), for polyolefin geosynthetics shall be performed. As a minimum, the lightest weight product in the product line should be tested. Unexposed and post-exposure specimens shall be tested for tensile properties (ASTM D4595 or ASTM D6637).

p. for geogrids, evaluation of geogrid brittleness per WSDOT Test Method T 926.

2. Installation Damage Data Requirements (RFID):

Installation damage testing and interpretation shall be conducted in accordance with Appendix A. As a minimum, for each product tested, the following information should be obtained:

a. Date tests were conducted.

b. Name(s), location(s), and telephone number(s) of laboratory(ies) conducting the testing and evaluation.

c. Identify whether installation damage testing was conducted as a site specific evaluation for an actual construction project or was conducted as a non-site specific evaluation.

d. Description of specific procedures used to conduct the installation damage testing, including installation procedures, sample size, method of specimen selection, sample removal procedures, etc. Identify any deviations in the installation procedures relative to typical installation practice in full scale structures, if the testing was not site specific.

q. Photographs illustrating procedures used and the conditions at the time of the testing, if available.

r. Measured mass/unit area per ASTM D5261 for the sample tested for installation damage and for the sample used to establish the undamaged strength. Also obtain product manufacturer Quality Control (QC) data on the uncoated product (i.e., “greige -good”) for the lot used for installation damage testing.

g. Tensile test results for the product before exposure to installation conditions (i.e., virgin material), and whether both virgin and damaged samples were taken from the same roll of material, or just from rolls within the same lot of material.

h. Tensile test results for specimens taken from the damaged material after installation.
i. Tensile test results for both virgin and damaged specimens should include individual test results for each specimen, typical individual load-strain curves which are representative of the specimens tested, including associated calibration data as necessary to interpret the curves (curves in which strain and load/unit width are already calculated are preferred), the average value for each sample, the coefficient of variation for each sample, and a description of any deviations from the standard tensile test procedures required by Appendix A.

j. Gradation curves for backfill material located above and below the installation damage geosynthetic samples, including the d50 size, maximum particle size, and a description of the angularity of the soil particles per ASTM D2488, including photographs illustrating the soil particle angularity, if available.

k. Photographs and/or a description of the type and extent of damage visually evident in the exhumed samples and specimens.

l. RFID, and a description of the data interpretation method used to determine RFID for each sample.

3. Creep Data Requirements (RF<sub>CR</sub> and Creep Stiffness J):

Creep testing and interpretation shall be conducted in accordance with Appendices B and C. As a minimum, for each product tested, the following information should be obtained:

a. Date tests were conducted.

b. Name(s), location(s), and telephone number(s) of laboratory(ies) conducting the testing and evaluation.

c. Photographs illustrating the creep testing equipment and procedures used, as available.

d. Tensile test results for the product before creep testing (i.e., virgin material), and whether both virgin and creep tested samples were taken from the same roll of material, or just from rolls within the same lot of material.

e. Tensile test results should include individual test results for each specimen, typical load-strain curves which are representative of the specimens tested, including associated calibration data as necessary to interpret the curves (curves in which strain and load/unit width are already calculated are preferred), the average value for each sample, the coefficient of variation for each sample, and a description of any deviations from the standard tensile test procedures required by Appendices B and C.

f. Creep test procedures used, especially any deviations from the procedures required in Appendices B and C.

g. If RF<sub>CR</sub> is determined using data obtained in accordance with Appendix B, provide load and time to rupture for each specimen as a minimum; however, strain data as a function of time is desirable if available.

h. If RF<sub>CR</sub> is determined using data obtained in accordance with Appendix C, provide strain data as a function of time, and strain at beginning of tertiary creep (if rupture occurred), in addition to load applied and time to rupture (if rupture occurred), is required.

j. Creep data plots should include both major and minor gridlines for ease in viewing and interpreting the data.

k. If elevated temperature testing is conducted, creep data before and after time/load shifting, including shift factors used and a description of how the shift factors were derived, must be provided.
l. Data illustrating the variability of the creep test environment, including temperature and humidity, during the creep test time period, or some assurance that the creep test environment was maintained within the variation of temperature prescribed within Standard Practice T 925, must be provided.

m. A detailed description of creep extrapolation procedures used (i.e., step-by-step procedures and theoretical/empirical justification) if procedures other than those outlined in Appendices B and C are used.

n. Description of statistical extrapolation procedures used in accordance with Appendices B and C, if statistical extrapolation is performed.

o. RF_{CR}, and a description of how RF_{CR} was determined for each product.

p. In addition, regardless of which approach is used to determine RF_{CR}, creep strain data at a load level that results in a strain of 2% and 4% at approximately 1,000 hours shall be submitted to determine the low strain (i.e., 2%) creep stiffness at 1,000 hours and at the specified design life (typically 75 years) using isochronous curves determined in accordance with Appendix C.

4. Long-Term Durability Data Requirements (RF_{D}):

As a minimum, the durability test data requested in part (1), which include molecular weight and CEG for PET, oven aging tests for polyolefins, and UV resistance for all polymers, shall be provided.

If it is desired to submit detailed durability performance test data to justify a lower RF_{D}, or to allow use in environments classified as chemically aggressive, durability testing and interpretation shall be conducted in accordance with Appendix D, and, as a minimum, for each product tested, the following information should be obtained:

a. Date tests were conducted.

b. Name(s), location(s), and telephone number(s) of laboratory(ies) conducting the testing and evaluation.

c. Photographs and drawings illustrating the durability testing equipment and procedures used, as well as a summary of the specific procedures used.

d. Tensile test results for the product before durability testing (i.e., virgin material), and whether both virgin and durability test samples were taken from the same roll of material, or just from rolls within the same lot of material.

e. Polymer characteristics for the lot or roll of material actually tested before long-term exposure in the laboratory, including, for example, molecular weight and carboxyl end group content for PET, melt flow index and OIT for polyolefins, percent crystallinity, SEM photographs of fiber surface, etc.

Note 1: Percent crystallinity can be determined using Differential Scanning Calorimetry (DSC). An appropriate test method is ASTM D3417-99. By definition, crystallinity (X) is calculated as follows:

\[ X = \Delta H / \Delta H^o \text{ (times 100 for %)} \]

where: \( \Delta H \) is the latent heat under the DSC melt curve

\( \Delta H^o \) is the latent heat for a 100% crystalline polymer

Temperature scan should start 10° C below, continue through, and stop 10° C above the melt range. Recommended test parameters are as follows:
### Homo-Polymer Sample Size

<table>
<thead>
<tr>
<th>Homo-Polymer</th>
<th>Sample Size (mg)</th>
<th>Melt Range (° C)</th>
<th>Latent Heat, ΔH° (cal/gm)</th>
<th>DSC Scan Speed (° C/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HDPE</td>
<td>5</td>
<td>100-145</td>
<td>68.4</td>
<td>10</td>
</tr>
<tr>
<td>PP</td>
<td>7.5</td>
<td>100-165</td>
<td>45</td>
<td>10</td>
</tr>
<tr>
<td>PET</td>
<td>10</td>
<td>200-245</td>
<td>30</td>
<td>10</td>
</tr>
</tbody>
</table>

*Other values of sample size, melt range, and DSC scan speed can be used with justification.*

f. Tensile test results for specimens taken for each retrieval from the incubation chambers.

g. Tensile test results, including tensile strength, strain at peak load, and 5 percent secant or offset modulus, for both virgin material and degraded material should include individual test results for each specimen, typical load-strain curves which are representative of the specimens tested, including associated calibration data as necessary to interpret the curves (curves in which strain and load/unit width are already calculated are preferred), the average value for each sample, the coefficient of variation for each sample, and a description of any deviations from the standard tensile test procedures required by Appendix D.

h. A detailed description of the data characterization and extrapolation procedures used, including data plots illustrating these procedures and their theoretical basis.

i. Results of any chemical tests taken (e.g., OIT or HPOIT, molecular weight, product weight/unit area, etc.), and any scanning electron micrographs taken, to verify the significance of any degradation in strength observed.

j. Results of biological degradation testing, if performed.

k. RF_D, and a description of the method used to determine RF_D for the product.

5. Evaluation of Product Lines

If determining the long-term strengths for a product line, the data required under “General Product Information” must be obtained for each product. Product specific information for creep and durability must be obtained for at least one product in the product line to qualify the product line for Class 1 structures or aggressive environments, or in the case of Class 2 structures to allow the use of a total long-term strength reduction factor of less than 7 (see description of environment aggressiveness and Class 1 and Class 2 structures in “Determination Of Long-Term Geosynthetic Strength” later in this Standard Practice). Additional product specific information for creep and durability shall also be obtained for each product in the product line in accordance with Appendices B, C and D regarding use of long-term data for “similar” products. This data is to be used to determine long-term strengths for each product in the product line.

In general, product specific installation damage data must be obtained for each product in the line. However, it is permissible to obtain installation damage data for only some of the products in the product line if interpolation of the installation damage reduction factor between products is feasible. Interpolation of the product specific installation damage reduction factor RF_D between tested products can be based on the weight per unit area or undamaged tensile strength of each product, provided that the progression of weight per unit area or tensile strength as compared to the progression of RF_D for each tested product is consistent. For coated geogrids, the weight of coating placed on the fibers or yarns may influence the amount of installation damage obtained (Sprague, et al., 1999). In that case, the installation damage reduction factor may need to be correlated to the coating weight instead.
Determination of Long-term Geosynthetic Strength for Initial Product Acceptance

1. Calculation of Long-Term Strength

Reinforcement elements in MSE walls and reinforced slopes should be designed to have a durability to ensure a minimum design life of 75 years for permanent structures. For ultimate limit state conditions:

\[ T_{al} = \frac{T_{ult}}{RF} \]  

(1)

where:

- \( RF = RF_{ID} \times RF_{CR} \times RF_{D} \)  
- \( T_{al} \) = The long-term tensile strength that will not result in rupture of the reinforcement during the required design life, calculated on a load per unit of reinforcement width basis
- \( T_{ult} \) = the ultimate tensile strength (MARV) of the reinforcement determined from wide width tensile tests
- \( RF \) = a combined reduction factor to account for potential long-term degradation due to installation damage, creep, and chemical/biological aging
- \( RF_{ID} \) = a strength reduction factor to account for installation damage to the reinforcement
- \( RF_{CR} \) = a strength reduction factor to prevent long-term creep rupture of the reinforcement
- \( RF_{D} \) = a strength reduction factor to prevent rupture of the reinforcement due to chemical and biological degradation

See Appendices A through D for protocols to use to determine RF from product specific data. Unless otherwise indicated in the contract specifications for a given project, the design temperature used to determine RF and \( T_{al} \) from product specific data shall be assumed to be 20° C (68° F).

The value selected for \( T_{ult} \) is the minimum average roll value (MARV) for the product to account for statistical variance in the material strength. \( T_{ult} \) should be based on a wide width tensile strength (i.e., ASTM D4595 for geotextiles or ASTM D6637 for geogrids). Other sources of uncertainty and variability in the long-term strength include installation damage (Appendix A), creep extrapolation (Appendices B and C), and chemical degradation (Appendix D). It is assumed that the observed variability in the creep rupture envelope is 100% correlated with the short-term tensile strength, as the creep strength is typically directly proportional to the short-term tensile strength within a product line (see Appendix B and Note 7 in Appendix B if this is not the case). Therefore, the MARV of \( T_{ult} \) adequately takes into account that source of variability. For additional discussion of this issue, see Note 2 below.

**Note 2:** The product strength variability is not taken into account by using the creep limited strength, \( T_{l} \), directly or in normalizing \( T_{i} \) by \( T_{tot} \) (see Appendix B). \( T_{i} \) only accounts for extrapolation uncertainty. Furthermore, \( T_{tot} \) is specific to the lot of material used for the creep testing. Normalizing by \( T_{tot} \) makes the creep reduction factor \( RF_{CR} \) applicable to the rest of the product line, as creep strength is typically directly proportional to the ultimate tensile strength, within a product line. As shown below, it is not correct to normalize the creep strength \( T_{i} \) using \( T_{ult} \), the MARV of the tensile strength for the product, nor is it correct to use \( T_{l} \) directly in the numerator to calculate \( T_{al} \).

\[ RF_{CR} = \frac{T_{tot}}{T_{i}} \neq \frac{T_{ult}}{T_{i}} \quad \text{and} \quad T_{al} \neq \frac{T_{i}}{RF_{ID} \times RF_{D}} \]
In the former case, the creep strength is not indexed to the actual tensile strength of the material used in the creep testing, and since there is a 50% chance that \( T_{ult} \) will be less than or equal to \( T_{lot} \), using \( T_{ult} \) in this case would result in an unconservative determination of \( RF_{CR} \). In the latter case, where \( T_l \) is used directly as a creep reduced strength, the product strength variability is not taken into account, since \( T_l \) is really a mean creep strength. Hence, \( RF_{CR} \) must be determined as shown in Equation B.4-1 (see Appendix B), and the MARV must be used for \( T_{ult} \) when determining \( T_{al} \). Note that the use of the MARV for \( T_{ult} \) may not fully take into account the additional variability caused by installation damage. For the typical degree of installation damage observed in practice, this additional variability is minor and can be easily handled through the overall safety factor used in design of reinforced structures. For durability (\( RF_d \)), additional variability does not come into play if a default reduction factor is used. If a more refined durability analysis is performed, additional variability resulting from chemical degradation may need to be considered.

The type and amount of data to be obtained, and the approach used to determine the long-term design strength, will depend on the geosynthetic wall or reinforced slope class and the aggressiveness of the environment.

2. Wall or Slope Class

The class of a given geosynthetic structure will be identified in the contract specifications. A Class 1 geosynthetic wall or reinforced slope typically includes walls or slopes that support bridge abutments, buildings, critical utilities, or other facilities for which the consequences of poor performance or failure would be severe. Examples of severe consequences include serious personal injury, loss of life, or significant property damage. Cost and impact to the public if a poorly performing wall or slope must be repaired or replaced may also be considered in the determination of wall or slope class. In general, geosynthetic walls greater than 6 m (20 ft) in height and reinforced slopes greater than 9.2 m (30 ft) in height will be considered to be Class 1. All other geosynthetic walls and reinforced slopes will in general be considered to be Class 2.

3. Environment Aggressiveness

A nonaggressive environment is defined based on soil gradation, chemical properties of the environment, and site temperature. Normally, the backfill pH will be the key chemical property that will affect the chemical aggressiveness of the geosynthetic environment. The gradation of the backfill will normally be specified in accordance with WSDOT Standard Specifications Section 9-03.14(1) for geosynthetic walls, and Section 9-03.14(2) for reinforced slopes. In some cases, backfill meeting the requirements of Section 9-03.14(2) may also be used for walls. Soils coarser than these gradations will be considered to be aggressive due to the potential for the geosynthetic to have a high installation damage reduction factor when installed in a soil with a coarser gradation. In general, these materials can be characterized as having a maximum \( d_{50} \) size of 4.75 mm. The gradation requirements specified in Sections 9-03.14(1) and 9-03.14(2) are as follows:
### Sieve Size Section 9-03.14(1) Percent Passing Section 9-03.14(2) Percent Passing

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Section 9-03.14(1) Percent Passing</th>
<th>Section 9-03.14(2) Percent Passing</th>
</tr>
</thead>
<tbody>
<tr>
<td>31.5 mm (1.25 in.)</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>4.75 mm (#4 sieve)</td>
<td>50 - 90</td>
<td>-</td>
</tr>
<tr>
<td>0.425 mm (#40 sieve)</td>
<td>30 max.</td>
<td>50 max.</td>
</tr>
<tr>
<td>0.075 mm (#200 sieve)</td>
<td>7.0 max.</td>
<td>10 max.</td>
</tr>
</tbody>
</table>

Other backfill gradations may be considered at the discretion of the approval authority (e.g., WSDOT).

Regarding chemical properties of the environment surrounding the geosynthetic in the wall or slope, the pH shall be between 4.5 and 9 to be considered nonaggressive. This applies both in the reinforced backfill and at the back of the face of walls.

Regarding temperature, the effective design temperature at the wall or slope site shall be less than 30°C (85°F) for the environment to be considered nonaggressive. In the state of Washington, all wall and slope sites are anticipated to have an effective design temperature that is below 30°C.

For most wall and slope sites in the State of Washington, the environment will likely be nonaggressive. A possible exception to this is immediately behind a concrete wall face, where pH levels could possibly be elevated above a pH of 9. However, recent research has indicated that for well drained backfills, the pH adjacent to a concrete face stays below 9 in the long-term (Koerner, et al., 2001, Koerner, et al., 2002). In any case, the long-term strength determination must account for the environment at the face.

The wall or slope contract specifications will identify if the environment is anticipated to be aggressive and the reason for the aggressive environment designation (i.e., backfill gradation, site chemistry, or site temperature). If aggressive conditions are not identified in the contract specifications, the environment should be considered to be nonaggressive to determine the long-term strength.

### 4. Requirements for Class 1 Walls and Slopes to Determine $T_{al}$

$RF_{id}$ and $RF_{cr}$ shall be determined from product specific data for all geosynthetics used in Class 1 walls and slopes. See submission requirements for installation damage and creep data provided in this document. The product specific data for these reduction factors shall be interpreted/extrapolated in accordance with Appendices A, B, and C. $RF_{d}$ shall be determined from long-term product specific data, or a default value may be used as described below. See submission requirements for durability data provided herein. Long-term product specific data for $RF_{d}$ should be interpreted in accordance with Appendix D. If adequate long-term durability data is not available, a default reduction factor for $RF_{d}$ may be used if the environment is nonaggressive and if the product meets the minimum polymer and physical property requirements provided in Table 1. In this case, a default value for $RF_{d}$ of 1.3 may be used for PET, HDPE, and PP geosynthetics.
Note 3: The default value for RF_D of 1.3, which can be used for products that meet the minimum property requirements in Table 1, was determined based on FHWA (1997) and Elias, et. al. (1997) and in consideration of the relatively cool climate which exists in the state of Washington, where effective design temperatures are always less than 20° C (68° F) and are likely to be on the order of 10° C (50° F) or less. A higher default value of 1.5 for products which meet the property requirements in Table 1 may be desirable for more temperate climates which still meet the requirements for a nonaggressive environment, especially to address polyolefin oxidative degradation, as the potential for this type of degradation, even for products which meet the property requirements in Table 1, becomes more uncertain at higher temperatures due to the lack of protocols which can accurately identify the amount or effectiveness of end use antioxidants present. The UV resistance criteria provided in Table 1 only provides a rough indication of the effectiveness of end use antioxidants in polyolefins (see additional commentary following Table 1).

If the environment is identified as aggressive due to the chemical regime or due to temperature, or if the geosynthetic product does not meet the requirements in Table 1, default reduction factors may not be used for RF_D. For chemically aggressive or elevated temperature environments, RF_D must be determined based on long-term product specific data for an environment that is as or more aggressive than the project specific environment in question. Aggressive environments need to be addressed in the product submittal only if specifically requested by the contracting agency or the geosynthetic supplier. Once the appropriate reduction factors are established, the long-term geosynthetic strength is determined using Equations 1 and 2, or as determined in Note 7 of Appendix B.

5. Requirements for Class 2 Walls and Slopes to Determine T_al

The strength reduction factors RF_D, RF_CR, and RF_D may be determined based on product specific data as described for Class 1 walls and slopes. If long-term product specific data is not available, the environment is nonaggressive, and the product meets the minimum requirements provided in Table 1, a default value of 7 may be used for RF to determine the long-term strength of the product in accordance with Equations 1 and 2.

6. Minimum Polymer and Physical Property Requirements to Allow Use of Default Reduction Factors for RF and RF_D in Nonaggressive Environments

If a default reduction factor is to be used, geosynthetic products that are likely to have good resistance to installation stresses and to long-term chemical degradation are required to minimize the risk of significant long-term degradation. The physical and polymer material requirements provided in Table 1 must be met if detailed product specific data as described in Appendices A, B, C and/or D is not obtained. Polymer materials not meeting the requirements in Table 1 could be used if detailed product specific data extrapolated to the design life intended for the structure (see Appendices A, B, C and D) is provided.
Table 1
Minimum Requirements for Geosynthetic Products
to Allow Use of Default Reduction Factor for Long-Term Degradation.

<table>
<thead>
<tr>
<th>Polymer Type</th>
<th>Property</th>
<th>Test Method</th>
<th>Criteria to Allow Use of Default RF*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP and HDPE</td>
<td>UV Oxidation Resistance</td>
<td>ASTM D4355</td>
<td>Min. 70% strength retained after 500 hrs in weatherometer</td>
</tr>
<tr>
<td>PET</td>
<td>UV Oxidation Resistance</td>
<td>ASTM D4355</td>
<td>Min. 50% strength retained after 500 hrs in weatherometer if geosynthetic will be immediately buried, 70% if left exposed for more than one week.</td>
</tr>
<tr>
<td>PP and HDPE</td>
<td>Thermo- Oxidation Resistance</td>
<td>ENV ISO 13438:1999, Method A (PP) or B (HDPE)</td>
<td>Min. 50% strength retained after 28 days (PP) or 56 days (HDPE)</td>
</tr>
<tr>
<td>PET</td>
<td>Hydrolysis Resistance</td>
<td>Inherent Viscosity Method (ASTM D4603 and GRI Test Method GG8), or Determine Directly Using Gel Permeation Chromatography</td>
<td>Min. Number Average Molecular Weight of 25,000</td>
</tr>
<tr>
<td>PET</td>
<td>Hydrolysis Resistance</td>
<td>GRI Test Method GG7</td>
<td>Max. Carboxyl End Group Content of 30</td>
</tr>
<tr>
<td>All Polymers</td>
<td>Survivability</td>
<td>¹Weight per Unit Area (ASTM D5261)</td>
<td>¹Min. 270 g/m²</td>
</tr>
<tr>
<td>All Polymers</td>
<td>% Post-Consumer Recycled Material by Weight</td>
<td>Certification of Materials Used</td>
<td>Maximum of 0%</td>
</tr>
</tbody>
</table>

*Polymers not meeting these requirements may be used if product specific test results obtained and analyzed in accordance with Appendices A, B, C, and D are provided.

¹Alternatively, a default RF₀ = 1.3 may be used if product specific installation damage testing is performed and it is determined that RF₀ ≤ 1.5 or less, and if the other requirements in Table 1 are met.

**Note 4:** The requirements provided in Table 1 utilize currently available index tests and are consistent with current AASHTO design specifications (AASHTO, 2004, 2002), with the exception of the oven aging test, which is a new requirement. These index tests can provide an approximate measure of relative resistance to long-term chemical degradation of geosynthetics. Values selected as “minimum” criteria to allow use without additional long-term testing are based on values for such properties reported in the literature. These values are considered indicative of good long-term performance or represent a readily available current standard within the industry that signifies that a product has been enhanced for long-term environmental exposure.

Though UV resistance (i.e., photo-oxidation resistance) is not a direct indicator of thermo-oxidation resistance for polypropylene and polyethylene, both photo-oxidation and thermo-oxidation are oxidation reactions, and many UV inhibitors also provide at least some long-term resistance to thermo-oxidation (Van Zanten, 1986). Regarding polyester requirements, maximum resistance to strength losses due to hydrolysis can be obtained by formulating to high molecular weights (> 25,000) and low (i.e., < 30) Carboxyl End Group numbers (Risseeuw and Schmidt, 1990; FHWA., 1997; and Elias, et. al., 1997).
Minimum weight/area requirements are based on the results of numerous exhumations of geosynthetics, in which it was determined that installation damage was minimal for products with a minimum of weight of 270 g/m² (8 oz/yd²) (Koerner and Koerner, 1990; Allen, 1991). This roughly corresponds to a Class 1 geotextile as specified in AASHTO M-288.

There is little long-term history or even laboratory data regarding the durability of geosynthetics containing a significant percentage of recycled material. Therefore, their potential long-term performance is unknown, and it is recommended that long-term data be obtained for products with significant recycled material to verify their performance before using them.

Quality Assurance Requirements for Products that have been Through Initial Acceptance

1. Data Verification Requirements

The following information about each product shall be submitted for verification purposes:

a. Geosynthetic type and structure.
b. Spacing and dimensions of geogrid elements.
c. Polymer(s) used for fibers, ribs, etc.
d. Polymer(s) used for coating, if present.
e. Roll size (length, width, and area).
f. Typical lot size.
g. Polymer source(s) used for product.
h. For HDPE and PP, primary resin ASTM type, class, grade, and category (for HDPE use ASTM D-1248, and for PP use ASTM D-4101).
j. % post-consumer recycled material by weight.
k. Minimum weight per unit area for product (ASTM D5261).
l. MARV for ultimate wide width tensile strength (ASTD D459 or ASTM D6637).

2. Quality Assurance (QA) Testing Approach

Results from index and performance tests will be compared to baseline index or performance test results obtained for initial product acceptance purposes. If the QA test results are within acceptable tolerances relative to the baseline results, the acceptance status of the product or product line will be maintained (e.g., the product will continue to be listed in the QPL). Re-testing must be done if there is any change in the product. If changes in the product identified through product data verification as described in part 1 above or identified through other means are such that the validity of the last complete assessment for initial acceptance is too questionable, WSDOT reserves the right to require a complete assessment of the product or product line in accordance with this Standard Practice instead of just a QA evaluation to maintain acceptance status.

3. Quality Assurance (QA) Sampling

All materials and/or products to be tested will be furnished by the manufacturer/supplier at no cost to the review/approval authority. Samples will be selected for testing by Department of Transportation personnel or designated parties. As a minimum, the following shall be obtained:

- a geosynthetic product sample of sufficient size to accommodate all of the specified testing;
- information showing the manufacturer’s name and description of product: (style, brand name, etc.);
• product roll and lot number;
• a sample of the polymer component(s) in sufficient quantity to conduct the specified polymer tests.

All samples for the specified QA testing shall be from the same roll of material for each product tested.

4. Quality Assurance (QA) Testing

Short-term ultimate tensile strength test results, and QA test results to verify the correctness of RF<sub>ID</sub>, RF<sub>CR</sub>, and RF<sub>D</sub> determined from initial product acceptance testing, shall be obtained.

Short-term tensile strength shall be determined in accordance with ASTM D4595 for geotextiles and ASTM D6637 for geogrids. QA testing required to verify the correctness of RF<sub>ID</sub>, RF<sub>CR</sub>, and RF<sub>D</sub> determined from initial product acceptance testing is as follows:

A. Installation Damage Testing

For installation damage evaluation, a field exposure trial conducted in accordance with Appendix A shall be conducted for the lightest weight material in the product line tested previously for initial product acceptance using the soil with a d<sub>50</sub> size which is equal to or larger than the d<sub>50</sub> size for WSDOT Gravel Borrow, 4.75 mm, or other d<sub>50</sub> size as determined by the approval authority that is consistent with the d<sub>50</sub> size used for initial product acceptance. Alternatively, reduced scale laboratory installation damage tests conducted in accordance with ISO/DIS 10722-1 may be used. In this case, these laboratory installation damage tests must also be conducted during initial product acceptance testing to establish a baseline value. The ultimate tensile strength of the lot or roll of material used in the installation damage testing obtained in accordance with ASTM D4595 or ASTM D6637 (or ISO 10319 if ISO/DIS 10722-1 is used) shall be obtained to normalize the installation damage test results in accordance with Appendix A. If it was determined during the initial product acceptance testing, for coated geogrids, that the installation damage factor was not a function of product weight or tensile strength, the coating weight shall also be evaluated. In this case, the mass/unit area of the sample tested shall be determined in accordance with ASTM D5261. The coating weight can then be established using the lot specific mass/unit area of the uncoated product from product manufacturer Quality Control (QC) data. The information required in part 2 of “Data Requirements for Initial Product Acceptance” as it applies to the QA testing shall be obtained and included in the test report for this QA testing.

B. Creep Testing

For creep rupture evaluation, a minimum of three creep-rupture points shall be obtained using SIM (GRI-GS10) or conventional ASTM D5262 tests (for which elevated test temperatures may be employed to accelerate creep – see Appendix B) at a load level established at the time of initial product acceptance testing that corresponds to a minimum rupture time of 100,000 hours at the reference temperature. If elevated temperature conventional creep testing using ASTM D5262 is performed, the shift factors obtained from the conventional creep testing for the temperatures used in the QA testing conducted for initial product acceptance shall be used to extrapolate the test data to the reference temperature. A fourth SIM test (or conventional ASTM D5262 test conducted at the reference temperature) shall be performed at a load level established at the time of initial product acceptance testing that corresponds to a minimum rupture time of 500 hours at the reference temperature. Note that if initial product acceptance was based on Appendix C (creep strain testing), creep strain measurements must be obtained, and the load levels selected for the QA creep testing should be equal to the load level that results in reaching a specified strain using the creep data used to establish the initial product acceptance envelope (see Appendix C, Section C.2.2) at 500 hours (one test) and 50,000
hours (three tests), at the reference temperature. The strain level used for this purpose shall preferably be 5 to 10% or more, and be as close to the instability limit strain as possible while catching as many of the creep curves as possible. See Section 5(d) for additional explanation.

For creep stiffness evaluation, two additional SIM tests (or ASTM D5262 tests after time shifting if elevated temperature is used) shall be conducted at the load level that corresponds to 1,000 hours on the initial product acceptance creep envelope (i.e., load level versus the logarithm of time to reach the specified strain) derived at 2% strain.

If SIM is used for this creep rupture testing, it shall have been demonstrated for the initial acceptance testing that the reduced specimen width typically used for SIM testing does not have a significant effect on the creep rupture results, and provided that the validity of SIM for the product through comparison of SIM data with “conventional” creep rupture data was established for the initial product acceptance testing.

The ultimate tensile strength of the lot or roll of material used in the creep testing obtained in accordance with ASTM D4595 or ASTM D6637 shall be obtained to normalize the creep rupture loads in accordance with Appendix B or C. The information required in part 3 of “Data Requirements for Initial Product Acceptance” as it applies to the QA testing shall be obtained and included in the test report for this QA testing.

**Note 5:** If “conventional” creep testing is performed for QA purposes, it is assumed that the product has not changed relative to what was tested for initial product acceptance purposes, thereby allowing the assumption to be made that the shift factors obtained through the initial product acceptance testing are valid for the QA testing. Requiring new “conventional” creep test shift factors to be re-established would result in the need to fully repeat the test program for the initial product acceptance, which would not be practical for QA purposes. Regarding the fourth creep test data point, the requirement to use only data obtained at the reference temperature if “conventional” creep testing is performed provides a second check that eliminates the need for this shift factor assumption and any inaccuracies associated with that assumption.

C. Durability Testing

If only index durability testing was conducted to allow use of a default value for RF₀ for the initial product acceptance testing, only index durability testing need be conducted for QA purposes. In this case, durability testing for QA purposes shall consist of the determination of molecular weight based on GRI-GG7 and carboxyl end group content based on GRI-GG8 for polyesters, UV resistance based on ASTM D4355 for polyolefins and PET’s), and an oven aging exposure test per ENV ISO 13438:1999 for polyolefin geosynthetics. Regarding the oven aging test, control and post-exposure specimens shall be tested for tensile properties (ASTM D4595 or ASTM D6637). The results of this oven aging testing will be used only to compare a product with itself, and to meet the minimum requirements in Table 1. In addition, geogrid brittleness shall be evaluated per WSDOT Test Method T 926.

If long-term performance durability testing was conducted to justify the use of a lower RF₀ or to justify use in aggressive environments for initial product acceptance, a minimum of five specimens shall be exposed to the most aggressive environment used in the initial product acceptance testing at the highest temperature tested, for a minimum of 2,000 hours. These specimens, and unexposed specimens from the same roll of material, shall be tested for tensile properties (ASTM D4595 or ASTM D6637). In addition, for polyolefins, either oxidative induction time per ASTM D 3895 or high pressure oxidative induction time per ASTM D 5885 shall be conducted for each specimen tested (before and after exposure), and for PET’s, molecular weight (ASTM D4603 and GRI:GG8) and specimen weight per unit area (ASTM D5261) shall be conducted for each specimen tested (before and after exposure).
5. Quality Assurance (QA) Criteria for Comparison to Initial Product Acceptance Test Results

The acceptability of the QA test results to allow a product or product line to maintain its prior acceptance status is established based on the statistical significance, or lack thereof, of the difference between the QA test results and the initial product acceptance test results. The criteria and methods for determining the statistical significance between the QA and initial product acceptance test results are as follows:

A. Short-term Index Tensile Testing

For wide width tensile strength, the mean of the test results for the sample for each product tested shall be greater than or equal to the MARV reported for the product.

B. Installation Damage Testing

If the mean of the average strength of the sample after damage as a percent of the undamaged strength is less than the average value obtained for the same product and condition during the product acceptance phase, the maximum difference between the two means shall be no greater that what is defined as statistically insignificant based on a one-sided student-t distribution at a level of significance of 0.05. In this case, t is determined as follows:

\[
t_{\alpha/2,n_1+n_2-2} = \frac{(\bar{P}_1 - \bar{P}_2) - \delta}{\sqrt{(n_1-1)s_1^2 + (n_2-1)s_2^2}} \sqrt{\frac{n_1n_2}{(n_1+n_2)}}
\]

where,

\[
t_{\alpha/2,n_1+n_2-2} = \text{value of the } t\text{-distribution for the installation damage samples}
\]

\[\bar{P}_1 = \text{the mean of the strength retained after installation damage (i.e., } T_{\text{dam}}/T_{\text{lot}}) \text{ obtained for initial product acceptance}\]

\[\bar{P}_2 = \text{the mean of the strength retained after installation damage (i.e., } T_{\text{dam}}/T_{\text{lot}}) \text{ obtained for QA testing}\]

\[\delta = \text{the difference in the means for the populations corresponding to the sample means } \bar{P}_1 \text{ and } \bar{P}_2 \text{ (assumed equal to zero for this test)}\]

\[s_1 = \text{the standard deviation corresponding to } \bar{P}_1\]

\[s_2 = \text{the standard deviation corresponding to } \bar{P}_2\]

\[n_1 = \text{the number of data points corresponding to } \bar{P}_1\]

\[n_2 = \text{the number of data points corresponding to } \bar{P}_2\]

\[t_{\alpha/2,n_1+n_2-2} \text{ calculated using Equation 3 shall be no greater than } t \text{ determined from the applicable Student } t\text{-table (or from the Microsoft EXCEL function TINV(}\alpha,n-2)) \text{ at } \alpha = 0.05 \text{ and } n_1+n_2-2 \text{ degrees of freedom. If this is not true, the difference between } \bar{P}_1 \text{ and } \bar{P}_2 \text{ is determined to be statistically significant, and } \bar{P}_1 > \bar{P}_2, \text{ two additional samples from the same installation condition shall be tested and } \bar{P}_2 \text{ recalculated and statistically compared to } \bar{P}_1. \text{ If the QA test results are still too low, a full installation damage study for initial product acceptance must be completed in accordance with Appendix A, and new values of RF_{ID} \text{ established.} \]
C. Creep Rupture Testing for Prediction of Creep Limit

For creep evaluation, the four creep-rupture points, one at a load level that results in an approximate rupture time, after time shifting, of 500 hours and three at a load level that results in an approximate rupture time, after time shifting, of 100,000 hours on the rupture envelope obtained for initial product acceptance purposes shall be compared to the creep data obtained for initial product acceptance purposes. The log of the rupture time for each of these four rupture points shall be equal to or greater than the 95% lower prediction limit of the variable, log time, established by the Student’s t test of the original product acceptance data set.

The prediction limit for the regression performed for initial product acceptance is given by (Wadsworth, 1998):

\[
\log t_L = \log t_{reg} - \left[ t_{t_{\alpha/2,n-2}} \left( 1 + \frac{1}{n} + \sum \left( \frac{P_i - P}{\bar{P}} \right)^2 \right) \right] \times \sigma \tag{4}
\]

and

\[
\sigma = \sqrt{\frac{\sum \left[ \log t_i - \log t_{\bar{P}} \right]^2 - \sum \left[ \frac{P_i - P}{\bar{P}} \right] \left[ \log t_i - \log t_{\bar{P}} \right]}{n-2}} \tag{5}
\]

where:

- \( \log t_L \) = lower bound prediction limit
- \( t_{reg} \) = time corresponding to the load level from the initial product acceptance creep rupture envelope at which QA creep tests were performed (e.g., at 500 and 100,000 hrs after time shifting)
- \( t_{t_{\alpha/2,n-2}} \) = value of the t distribution determined from applicable Student t table (or from the Microsoft EXCEL function TINV(\( \alpha/n-2 \)) at \( \alpha/2 = 0.05 \) and \( n-2 \) degrees of freedom (this corresponds to the 95% one-sided prediction limit)
- \( n \) = the number of rupture or allowable run-out points in the original test sample (i.e., for initial product acceptance)
- \( P \) = load level obtained at \( t_{reg} \) from the regression line developed from the initial product acceptance testing
- \( \bar{P} \) = the mean rupture load level for the original test sample (i.e., all rupture or run-out points used in the regression to establish the rupture envelope for initial product acceptance)
- \( P_i \) = the rupture load level of the i'th point for the rupture points used in the regression for establishing the rupture envelope for initial product acceptance
- \( \log t_i \) = the mean of the log of the rupture time for the original test sample (i.e., all rupture or run-out points used in the regression to establish the rupture envelope for initial product acceptance)
- \( t_i \) = the rupture time of the i'th point for the rupture points used in the regression for establishing the rupture envelope for initial product acceptance
The comparison between the QA test results and the initial product acceptance test results is illustrated conceptually in Figure 1. Once log $t_L$ has been determined at each specified load level, compare this value to the log rupture time (i.e., log $t_{QA}$) obtained for each QA creep rupture test at the specified load level (e.g., 500 and 100,000 hours). If log $t_{QA} < \log t_L$ for any of the QA creep rupture test results, perform two additional tests at the load level $P$ for the specified $t_{reg}$ where this QA criteria was not met and compare those results to log $t_L$. If for these two additional tests this criterion is not met, perform adequate additional creep rupture testing to establish a new rupture envelope for the product in accordance with initial product acceptance requirements (Appendix B). This new rupture envelope will form the baseline for any future QA testing.

![Figure 1. Conceptual illustration of the comparison of QA creep rupture test results to initial product acceptance creep rupture test results.](image)

**D. Creep Strain Testing for Prediction of Creep Limit**

The comparison between the creep data obtained for the initial product acceptance testing and the QA creep data shall be performed at a specified strain. The specified strain will depend on the strains observed in all of the creep tests (initial product acceptance and QA). Select a strain that will intercept all of the creep curves as much as possible. Preferably, the strain level should be approximately 5 to 10% or more, and as close to the instability limit strain as possible. Where the selected strain level intersects each creep curve, determine the time required to reach the specified strain. Plot the load level as a function of the logarithm of time to reach the specified strain for the initial product acceptance data, and perform a regression for this data set. The log times to the specified strain level for the QA creep data shall be determined at a load level that corresponds to 500 hours and 50,000 hours on the initial product acceptance creep envelope. This is illustrated conceptually in Figure 2. The log of the time to reach the same specified strain for each of the four QA creep data points shall be equal to or greater than the 95% lower prediction limit of the variable, log time, established by the Student’s t test of the original product acceptance data set, using Equations 4 and 5 (see part “c” above).
Once log \( t_L \) has been determined at each specified load level, compare this value to the log time to reach the specified strain (i.e., log \( t_{QA} \)) obtained for each QA creep test at the specified load level (e.g., 500 and 50,000 hours). If log \( t_{QA} < \) log \( t_L \) for any of the QA creep rupture test results, perform two additional tests at the load level \( P \) for the specified \( t_{reg} \) where this QA criteria was not met and compare those results to log \( t_L \). If for these two additional tests this criterion is not met, perform adequate additional creep testing to establish a new creep stiffness curve for the product in accordance with initial product acceptance requirements (Appendix C). This new creep stiffness curve will form the baseline for any future QA testing.

Figure 2. Conceptual illustration of the comparison of QA creep strain test results to initial product acceptance creep strain test results (a) creep strain curves, and (b) envelope of time to the specified strain.
E. **Assessment of the Creep Stiffness at Low Strain**

The comparison between the creep data obtained for the initial product acceptance testing and the QA creep data shall be performed at a specified strain, in this case typically 2%. Where the selected strain level intersects each creep curve, determine the time required to reach the specified strain. Plot the load level as a function of the logarithm of time to reach the specified strain for the initial product acceptance data, and perform a regression for this data set. The log times to the specified strain level for the QA creep data shall be determined at a load level that corresponds to 1,000 hours on the initial product acceptance creep envelope. The time to reach the same specified strain for each of the two QA creep data points shall be equal to or greater than the 95% lower prediction limit of the variable, log time, established by the Student’s $t$ test of the original product acceptance data set, using Equations 4 and 5 (see part “c” above).

Once log $t_{L}$ has been determined at the specified load level, compare this value to the log time to reach the specified strain (i.e., log $t_{QA}$) obtained for each QA creep test at the specified load level (e.g., 1,000 hours). If log $t_{QA} < log t_{L}$ for any of the QA creep rupture test results, perform two additional tests at the same load level $P$ for the specified and compare those results to log $t_{L}$. If for these two additional tests this criterion is not met, perform adequate additional creep testing to establish a new low strain creep stiffness value for the product in accordance with initial product acceptance requirements (Appendix C). This new low strain creep stiffness value will form the baseline for any future QA testing.

F. **Durability Testing**

For UV resistance (all polymers), molecular weight and CEG (PET only), and oven aging (PP and HDPE), the QA test results shall meet the minimum requirements provided in Table 1. For the oven aging tests (polyolefins only), compare the tensile strength retained (i.e., strength after oven exposure divided by the strength of the control specimens) to the strength observed during initial product acceptance testing. The maximum difference between the values of the changes shall be no greater that what is defined as statistically insignificant based on a one-sided student-$t$ distribution at a level of significance of 0.05, as determined using Equation 3. In this case, $P_1$ and $P_2$ are defined as the strength retained after oven aging.

$t_{\alpha/2,n_1+n_2-2}$ calculated using Equation 3 shall be no greater than $t$ determined from the applicable Student $t$ table (or from the Microsoft EXCEL function TINV($\alpha$,n-2)) at $\alpha/2 = 0.05$ and $n_1+n_2-2$ degrees of freedom. If this is not true, and the difference between $P_1$ and $P_2$ is determined to be statistically significant, and $P_1 > P_2$, two additional samples from the same installation condition shall be tested and recalculated and statistically compared to $P_1$. If the QA test results are still unacceptable, or if the product loses more than 50% of its tensile strength during the QA test, a more complete investigation performed in accordance with Appendix D shall be performed.

If long-term performance durability testing was conducted to justify the use of a lower RF$_D$ or to justify use in aggressive environments for initial product acceptance, the statistical methodology and criteria provided above for index oven aging (i.e., that there be no statistically significant difference between the initial product acceptance test results and the QA test results at a level of significance of 0.05) shall be applied to the oxidation or hydrolysis performance test results at the maximum exposure time and environmental conditions used for the QA testing.
References


WSDOT Test Method No. 925, Appendix A

Product Specific Testing and Data Interpretation Requirements to Determine $RF_{ID}$ for Geosynthetic Reinforcements

The effect of installation damage on geosynthetic reinforcement strength and deformation shall be determined from the results of full scale installation damage tests in accordance with ASTM D5818, except as augmented herein:

1. The locations of specimens within the sample to be removed for testing after installation and exhumation should be predetermined before installation through the use of a template. The specimens should be large enough to be used for wide width testing (ASTM D4595 for geotextiles or ASTM D6637 for geogrids). A minimum of 20 specimen locations should be identified using a template for each installation condition evaluated. No specimen should be located closer than 150 mm (6 in.) from the edge of the sample. The locations of the specimens should be evenly distributed throughout the sample. Each specimen should be consecutively numbered before installation.

2. Place and compact 150 mm (6 in.) or more of soil (same soil as used to cover the geosynthetic) on a flat, level, relatively incompressible subgrade. The compacted layer shall simulate the roughness and compressibility of the backfill conditions in which the geosynthetic layer is likely to be placed in full scale structures.

3. Place the geosynthetic on top of the compacted soil pad. The geosynthetic shall be pulled taught with no wrinkles or folds. It may be necessary to pin the corners of the geosynthetic to maintain its position as soil is placed over the geosynthetic.

4. Place and compact 150 to 200 mm (6 to 8 in.) of backfill material (or other previously agreed upon depth to simulate actual installation conditions likely to be encountered) over the geosynthetic using the type of spreading and compacting equipment that is likely to be used in full scale structures. The backfill should be compacted in accordance with WSDOT Standard Specifications Section 2-03.3(14)C as described below, or to another compaction standard typically used for geosynthetic reinforced structures if agreed upon in advance by the approval authority (e.g., 95% of Modified Proctor per ASTM D1557). WSDOT Standard Specifications Section 2-03.3(14)C requires a soil moisture content that is within 3% of optimum, and compaction to 95% of maximum density per WSDOT test procedures (this is approximately equivalent to 90 to 95% of maximum density as determined per ASTM D1557).

5. The sample shall be removed from the compacted fill in a way that minimizes damage to the sample caused by the excavation process. The sample removal process is generally described in ASTM D5818. In addition to the removal methods described therein, a lifting plate may be placed below the compacted soil pad below the geosynthetic layer to facilitate easy removal of the soil above the geosynthetic.

6. The first nine prenumbered specimens identified on the exhumed sample shall be selected for testing. If any of these specimens were damaged due to the exhumation process, that specimen(s) shall be skipped, and the next consecutively numbered specimen(s) shall be selected for testing. If the coefficient of variation for the tensile test results of these first nine prenumbered specimens is greater than 5%, the required number of specimens shall be recomputed using the one-sided student $t$ distribution as required by ASTM D4595 or ASTM D6637. The additional specimens shall be selected from the next consecutively numbered specimens.
7. Samples subjected to installation damage shall be tested for tensile strength and deformation characteristics in accordance with ASTM D4595, or ASTM D6637. The number of specimens tested should be in accordance with ASTM D4595 or D6637. Single rib tests such as GRI: GG1 shall not be used for installation damage evaluation, as it is difficult to assess the effect of severed ribs on the strength and stiffness of damaged materials. Test results from damaged specimens shall be compared to tensile test results obtained from undamaged (i.e., not exposed to installation conditions) specimens taken from the same lot, and preferably the same roll, of material as the damaged specimens.

8. The installation damage reduction factor RF\textsubscript{ID} is then determined as follows:

\[
RF_{ID} = \frac{T_{lot}}{T_{dam}} \tag{A-1}
\]

where, \(T_{lot}\) is the average lot specific tensile strength before exposure to installation, and \(T_{dam}\) is the average lot specific tensile strength after installation. In no case should RF\textsubscript{ID} be less than 1.1.

9. To select an appropriate reduction factor for design, the project site installation conditions must be related to the installation test conditions. To relate the installation damage test conditions to the actual site conditions, primary consideration will be given to the backfill characteristics (d\textsubscript{50} particle size, potential for oversize material, particle angularity, and overall gradation), and to a lesser degree the method of spreading the backfill over the geosynthetic, the type of compaction equipment, and initial backfill lift thickness over the geosynthetic, provided that the initial lift thickness is 150 mm (6 in.) or more. The actual installation conditions used in the test must be clearly stated in the test report, specifically identifying any deviations from typical geosynthetic reinforcement installation practices in full scale structures, and the impact those deviations may have on the values of RF\textsubscript{ID} determined. Typical installation conditions, for the purposes of this protocol, are generally described as follows:

- The geosynthetic is placed on a subgrade prepared as described in step 2.
- Backfill material is placed and spread on the geosynthetic using full scale spreading and compaction equipment commonly used in wall or reinforced slope construction (e.g., loaded dump trucks, dozers such as a D-6 or larger, etc.).
- Backfill is compacted using a full scale vibratory drum roller (i.e., one that someone can drive, not a smaller walk behind unit typically used to compact near the face of a wall to prevent distortion of facing elements during compaction).

If it is not possible to conduct the installation damage test in a way that fully simulates these “typical” installation conditions (i.e., installation conditions in the test are less severe than the conditions described above, for example, use lighter equipment, less movement of soil over the geosynthetic as the backfill is spread, etc.), data must be provided that demonstrates the effect the use of these less severe installation conditions would have on RF\textsubscript{ID}.

10. It is recommended that installation damage data be obtained for more than one backfill gradation to permit interpolation as needed to match the specific soil to be used as reinforced soil backfill. For WSDOT projects, at least one of the backfill soils tested should meet the requirements of Standard Specifications Sections 9-03.14(1) and 9-03.14(2). These materials will generally result in a d\textsubscript{50} size of 0.425 mm (#40 sieve) to 4.75 mm (#4 sieve). As a minimum, one installation damage test using backfill with a d\textsubscript{50} of approximately 4.75 mm
(#4 sieve) or greater should be conducted. At the discretion of the approval authority, if another backfill specification is selected for use, the installation damage testing should be conducted on at least one backfill soil meeting the gradation requirements for that backfill, or on at least two soils that bracket the desired gradation closely enough to permit interpolation between the soils.

11. Values of RF_{ID} may be estimated for a specified soil gradation using interpolation as illustrated in Figure A-1. The d_{50} size of the soil has commonly been used for interpolating between soil backfills for determination of RF_{ID}. Other combinations of soil particle size and factors that account for soil angularity may be considered for this correlation and interpolation procedure per mutual agreement between the geosynthetic manufacturer and the approval authority.

![Figure A-1. Interpolation of RF_{ID} for a soil with d_{50} = 2 mm from installation damage data obtained for soils with d_{50} = 0.02, 0.5 and 10 mm (Note: 1in. = 0.0254 mm).](attachment:image.png)

12. Not all products within a product line need to be tested for installation damage. As a minimum, the weakest (lightest) and strongest (heaviest) products within the product line, plus at least one additional intermediate strength (weight) product should be used to characterize RF_{ID} for the product line (note: if the strongest product within the product line is not tested, all heavier products will be assumed to have the same value of RF_{ID} as the heaviest product tested). An example of this interpolation procedure is provided in Figure 2. For coated polyester geogrids, the coating thickness or coating mass per unit area relative to the mass per unit area of the product should be considered for the purpose of correlating RF_{ID} between products rather than product unit weight or tensile strength alone. It is acceptable to obtain the coating mass/unit area through the use of manufacturer Quality Control (QC) data on the lot specific mass/unit area of the uncoated material (i.e., the weight of the “greige-good”), subtracting that mass/unit area from the total mass/unit area of the finished product. Therefore, the total mass per unit area of the sample used in the installation damage testing should be obtained in accordance with ASTM D5261.
Fig. A-2. Interpolation of $R_{F_{ID}}$ from damage measurements on products from the same line but with different weights. For a product of weight 300 g/m², $R_{F_{ID}} = 1.42$ (Note: 1 oz/yd² = 0.0295 gm/m²).
WSDOT Test Method No. 925, Appendix B

Creep Rupture Testing and Extrapolation Procedures

The effect of long-term load/stress on geosynthetic reinforcement strength and deformation characteristics shall be determined from the results of product specific, controlled, long-term laboratory creep tests conducted for a range of load levels and durations in accordance with ASTM D5262 adequate for extrapolation purposes to the desired design life, carried out to rupture of the geosynthetic when possible. Creep testing in accordance with ASTM D5262, but carried out to rupture where feasible, is described herein as the “conventional method.” A limited number of conventional creep tests may be supplemented and extended to longer creep rupture times using GRI-GS10 (Stepped Isothermal Method, or SIM) as described in this appendix. Specimens should be tested in the direction in which the load will be applied in use. Test results shall be extrapolated to the required structure design life. Based on the extrapolated test results, for ultimate limit state design, determine the highest load level, designated T₁, which precludes both ductile and brittle creep rupture within the required lifetime. T₁ should be determined at the required design life and at the design site temperature.

In both cases, unless otherwise specified or mutually agreed upon by the geosynthetic supplier, the testing laboratory, and the owner, a baseline testing temperature of 68°F (20°C) shall be used for this testing. Higher test temperatures shall be considered as elevated temperatures to be used for the purpose of time extrapolation. ASTM D5262 requires that the testing temperature be maintained at ±3.6°F (2°C). For some polymers, this degree of variance could significantly affect the accuracy of the shift factors and extrapolations determined in accordance with this appendix. For polymers that are relatively sensitive to temperature variations, this issue should be considered when extrapolating creep data using time-temperature superposition techniques, or minimized by using a tighter temperature tolerance.

Note that the procedures provided in this appendix are for in-air creep rupture testing. The procedures provided herein can likely also be applied to creep tests conducted confined in-soil. However, there is little information available, at present, on which to confirm that these procedures can be applied to in-soil creep data without modification. Therefore, procedures for use and interpretation of in-soil creep testing in compliance with this standard practice are not provided.

Note 1: Creep results in time dependent deformation that may continue to occur as long as the reinforcement is loaded. At low to intermediate load levels, depending on the polymer type, the creep rate will continue to decrease with time and may eventually stabilize, at least within the ability to measure creep. At higher load levels, creep will continue until rupture occurs.

In general, two types of creep tests are conducted: stress or creep rupture, and creep strain. Creep strains do not have to be monitored in creep rupture testing (strain measurement in this type of test is recommended, however), though creep strain tests can be carried to rupture. Rupture data is necessary if the creep reduction factor for ultimate limit state conditions, $RF_{CR}$, is to be determined. Creep rupture test results, if properly accelerated and extrapolated, can also be used to investigate the effects of stress cracking and the potential for a ductile to brittle transition to occur. This transition to brittle behavior, if it occurs, can cause a geosynthetic to fail in creep at lower loads and strains than anticipated from evaluating only creep strain and rate data.

Single ribs for geogrids or yarns for woven geotextiles may be used for creep testing for the determination of $RF_{CR}$ provided that it can be shown through a limited 1,000 hour creep testing program that the rupture behavior and envelope for the single ribs or yarns are the same as that for the full product. This comparison must demonstrate that there is no statistical difference between the full product regression line and the single rib or yarn regression line at a time of 1,000 hours using a student-t distribution at a confidence level of 0.10 (see Equation B.3-1 later in this appendix).
B.1 Overview of Extrapolation Approach to Determine the Ultimate Limit State Creep Limit, $T_1$

Considering that typical design lives for permanent MSE structures are 75 years or more, extrapolation of creep data will be required. Current practice allows creep data to be extrapolated up to one log cycle of time beyond the available data without some form of accelerated creep testing, or possibly other corroborating evidence (Jewell and Greenwood, 1988; Koerner, 1990). Based on this, unless one is prepared to obtain 7 to 10 years of creep data, temperature accelerated creep data, or possibly other corroborating evidence, must be obtained.

It is well known that temperature accelerates many chemical and physical processes in a predictable manner. In the case of creep, this means that the creep strains under a given applied load at a relatively high temperature and relatively short times will be approximately the same as the creep strains observed under the same applied load at a relatively low temperature and relatively long times. Temperature affects time to rupture at a given load in a similar manner. This means that the time to a given creep strain or to rupture measured at an elevated temperature can be made equivalent to the time expected to reach a given creep strain or to rupture at in-situ temperature through the use of a time shift factor.

The ability to accelerate creep with temperature for polyolefins such as polypropylene (PP) or high density polyethylene (HDPE) has been relatively well defined (Takaku, 1981; Bush, 1990; Popelar, et. al., 1991). Also for polyolefins, there is some risk that a “knee” in the stress rupture envelope due to a ductile to brittle transition could occur at some time beyond the available data (Popelar, et. al., 1991). Therefore, temperature accelerated creep data is strongly recommended for polyolefins. However, in practice, a ductile to brittle transition for polyolefin geosynthetic reinforcement products has so far not been observed, likely due to the highly oriented nature of polymer resulting from the processing necessary to make fibers and ribs. In general, the degree of orientation of the polymer is an important factor regarding the potential for ductile to brittle transitions.

For polyester (PET) geosynthetics, available evidence indicates that temperature can also be used to accelerate PET creep, based on data provided by den Hoedt, et. al., 1994 and others. However, the creep rupture envelopes for PET geosynthetics tend to be flatter than polyolefin creep rupture envelopes, and accurate determination of time-shift factors can be difficult for PET geosynthetics because of this. This may require greater accuracy in the PET stress rupture data than would be required for polyolefin geosynthetics to perform accurate extrapolations using elevated temperature data. This should be considered if using elevated temperature data to extrapolate PET stress rupture data. Note that a “knee” in the stress rupture envelope of PET does not appear to be likely based on the available data and the molecular structure of polyester.

If elevated temperature is used to obtain accelerated creep data, it is recommended that minimum increments of 10°C be used to select temperatures for elevated temperature creep testing. The highest temperature tested, however, should be below any transitions for the polymer in question. If one uses test temperatures below 70 to 75°C for polypropylene (PP), high density polyethylene (HDPE), and PET geosynthetics, significant polymer transitions will be avoided. If higher temperatures must be used, the effect of any transitions on the creep behavior should be carefully evaluated. One should also keep in mind that at these high temperatures, significant chemical interactions with the surrounding environment are possible, necessitating that somewhat lower temperatures or appropriate environmental controls be used. These chemical interactions are likely to cause the creep test results to be conservative. Therefore, from the user’s point of view, potential for chemical interactions is not detrimental to the validity of the data for predicting creep limits.

However, exposure to temperatures near the upper end of these ranges could affect the stress-strain behavior of the material due to loss of molecular orientation, or possibly other effects that are not the result of chemical degradation. Therefore, care needs to be exercised when interpreting results from tests performed at temperatures near the maximum test temperatures indicated above. In general, if the stiffness of the material after exposure to the environment is significantly different from that of
the virgin material, the stress-strain properties, and possibly the strength, of the material may have been affected by the exposure temperature in addition to the chemical environment. If the stiffness has been affected, the cause of the stiffness change should be thoroughly investigated to determine whether or not the change in stiffness is partially or fully due to the effect of temperature, or alternatively not use the data obtained at and above the temperature where the stiffness was affected.

Unless otherwise specified or required by site specific temperature data, an effective design temperature of 20°C (T_amb) should be assumed.

A number of extrapolation and creep modeling methods have been reported in the literature (Findley, et. al., 1976; Wilding and Ward, 1978; Wilding and Ward, 1981; Takaku, 1981; McGown, et. al., 1984; Andrawes, et. al., 1986; Murray and McGown, 1988; Bush, 1990; Poplar, et. al., 1991; Helwany and Wu, 1992). Many of the methods discussed in the literature are quite involved and mathematically complex.

Two creep extrapolation techniques are provided herein for creep rupture evaluation: the conventional method, which utilizes a simplified visual/graphical approach, temperature acceleration of creep, regression techniques, and statistical extrapolation, and the Stepped Isothermal Method (SIM). This does not mean that the more complex mathematical modeling techniques cannot be used to extrapolate creep of geosynthetics; they are simply not explained herein. These two techniques are described in more detail as follows:

**B.2 Step-By-Step Procedures for Extrapolating Creep Rupture Data – Conventional Method**

**Step 1:** Plot the creep rupture data as log time to rupture versus log load level, as shown in Figure B.2-1. Do this for each temperature in which creep rupture data is available. For some materials, a semi-log rather than a log-log plot could be used. The plotting method that provides the best and most consistent fit of the data should be used. In general, approximately 12 to 18 data points (i.e., combined from all temperature levels tested to produce the envelope, with a minimum of 4 data points at each temperature) are required to establish a rupture envelope (Jewell and Greenwood, 1988; ASTM D2837). The data points should be evenly distributed through each log cycle of time. Rupture points with a time to rupture of less than 5 hours should in general not be used, unless it can be shown that these shorter duration points are consistent with the rest of the envelope (i.e., they do not contribute to non-linearity of the envelope). As a guide, three of the test results should have rupture times (not shifted by temperature acceleration) of 10 to 100 hours, four of the test results should have rupture times between 100 and 1,000 hours, and four of the test results should have rupture times of 1,000 to 10,000 hours, with at least one additional test result having a rupture time of approximately 10,000 hours (1.14 years) or more. It is recommended that creep strain be measured as well as time to rupture, since the creep strain data may assist with conventional time-temperature shifting and in identifying any change in behavior that could invalidate extrapolation of the results.

**Step 2:** Extrapolate the creep rupture data. Elevated temperature creep rupture data can be used to extrapolate the rupture envelope at the design temperature through the use of a time shift factor, a_T. If the rupture envelope is approximately linear as illustrated in Figure B.2-1(a), the single time shift factor a_T should be adequate to perform the time-temperature superposition.

**Note 2:** This time-temperature superposition procedure assumes that the creep-rupture curves at all temperatures are linear on a semi-logarithmic or double logarithmic scale and parallel. It has been found empirically that the curves for PET are semi-logarithmic and approximately parallel, or double logarithmic and approximately parallel in the case of HDPE and PP. It should be pointed out that the theory of Zhurkov (1965), which assumes that the fracture process is activated thermally with the additional effect of applied stress, predicts that the creep-rupture characteristics should be straight when plotted on a double logarithmic diagram, and that their gradients should be stress-dependent.
Use of a single time shift factor to shift all the creep rupture data at a given temperature, termed “block shifting,” assumes that the shift factor $a_T$ is not highly stress level dependent and that the envelopes at all temperatures are parallel, allowing an average value of $a_T$ to be used for all of the rupture points at a given temperature. While research reported in the literature indicates that $a_T$ may be somewhat stress level dependent and that the curves at all temperatures are not completely parallel, this assumption tends to result in a more conservative assessment of the creep reduction factor $RF_{CR}$ (Thornton and Baker, 2002).

Figure B.2-1. Typical Stress Rupture Data for Geosynthetics, and the Determination of Shift Factors for Time-Temperature Superposition.

Figure B.2-2. Extrapolation of Stress Rupture Data and the Determination of the Creep Limit Load.

The time to rupture for the elevated temperature rupture data is shifted in accordance with the following equation:

$$t_{amb} = (t_{elev})(a_T)$$  \hspace{1cm} (B.2-1)
where, \( t_{\text{amb}} \) is the predicted time at the ambient or temperature to reach rupture under the specified load, \( t_{\text{elev}} \) is the measured time at elevated temperature to reach a rupture under the specified load, and \( a_T \) is the time shift factor. \( a_T \) can be approximately estimated using a visual/graphical approach as illustrated in figures B.2-1 and B.2-2. The preferred approach, however, is to use a computer spreadsheet optimization program to select the best shift factors for each constant temperature block of data to produce the highest \( R^2 \) value for the combined creep rupture envelope to produce the result in Figure B.2-2.

**Note 3:** Incomplete tests may be included, with the test duration replacing the time to rupture, but should be listed as such in the reported results, provided that the test duration, after time shifting, is 10,000 hours or more. The rule for incomplete tests is as follows. The regression should be performed with and without the incomplete tests included. If the incomplete test results in an increase in the creep limit, keep the incomplete tests in the regression, but if not, do not include them in the regression, in both cases for incomplete tests that are 10,000 hours in duration after time shifting or more. Record the duration of the longest test which has ended in rupture, or the duration of the longest incomplete test whose duration exceeds its predicted time to failure; this duration is denoted as \( t_{\text{max}} \).

It is preferred that creep rupture data be extrapolated statistically beyond the elevated temperature time shifted data using regression analysis (i.e., curve fitting) up to a maximum of one log cycle of time for all geosynthetic polymers (greater extrapolation using only statistical methods is feasible, but uncertainty in the result increases substantially and must be taken into account). Therefore, adequate elevated temperature data should be obtained to limit the amount of statistical extrapolation required.

**Note 4:** There may be situations where extrapolation to create a creep rupture envelope at a lower temperature than was tested is necessary. Situations where this may occur include the need to elevate the ambient temperature to have greater control regarding the temperature variations during the creep testing (i.e., ambient laboratory temperature may vary too much), or for sites where the effective design temperature is significantly lower than the “standard” reference temperature used for creep testing (e.g., northern or high elevation climates). In such cases, it is feasible to use lower bound shift factors based on previous creep testing experience to allow the creep rupture envelope to be shifted to the lower temperature, as shift factors for the materials typically used for geosynthetic reinforcement are reasonably consistent. Based on previous creep testing experience and data reported in the literature (Chow and Van Laeken 1991; Thornton, et al. 1998; Thornton, et al. 1999a; Lothspeich and Thornton 2000; Takemura 1959; Bush, 1990; Popelar, et al. 1990; Wrigley, et al. 2000; Takaku 1980; Thornton and Baker 2000), shift factors for HDPE and PP geosynthetics are typically in the range of 0.05 to 0.18 decades (i.e., log cycles of time) per \( 1^o \) C increase in temperature (i.e., a \( 10^o \) C increase would result in a time shift factor of 12 to 15) and 0.05 to 0.12 decades per \( 1^o \) C increase in temperature for PET geosynthetics. It is recommended that if shifting the creep rupture envelope to temperatures below the available data is necessary, that a shift factor of 0.05 decades per \( 1^o \) C increase in temperature for PP, HDPE, and PET be used. This default shift factor should not be used to shift the creep rupture data more than \( 10^o \) C.

**Step 3:** Once the creep data has been extrapolated, determine the design, lot specific, creep limit load by taking the load level at the desired design life directly from the extrapolated stress rupture envelope as shown in Figure B.2-2. If statistical extrapolation beyond the time shifted stress rupture envelopes (PP or HDPE), or beyond the actual data if temperature accelerated creep data is not available, is necessary to reach the specified design life, the calculated creep load \( T_l \) should be reduced by an extrapolation uncertainty factor as follows:

\[
T_l = \frac{P_{cl}}{(1.2)^{x-1}}
\]  

(B.2-3)

where \( P_{cl} \) is the creep limit load taken directly from the extrapolated stress rupture envelope, and “\( x \)” is the number of log cycles of time the rupture envelope must be extrapolated beyond the actual or...
time shifted data, and is equal to \( \log t_d - \log t_{\text{max}} \) as illustrated in Figure B.2-2. The factor 
\((1.2)^{x-1}\) is the extrapolation uncertainty factor. If extrapolating beyond the actual or time shifted 
data less than one log cycle, set “x-1” equal to “0”. This extrapolation uncertainty factor only applies 
to statistical extrapolation beyond the actual or time shifted data using regression analysis and 
assumes that a “knee” in the rupture envelope beyond the actual or time shifted data does not occur.

**Note 5**: A condition on the extrapolation is that there is no evidence or reason to believe that 
the rupture behavior will change over the desired design life. It should be checked that at long 
durations, and at elevated temperatures if used:

- There is no apparent change in the gradient of the creep-rupture curve
- There is no evidence of disproportionately lower strains to failure
- There is no significant change in the appearance of the fracture surface.

Any evidence of such changes, particularly in accelerated tests, should lead to the exclusion of any reading where either the gradient, strain at failure or appearance of the failure is different to those in the test with the longest failure duration. Particular attention is drawn to the behavior of unoriented thermoplastics under sustained load, where a transition in behavior is observed in long-term creep-rupture testing (i.e., the so called “ductile to brittle transition – Popelar, et al., 1991). The effect of this transition is that the gradient of the creep-rupture curve becomes steeper at the so-called “knee” such that long-term failures occur at much shorter lifetimes than would otherwise be predicted. The strain at failure is greatly reduced and the appearance of the fracture surface changes from ductile to semi-brittle. If this is observed, any extrapolation should assume that the “knee” will occur. For the method of extrapolation reference should be made to ISO/FDIS 9080: 2001, ASTM D2837, and Popelar; et al. (1991).

This extrapolation uncertainty factor also assumes that the data quality is good, data scatter is reasonable, and that approximately 12 to 18 data points which are well distributed (see Step 1 for a definition of well distributed) defines the stress rupture envelope. If these assumptions are not true for the data in question, this uncertainty factor should be increased. The uncertainty factor may also need to be adjusted if a method other than the one presented in detail herein is used for extrapolation. This will depend on how well that method compares to the method provided in this appendix. This extrapolation uncertainty factor should be increased to as much as \((1.4)^x\) if there is the potential for a “knee” in the stress rupture envelope to occur beyond the actual or time shifted data, or if the data quality, scatter, or amount is inadequate. Furthermore, if the data quantity or distribution over the time scale is inadequate, it may be necessary to begin applying the extrapolation uncertainty factor before the end of the time shifted data.

**Note 6**: Based on experience, the \(R^2\) value for the composite (i.e., time shifted) creep rupture 
envelope should be approximately 0.8 to 0.9 or higher to be confident that Equation B.2-3 will 
adequately address the extrapolation uncertainty. If the \(R^2\) value is less than approximately 0.6 
to 0.7, extrapolation uncertainty is likely to be unacceptably high, and additional testing and investigation should be performed. In general, such low \(R^2\) values are typically the result of data that is too bunched up, unusually high specimen to specimen variability, or possibly poor testing technique.

### B.3 Procedures for Extrapolating Creep Rupture Data – Stepped Isothermal Method (SIM)

An alternative creep strain/rupture analysis and extrapolation approach that has recently become available for geosynthetics is the Stepped Isothermal Method (SIM) proposed, illustrated, and
investigated by Thornton, et. al. (1997), Thornton, et. al. (1998), Thornton, et. al. (1998), and
Thornton and Baker (2002). SIM has been applied successfully to PET geogrids and PP geotextiles.
SIM utilizes an approach similar to the Williams-Landell-Ferry, or WLF, approach to creep
extrapolation (Ferry, 1980), where master creep curves for a given material are produced from
a series of short-term tests (i.e., creep test durations on the order of a few hours) on the same
specimen over a wide range of temperatures (i.e., while the load on the specimen is held constant,
the temperature is increased in steps). The sections of creep curve at the individual temperatures
are shifted in time and combined to form a continuous prediction of the creep strain at the starting
temperature.

Though the general principles of this method have been in use for many years in the polymer
industry (Ferry, 1980), it has been only recently that this approach has been used for geosynthetics.
Though this approach was initially developed to extrapolate creep strain data, it has been adapted
to produce stress rupture data by taking the specimen to rupture once the highest test temperature is
reached. In effect, through time shifting of the creep strain data obtained prior to rupture, the rupture
point obtained has an equivalent shifted time that is several orders of magnitude greater than the
actual test time, which could be on the order of only a few days.

The method is conducted in accordance with GRI:GS10. Key issues are the very short test time
used for this method, potential use of temperatures that are well above transitions in the geosynthetic
material, and its complexity. Key technical advantages of the method, however, include more
accurate determination of time shift factors, since the same specimen is used at the same load
level at all of the temperatures (the “conventional” method must deal with the effect of specimen
to specimen variability when determining the shift factors), and that time shift factors between
temperatures are determined at the same load level, eliminating the effect of load level in the
determination of the shift factors (in the “conventional” method, the shift factors used are in
fact an average value for a wide range of loads).

SIM can be considered for use in generating and extrapolating geosynthetic creep and creep rupture
data provided this method is shown to produce results which are consistent with the “conventional”
extrapolation techniques recommended in this appendix. To this end, creep-rupture testing shall be
conducted using conventional tests (ASTM D5262) and SIM tests (GRI GS10). At least six SIM
rupture tests and six conventional rupture tests shall be conducted on the product or at least
one of the products of the product line being evaluated. Of the six SIM rupture tests, four shall
have rupture times (shifted as appropriate) between 100 and 2000 hours and two shall have rupture
times greater than 2000 hours. All of the conventional creep rupture points shall be obtained at
the reference temperature (i.e., not temperature shifted). Creep rupture plots shall be constructed,
regression lines computed and the log times to rupture determined at a load level that corresponds
to 1,000 hours and 50,000 hours on the conventional creep rupture envelope, for the two data sets.
The log time to rupture for the SIM regression at this load level shall be within the upper and lower
90% confidence limits of the mean conventional regressed rupture time at the same load level using
Student’s t test.

The confidence limit for the regression performed for the conventional creep rupture data is given
by (Wadsworth, 1998):

\[
\log t_L = \log t_{reg} \pm t_{\alpha, n-2} \left[ \frac{1}{n} + \frac{(P - \bar{P})^2}{\sum (P_i - \bar{P})^2} \right] \times \sigma
\]  

(B.3-1)
and

\[ \sigma = \sqrt{\frac{\sum [\log t_i - \log \bar{t}]^2 - \left( \sum (P_i - \bar{P})(\log t_i - \log \bar{t}) \right)^2}{\sum (P_i - \bar{P})^2}} \]  \hspace{1cm} (B.3-2)

where:

\[ \log t_i = \text{lower and upper bound confidence limit. The } + \text{ or } - \text{ term in Equation B.2-1 results in the lower and upper bound confidence limits, respectively.} \]

\[ t_{\text{reg}} = \text{time corresponding to the load level from the conventional creep rupture envelope at which the comparison between the two envelopes will be made (e.g., at 1,000 and 50,000 hrs after time shifting).} \]

\[ t_{\alpha,n-2} = \text{value of the } t \text{ distribution determined from applicable Student } t \text{ table (or from the Microsoft EXCEL function TINV(} \alpha, n-2)) \text{ at } \alpha = 0.10 \text{ and } n-2 \text{ degrees of freedom (this corresponds to the } 90\% \text{ two-sided prediction limit).} \]

\[ n = \text{the number of rupture or allowable run-out points in the original test sample (i.e., the conventional creep rupture data)} \]

\[ P = \text{load level obtained at } t_{\text{reg}} \text{ from the regression line developed from the conventional creep rupture testing} \]

\[ \bar{P} = \text{the mean rupture load level for the original test sample (i.e., all rupture or run-out points used in the regression to establish the conventional creep rupture envelope)} \]

\[ P_i = \text{the rupture load level of the } i\text{'th point for the rupture points used in the regression for establishing the conventional creep rupture envelope} \]

\[ \log t = \text{the mean of the log of rupture time for the original test sample (i.e., all rupture or run-out points used in the regression to establish the conventional creep rupture envelope)} \]

\[ t_i = \text{the rupture time of the } i\text{'th point for the rupture points used in the regression for establishing the conventional creep rupture envelope} \]

Once \( \log t_i \), both upper and lower bound, has been determined at the specified load level, compare these values to the log rupture time (i.e., \( \log t_{\text{SIM}} \)) obtained for the SIM creep rupture envelope test at the specified load level (e.g., 1,000 and 50,000 hours). The value of \( \log t_{\text{SIM}} \) at the two specified load levels must be between the upper and lower bound confidence limits (\( \log t_i \)). If this requirement is not met, perform two additional SIM tests at each load level \( P \) for the specified \( t_{\text{reg}} \) where this comparison was made and develop a new SIM creep rupture envelope using all of the SIM data. If for the revised SIM regression envelope resulting from these additional tests this criterion is still not met, perform adequate additional conventional creep rupture testing to establish the complete rupture envelope for the product in accordance with this appendix.

If the criterion provided above is met, the SIM testing shall be considered to be consistent with the conventional data, and SIM may be used in combination with the conventional data to meet the requirements of Section B.2 regarding the number of rupture points and their distribution in time and maximum duration. Therefore, the combined data can be used to create the creep rupture envelope
as shown in Figure B.2-2. In that figure, the SIM data shall be considered to already be time shifted. Equation B.2-3 is then used to determine \( T_1 \).

**B.4 Determination of \( R_{FCR} \)**

The creep reduction factor, \( R_{FCR} \), is determined by comparing the long-term creep strength, \( T_1 \), to the ultimate tensile strength (ASTM D4595 or ASTM D6637) of the sample tested for creep (\( T_{ul} \)). The sample tested for ultimate tensile strength should be taken from the same lot, and preferably the same roll, of material that is used for the creep testing. For ultimate limit state design, the strength reduction factor to prevent long-term creep rupture is determined as follows:

\[
R_{FCR} = \frac{T_{lot}}{T_1}
\]

(B.4-1)

where, \( T_{lot} \) is the average lot specific ultimate tensile strength (ASTM D4595 or ASTM D6637) for the lot of material used for the creep testing. Note that this creep reduction factor takes extrapolation uncertainty into account, but does not take into account variability in the strength of the material. Material strength variability is taken into account when \( R_{FCR} \), along with \( R_{ID} \) and \( R_{FD} \), are applied to \( T_{ul} \) to determine the long-term allowable tensile strength, as \( T_{ul} \) is a minimum average roll value. The minimum average roll value is essentially the value that is two standard deviations below the average value.

**B.5 Use of Creep Data from “Similar” Products and Evaluation of Product Lines**

Long-term creep data obtained from tests performed on older product lines, or other products within the same product line, may be applied to new product lines, or a similar product within the same product line, if one or both of the following conditions are met:

- The chemical and physical characteristics of tested products and proposed products are shown to be similar. Research data, though not necessarily developed by the product manufacturer, should be provided which shows that the minor differences between the tested and the untested products will result in equal or greater creep resistance for the untested products.
- A limited testing program is conducted on the new or similar product in question and compared with the results of the previously conducted full testing program.

For polyolefins, similarity could be judged based on molecular weight and structure of the main polymer (i.e., is the polymer branched or crosslinked, is it a homopolymer or a blend, percent crystallinity, etc.), percentage of material reprocessed, tenacity of the fibers and processing history, and polymer additives used (i.e., type and quantity of antioxidants or other additives used). For polyesters and polyamides, similarity could be judged based on molecular weight or intrinsic viscosity of the main polymer, carboxyl end group content, percent crystallinity, or other molecular structure variables, tenacity of the fibers and processing history, percentage of material reprocessed or recycled, and polymer additives used (e.g., pigments, etc.). The untested products should also have a similar macrostructure (i.e., woven, nonwoven, extruded grid, needlepunched, yarn structure, etc.) and fiber dimensions (e.g., thickness) relative to the tested products. It should be noted that percent crystallinity is not a controlled property and there is presently no indication of what an acceptable value for percent crystallinity should be.

For creep evaluation of a similar product not part of the original product line, this limited testing program should include creep tests taken to at least 1,000 to 2,000 hours in length before time shifting if using the “conventional” creep testing approach, with adequate elevated temperature data to permit extrapolation to 50,000 hours or more. If it has been verified that SIM can be used, in accordance with Section B.3, durations after time shifting due to elevated temperature up to a
minimum of 50,000 hours are required. A minimum of 4 data points per temperature level tested should be obtained to determine time shift factors and to establish the envelope for the similar product. These limited creep test results must show that the performance of the similar product is equal to or better than the performance of the product previously tested. This comparison must demonstrate that there is no statistical difference between the old product regression line and the regression line obtained for the similar product at a time of 2,000 hours (not temperature accelerated) and 50,000 hours (after time shifting) using a student-t distribution at a confidence level of 0.10 (see Equation B.3-1). If no statistical difference is observed, the results from the full testing program on the older or similar product could be used for the new/similar product. If this is not the case, then a full testing and evaluation program for the similar product should be conducted.

Similarly, for extension of the creep data obtained on one product in the product line to the entire product line as defined herein, a limited creep testing program must be conducted on at least two additional products in the product line. The combination of the three or more products must span the full range of the product line in terms of weight and/or strength. The limited test program described in the preceding paragraph should be applied to each additional product in the product line. The loads obtained for the data in each envelope should then be normalized by the lot specific ultimate tensile strength, $T_{\text{lot}}$. All three envelopes should plot on top of one another, once normalized in this manner, and the two additional product envelopes should be located within the confidence limits for the product with the more fully developed creep rupture envelope as described above for “similar” products. If this is the case, then the creep reduction factor for the product line shall be the lesser of the reduction factor obtained for the product with the fully developed rupture envelope and the envelope of all three products combined, and normalization using the ultimate tensile strength shall be considered acceptably accurate.

If this is not the case, then the creep rupture envelopes for the other two products, plus enough other products within the product line, to establish the trend in $R_{\text{CR}}$ as a function of product weight or ultimate tensile strength, so that the $R_{\text{CR}}$ for the other products within the product line can be accurately interpolated. Furthermore, $T_{\text{al}}$ must be determined in accordance with Note 7.

**Note 7:** Note that normalization using the ultimate lot specific tensile strength may not be completely accurate for some geosynthetic products regarding characterization of creep rupture behavior, and other normalization techniques may be needed (Wrigley, et al., 1999). In such cases, individual creep reduction factors for each product in the product line may need to be established through fully developed creep rupture envelopes for representative products obtained at the low, middle, and high strength end of the product series. Once the creep limited strength, $P_{\text{al}}$, and the creep reduction factors are established for each product, in this case, product variability must still be taken into account. In such cases, $T_{\text{al}}$ must be the lesser of the determination from Equation 1 and the following determination:

$$T_{\text{al}} = \frac{P_{95}}{R_{\text{ID}} \times R_{F}}$$

where,

$P_{95} = \text{the tensile strength determined from the 95\% lower bound prediction limit for the creep rupture envelope at the specified design life (see Equations 4 and 5 in “Quality Assurance (QA) Criteria for Comparison to Initial Product Acceptance Test Results”)}$

**References**


*Sixth International Conference on Geotextiles, Geomembranes, and Related Products*, Atlanta, GA, pp.699-706.


WSDOT Test Method No. 925, Appendix C

Strain Based Creep Testing and Extrapolation

This appendix provides supplementary information to Appendix B regarding the use and extrapolation of creep strain data.

As is true for stress rupture testing, in-air long-term laboratory creep tests should be conducted for a range of load levels in accordance with ASTM D5262, adequate for extrapolation to the required design life as described in this appendix. Specimens should be tested in the direction in which the load will be applied in use. Test results should be extrapolated to the required structure design life. Based on the extrapolated test results, for ultimate limit state design, determine the highest load level, designated Tl, at which the log time creep rate continues to decrease with time and which precludes both ductile and brittle creep rupture within the required lifetime. Tl should be determined at the required design life and at the effective design site temperature. Unless otherwise specified or required by site specific temperature data, an effective design temperature of 20°C (Tamb) should be assumed.

As stated above, the application of the extrapolated creep data as described in this appendix is to estimate the highest load level within the specified design lifetime that precludes creep rupture. Creep strain data can be used for other purposes, such as to estimate long-term deformations or to estimate long-term stiffness values. The extrapolated creep strain data developed in accordance with this Appendix can be used for these purposes as well.

C.1 Creep Strain Assessment Tools and Concepts

Creep strain curves are typically plotted as a function of time or the logarithm of time. In general, there are up to three stages of creep observed in polymeric materials – these include primary, secondary or steady-state, and tertiary creep. Primary creep strains are characteristically linear when plotted against a logarithmic time scale and increase at a decreasing rate on an arithmetic time scale. Secondary creep strains are typically linear when plotted against an arithmetic time scale. Tertiary creep is the rupture phase of creep and is characterized by a rapidly increasing creep rate with time. Geosynthetic structure tends to dominate primary creep (at least for nonwoven geotextiles, but much less so for woven geotextiles and not at all for geogrids), and the polymer characteristics tend to dominate secondary and tertiary creep mechanisms (Allen 1991). Polyolefins (HDPE and PP) tend to exhibit all three stages of creep, depending on the load level, whereas PET tends to only exhibit primary and tertiary creep. Figure C.1-1 illustrates these concepts.
The instability limit strain is defined as the strain beyond which the material exhibits signs of instability, i.e., approaches failure (Andrawes, et. al., 1986). When extrapolating creep strain data, it is important to not extrapolate the data to strain levels that are in excess of the instability limit strain, as doing so would produce invalid results. The determination of this limit strain can be the most difficult part of assessing the creep rupture limit from creep strain data. The actual rupture strain, which occurs at the end of tertiary creep, for a given material at a given load level is difficult to measure as well as to identify. A more consistent and more easily measured instability limit strain would be the strain level where tertiary creep begins as illustrated in Figure C.1-1. However, if rupture occurs during the primary or secondary creep stage (e.g., PET) the instability limit strain is the rupture strain.

Another tool that can be useful in determining the strain at the beginning of tertiary creep is the Sherby-Dorn plot, as illustrated in Figure C.1-2. A Sherby-Dorn plot is a well known plotting technique used in polymer science (McGown et al. 1984a). Each curve represents a specific geosynthetic layer in a wall or a specific geosynthetic specimen tested at a specific load level. Creep strain rates observed under constant load are plotted against the total strain in the specimen or layer measured at the time the creep strain rate was calculated. The creep strain rate is simply the slope of the creep strain curve at a given point in time (see Figure C.1-2a). Curves that are linear or concave downward indicate that only primary creep is occurring, and that stabilization (no rupture) is likely. Curves that are concave upward indicate secondary or tertiary creep is occurring, and that rupture is likely. The closer the curves are located to the bottom left corner of the plot (Figure C.1-2), the better the creep performance of the material. The closer the curves are to the upper right corner, the more likely creep rupture will occur.
(a) Determination of creep strain rate.

\[ \text{Creep strain rate} = \frac{\Delta \varepsilon}{\Delta t} \]

Load level = \( P_2 \)

(b) Interpretation of Sherby-Dorn plots.

Figure C.1-2. Development and Use of Sherby-Dorn Plots to Analyze Creep Strain Data.
Note that some interpretation of the creep curves through curve fitting is required to determine strain rates, since local jumps in the measured creep strain curves can cause wide variations in calculated creep strain rates. The jumps in the curves are typically the result of the short increments of time used in the calculations and the small magnitude of changes in strain readings that may be at the limit of the resolution of the measuring devices. Hence, the slope of the measured creep curves must be taken over fairly long increments of time to be meaningful.

Tertiary creep begins where the creep strain rate (based on an arithmetic time scale) begins to increase. The strain at the beginning of tertiary creep is located where the creep strain rate begins to increase after reaching a minimum value, at least for polyolefin geosynthetics. A minimum and then an increasing creep strain rate is very difficult to see for polyester geosynthetics on this type of plot. Therefore, Sherby-Dorn plots tend to not be very useful for polyester geosynthetics.

To determine the long-term instability limit strain, the measured creep strain and time near creep rupture (i.e., at the beginning of tertiary creep) for various load levels must be obtained. These strains are plotted versus time to the beginning of tertiary creep (the rupture phase) on a semi-log plot (i.e., log time), or possibly a log-log plot, to establish the trend in the data. In general, strains near rupture for times to the rupture phase from 10 hours up to approximately 10,000 hours should be obtained so that the data need only be extrapolated two log cycles of time or less. A minimum of one data point per log cycle of time should be obtained to define the trend, but more data points are likely to be needed to establish the trend.

Typical near rupture strain trends for various geosynthetic polymers are illustrated in Figure C.1-3. Strain near rupture which increases as time to the rupture phase increases may be indicative of ductile behavior, whereas strain near rupture which decreases as time to the rupture phase increases may be indicative of brittle behavior (i.e., localized crack growth). If ductile behavior is observed, a transition to brittle behavior is possible. If such a shift to brittle behavior occurred, the strain near rupture increase occurring as time to the rupture phase increases (ductile behavior) could be lost as behavior becomes more brittle. Until more is known, it is recommended that strains near rupture which appear to be increasing as time to the rupture phase increases not be depended upon when assessing the long-term instability limit strain. Therefore, if the strain near rupture increases as time to the rupture phase increases, which appears to be typical of polypropylene (PP) geosynthetics (Takaku, 1981; Allen, 1991, Thornton and Baker, 2002), the short-term (i.e., rupture times on the order of 10 hours) strain near rupture should be used as the instability limit strain. If the strain near rupture decreases as time to the rupture phase increases, which appears to be typical of high density polyethylene (HDPE) geosynthetics (Ingold, et. al., 1994; Allen and Bathurst, 1996) and polyester (PET) geosynthetics (Krumm, 1988; Allen and Bathurst, 1996), extrapolate (up to a maximum of two log cycles) to the strain near rupture at the specified design life using visual extrapolation, regression analysis, or time-temperature superposition if elevated temperature data is available, and reduce that strain by a reduction factor of 1.1 to account for potential uncertainty. This reduction factor could be adjusted depending on the amount and quality of the strain near rupture data. The determination of the long-term instability strain limit is conceptually illustrated in Figure C.1-3.
If inadequate data is available to extrapolate measured strains near rupture to the long-term strain near rupture and the material type is likely to have strains near rupture which decrease with increasing time to the rupture phase (e.g., HDPE and PET), an acceptable alternative to estimate the instability limit strain is to take the peak strain measured from a wide width load-strain test (ASTM D4595 or ASTM D6637) and reduce it by a factor of 2.0.

Another tool that is useful for characterizing and working with creep strain data is the isochronous curve. Each isochronous (i.e., constant time) curve is created by taking load and strain levels from each creep curve at a given constant time and plotting them to form an isochronous curve. Some curve fitting may be necessary to account for specimen variability and to create a reasonably smooth curve. For HDPE, this curve fitting is reasonably straight-forward to do. However, for PET, the isochronous curves characteristically have an “s” shape at low strains resulting from load and strain dependent changes in the crystalline and between crystal arrangement of molecules within the polymer (Jewell and Greenwood, 1988; den Hoedt, et al, 1994). PP geosynthetics may also have inherent abrupt non-linearities in their isochronous curves at certain strain or load levels as observed by McGown, et al. (1984). Such changes in the curves resulting from physical processes in the polymer must be considered when constructing isochronous from creep strain data. Note that isochronous curves are not an extrapolation tool, but instead are an interpolation tool. Isochronous curves provide a convenient method of interpolating between creep curves, which will be necessary to accomplish the creep extrapolation steps that follow.
Creep stiffness curves as a function of time are also very useful to use for extrapolation purposes, or to simply estimate the long-term stiffness of the reinforcement for purposes of estimating deformation behavior. Select a constant strain or a constant load level from which to calculate the creep stiffness from the isochronous creep curves. At each point where the selected strain or load level intersects an isochronous curve, calculate the creep stiffness, $J$, as shown in Figure C.1-5, using the general form of the equation shown below:

$$J = \frac{P}{\epsilon}$$  \hspace{1cm} \text{(C.1-1)}

where, “$P$” is the load, and “$\epsilon$” is the strain. Each isochronous curve represents the specific time associated with the calculated creep stiffness. The creep stiffness values calculated should be plotted as a function of log time as shown in Figure C.1-5. Note that creep stiffness values can be calculated directly from the creep curves (Figure C.1-1) by identifying where the specified strain intersects each creep curve, or for each creep curve estimate the strain at each selected time. As shown in Figure C.2.1-1, the load or strain level at which the creep stiffness values are calculated should be selected such that the strain near the end of the specified design life is approximately equal to or less than the instability limit strain, if it is desired to predict the maximum load level that will preclude rupture, $T_l$. Note that other strain or load levels could be selected to calculate the creep stiffness curves if it is desired to estimate deformations at working stress conditions or to estimate the long-term creep stiffness at working strains.
C.2 Creep Strain Data Extrapolation

The ability to accelerate creep with temperature for polyolefins such as polypropylene (PP) or high density polyethylene (HDPE) has been relatively well defined (Takaku, 1981; Bush, 1990; Popelar, et. al., 1991). Since the focus of the creep strain extrapolation method provided in this appendix is on the creep limit to prevent rupture during the design lifetime required, the issues affecting creep rupture data extrapolation also affect creep strain data extrapolation. Therefore, as is true for creep rupture testing, temperature accelerated creep data is strongly recommended for polyolefins. For polyester (PET) geosynthetics, evidence indicates that temperature can also be used to accelerate PET creep, based on data provided by den Hoedt, et. al., (1994), and others.

If elevated temperature is used to obtain accelerated creep data, it is recommended that minimum increments of 10°C be used to select temperatures for elevated temperature creep testing. The highest temperature tested, however, should be below any transitions for the polymer in question. If one uses test temperatures below 70 to 75°C for polypropylene (PP), high density polyethylene (HDPE), and PET geosynthetics, significant polymer transitions will be avoided. One should also keep in mind that at these high temperatures, significant chemical interactions with the surrounding...
environment are possible, necessitating that somewhat lower temperatures or appropriate environmental controls be used. These chemical interactions are likely to cause the creep test results to be conservative. Therefore, from the user’s point of view, potential for chemical interactions is not detrimental to the validity of the data for predicting creep limits. However, exposure to temperatures near the upper end of these ranges could affect the stress-strain behavior of the material due to loss of molecular orientation, or possibly other effects that are not the result of chemical degradation. Therefore, care needs to be exercised when interpreting results from tests performed at temperatures near the maximum test temperatures indicated above. In general, if the stiffness of the material after exposure to the environment is significantly different from that of the virgin material, the stress-strain properties, and possibly the strength, of the material may have been affected by the exposure temperature in addition to the chemical environment. If the stiffness has been affected, the cause of the stiffness change should be thoroughly investigated to determine whether or not the change in stiffness is partially or fully due to the effect of temperature, or alternatively not use the data obtained at and above the temperature where the stiffness was affected.

A number of extrapolation and creep modeling methods have been reported in the literature (Findley, et. al., 1976; Wilding and Ward, 1978; Wilding and Ward, 1981; Takaku, 1981; McGown, et. al., 1984a; Andrawes, et. al., 1986; Murray and McGown, 1988; Bush, 1990; Popelar, et. al., 1991; Helwany and Wu, 1992). Many of the methods discussed in the literature are quite involved and mathematically complex.

Two creep extrapolation techniques are provided herein for creep rupture evaluation: the conventional method, which utilizes a simplified visual/graphical approach, temperature acceleration of creep, regression techniques, and statistical extrapolation, and the Stepped Isothermal Method (SIM). This does not mean that the more complex mathematical modeling techniques cannot be used to extrapolate creep of geosynthetics; they are simply not explained herein.

The two techniques identified above are described in more detail in Appendix B, and as follows:

C.2.1 Step-By-Step Procedures for Extrapolating Creep Strain Data – Conventional Method

**Step 1:** Plot the creep data. Plot the data as a semilog plot (log of time) or as an arithmetic plot (time). Do this separately for each temperature if data at elevated temperatures is available. For examples, see Figure C.1-1.

**Step 2:** Determine the instability limit strain (see Section C.1).

**Step 3:** Construct isochronous curves, as shown in Figure C.1-4.

**Step 4:** Develop creep stiffness curves for each temperature in which creep data is available, all at the same load level or strain level (see Figure C.1-5). Develop these stiffness curves at a strain level near the instability strain limit, or at a load level that results in a strain near the end of the specified design life that is approximately equal to the instability strain limit. These creep stiffness curves can then be used to perform time-temperature superposition for the purpose of creep extrapolation. Note that more accurate time-temperature shift factors are likely to be obtained if the creep stiffness curves are produced at a constant load level (Figure C.1-5b) rather than constant strain level (Figure C.1-5a), as doing so avoids the additional uncertainty caused by the stress level dependence of the shift factors. In fact, using constant load level creep stiffness curves to determine shift factors should produce nearly identical results to the Stepped Isothermal Method (SIM), except that specimen to specimen variability will still be present (SIM eliminates the specimen to specimen variability when determining shift factors, since only one specimen is used – See Appendix B for more information on SIM).
**Step 5:** Extrapolate the creep data. For all geosynthetics, creep strain or stiffness data can be extrapolated statistically using regression analysis (i.e., curve fitting), or creep data can be accelerated by temperature to allow extrapolation using time-temperature superposition principles. It is well known that temperature accelerates many chemical and physical processes in a predictable manner. In the case of creep, this means that the creep strains under a given applied load at a relatively high temperature and relatively short times will be approximately the same as the creep strains observed under the same applied load at a relatively low temperature and relatively long times. This means that the time to a given creep strain or stiffness measured at an elevated temperature can be made equivalent to the time expected to reach a given creep strain or stiffness at in-situ temperature through the use of a time shift factor. Therefore, elevated temperature creep strain or stiffness data is made into equivalent in-situ temperature data as follows:

\[ t_{\text{amb}} = (t_{\text{elev}})(a_T) \]  

(C.2.1-1)

where, \( t_{\text{amb}} \) is the predicted time at in-situ temperature to reach a specified creep stiffness or strain under the specified load, \( t_{\text{elev}} \) is the measured time at elevated temperature to reach a specified creep stiffness or strain under the specified load, and \( a_T \) is the time shift factor. For example, this means that if the time to reach a creep stiffness \( J_{cl} \) at elevated temperature is 10,000 hours, the creep stiffness will also be \( J_{cl} \) at in-situ temperature at a time of \((10,000)(a_T)\) hours. In this way, the creep stiffness curve at in-situ temperature can be extrapolated to longer times.

The magnitude of the time shift factor can be determined graphically as illustrated in Figure C.2.1-1, or regression analysis of the composite creep stiffness curve can be used to optimize the shift factors to produce the highest \( R^2 \) value for all of the time shifted data. Adjust \( a_T \) such that the creep stiffness curves at elevated temperature line up with the creep stiffness curve at the design (in-situ) temperature. Note that the magnitude of the shift factors for a given product can be different at different stages of creep (i.e., primary, secondary, or tertiary creep) and different load levels (see Appendix B). This should be considered when determining shift factors. See Note 4 in Appendix B, Section B.2 for additional considerations regarding the use of time shift factors.

**Step 6:** Once the creep data has been extrapolated, determine the design, lot specific, creep limit load as follows:

\[ T_1 = (J_{cl})(\varepsilon_i) \]  

(C.2.1-2)

where, \( J_{cl} \) is the creep stiffness at the desired design life and temperature, and \( \varepsilon_i \) is the instability limit strain. If statistical extrapolation beyond the time shifted creep stiffness curves, or beyond the actual data if temperature accelerated creep data is not available, is necessary to reach the specified design life, the calculated creep load \( T_1 \) should be reduced by an extrapolation uncertainty factor as follows:

\[ T_1 = (J_{cl})(\varepsilon_i)/(1.2)^{x-1} \]  

(C.2.1-3)

where “\( x \)” is the number of log cycles of time the creep stiffness data must be extrapolated. Extrapolations greater than two log cycles of time are not recommended (see Appendix B for a more detailed explanation). The factor \((1.2)^{x-1}\) is the extrapolation uncertainty factor. If extrapolating beyond the actual or time shifted data less than one log cycle, set “\( x-1 \)” equal to “0”. This extrapolation uncertainty factor only applies to statistical extrapolation beyond the actual or time shifted data using regression analysis and assumes that a shift from ductile to brittle behavior beyond the actual or time shifted data does not occur. This extrapolation uncertainty factor also assumes that the data quality is good, data scatter is reasonable, and that a minimum of 5 load levels for each temperature are tested. The load levels should be well distributed within the load range tested, and at least one to two load levels should be high enough to produce rupture within the test time at each test temperature. If these assumptions are not true for the data in question, this uncertainty factor
should be increased. The uncertainty factor may also need to be adjusted if a method other than the one presented in detail herein is used for extrapolation. This will depend on how well that method compares to the method provided in this appendix. This extrapolation uncertainty factor should be increased to as much as \((1.4)^n\) if there is the potential for a shift from ductile to brittle behavior to occur beyond the actual or time shifted data, or if the data quality, scatter, or amount is inadequate (see Appendix B, Notes 5 and 6). Furthermore, if the data quality is inadequate, it may be necessary to begin applying the extrapolation uncertainty factor before the end of the time shifted data.

Figure C.2.1-1. Use of Time-Temperature Superposition to Extrapolate Creep Stiffness Curves.

(a) Creep stiffness curves and determination of shift factors.

(b) Extrapolation of creep stiffness curves and determination of creep limit.

Figure C.2.1-1. Use of Time-Temperature Superposition to Extrapolate Creep Stiffness Curves.
C.2.2 Step-By-Step Procedures for Extrapolating Creep Strain Data – Stepped Isothermal Method (SIM)

See Appendix B for details regarding the description of SIM and issues regarding its use. SIM can be considered for use in generating and extrapolating geosynthetic creep data provided this method is shown to produce results which are consistent with the “conventional” extrapolation techniques recommended in this appendix. To this end, creep testing shall be conducted using conventional tests (ASTM D5262) and SIM tests (GRI GS10). At least six SIM creep tests and six conventional creep tests and shall be conducted on the product or at least one of the products of the product line being evaluated. The load levels selected for the SIM tests and the conventional tests shall be the same. The range of load levels for both types of tests shall be evenly distributed between loads near the creep limit typical for the type of geosynthetic and polymer tested and load levels high enough to result in failure in approximately 100 hours or less (shifted time for SIM, unshifted time for conventional tests). All of the conventional creep tests shall be obtained at the reference temperature (i.e., not temperature shifted). The minimum duration of all of the tests shall be 1,000 hours or more (unshifted for conventional tests and time shifted for SIM).

The comparison between the SIM and conventional creep tests data shall be performed at a specified strain. The specified strain will depend on the strains observed in all of the creep tests (SIM and Conventional). Select a strain that will intercept all of the creep curves as much as possible. Preferably, the strain level should be approximately 5 to 10% or more. Where the selected strain level intersects each creep curve, determine the time required to reach the specified strain. Plot the load level as a function of the logarithm of time to reach the specified strain for each set of data, and perform a regression for each data set. Use the confidence limit test for comparing SIM to conventional data as described in Appendix B, using Equations B.3-1 and B.3-2. The log times to the specified strain level shall be determined at a load level that corresponds to 1,000 hours and 50,000 hours on the conventional creep envelope, for both data sets. The log time to rupture for the SIM regression at this load level shall be within the upper and lower 90% confidence limits of the mean conventional regressed time to the specified strain at the same load level using Student’s \( t \) test (see Appendix B for details regarding how to calculate this).

If this requirement is not met, perform two additional SIM tests at each load level \( P \) for the specified \( t_{\text{reg}} \) where this comparison was made and develop a new SIM time to the specified creep strain envelope using all of the SIM data. If for the revised SIM regression envelope resulting from these additional tests this criterion is still not met, perform adequate additional conventional creep rupture testing to adequately establish \( T_l \) for the product in accordance with this appendix.

If the criterion provided above is met, the SIM testing shall be considered to be consistent with the conventional data, and SIM may be used in combination with the conventional data to meet the requirements of this appendix regarding the number and duration of creep tests. In this case, the SIM data can be used to establish an isochronous curve at the specified design life (see Figure C.1-4). Once established, using a strain level equal to the instability strain limit for the product determined in accordance with Figure C.1-3 and Section C.1, determine the load \( P_o \) directly from the isochronous curve and calculate the creep limit load \( T_l \).

C.3 Determination of \( RF_{CR} \)

The creep reduction factor, \( RF_{CR} \), is determined by comparing the long-term creep strength, \( T_l \), to the ultimate tensile strength (ASTM D4595 or ASTM D6637) of the sample tested for creep. The sample tested for ultimate tensile strength should be taken from the same lot, and preferably the same roll, of material that is used for the creep testing. For ultimate limit state design, the strength reduction factor to prevent long-term creep rupture is determined as follows:

\[
RF_{CR} = \frac{T_{\text{tot}}}{T_l}
\]  

(C.1-5)
where, $T_{lot}$ is the average lot specific ultimate tensile strength (ASTM D4595 or ASTM D6637) for the lot of material used for the creep testing. Note that this creep reduction factor takes extrapolation uncertainty into account, but does not take into account variability in the strength of the material. Material strength variability is taken into account when $RF_{CR}$, along with $RF_{ID}$ and $RF_{D}$, are applied to $T_{lot}$ to determine the long-term allowable tensile strength, as $T_{ult}$ is a minimum average roll value. The minimum average roll value is essentially the value that is two standard deviations below the average value.

C.4 Estimation of Long-Term Creep Deformation

In-isolation creep strain data can be used to estimate post-construction strains and deformations (see Allen and Bathurst, 2002b). Since load levels in full scale structures as a percent of the ultimate tensile strength are generally quite low, adequate creep data must be obtained at low load levels (typically in the range of 2 to 20% of ultimate). The key to accurate estimation of creep strains in full scale structures is an accurate prediction of the load level.

**Step 1:** Estimate the load levels in the reinforcement layer(s). Current design specifications (AASHTO 2004, 2002) use the Simplified Method to estimate reinforcement loads in walls, or slope stability analysis techniques (Elias, et al., 2001) to assess reinforcement loads in reinforced slopes. Loads should be estimated for this purpose without any factor of safety or load factor applied. Based on the results obtained by Allen and Bathurst (2002a), the methods provided in current design specifications and guidelines tend to significantly over-predict reinforcement load levels in geosynthetic structures. A new method (the K-Stiffness Method) has been developed by Allen et al. (2003) that appears to predict reinforcement loads in geosynthetic structures much more accurately.

**Step 2:** From the available creep strain data, create isochronous curves (see Section C.1 and Figure C.1-4).

**Step 3:** Use the isochronous curves to create a creep strain curve at the desired load level, by selecting the strains at each time at the selected constant load level.

**Step 4:** Select the portion of the creep strain curve that is applicable to the post-construction strains in the full scale structure, accounting for the time required to build the structure (see Figure C.4-1).

![Figure C.4-1. Conceptual illustration showing adjustment of reinforcement creep strains measured in walls to in-isolation laboratory creep data (after Allen and Bathurst, 2002b).](image-url)
Step 5: The creep strain is calculated as the strain at the desired design life plus the time required for the construction of the geosynthetic structure minus the strain at the end of the geosynthetic structure construction.

C.5 Estimation of Creep Stiffness for Working Stress Design

Strains for working stress design, such as when using the K-Stiffness Method (Allen et al., 2003) are typically small (i.e., approximately 2%). Since reinforcement stiffness is generally a nonlinear function of strain, it is important to obtain the creep stiffness at the appropriate strain level. The primary purpose of this stiffness calculation is to provide input data for working stress methods such as the K-Stiffness method (Allen et al., 2003), or for more sophisticated analyses such as finite element or finite difference numerical simulations. Typically, the stiffness at the end of structure construction ($J_{EOC}$) or at the end of the structure design life ($J_{DL}$) would be required as input for this type of analysis. Either “conventional” creep testing may be used, or SIM may be used if SIM is determined to be consistent with the conventional data as described in Section C.2.2.

Step 1: Obtain creep strain data for at least one load level low enough to produce a strain level at the end of structure construction (assume to be 1,000 hrs unless otherwise specified by the approval authority), and a second load level low enough to produce a strain level of approximately 4% at end of structure construction (i.e., 1,000 hrs). A minimum of two specimens per product tested at each strain level (i.e., two specimens each at the 2 and 4% strain levels) shall be tested.

Step 2: Use this, and any other available creep strain data for the same lot of material, to create isochronous creep curves as shown in Figure C.1-4.

Step 3: Create creep stiffness curves at the desired constant strain level (typically 2%) as shown in Figure C.1.5, and extrapolate to the desired time as necessary using the “conventional” approach illustrated in Figure C.2.1-1 or using SIM. Be sure to extrapolate the creep stiffness curve to the desired effective design temperature. The “conventional” approach (ASTM D 5262) shall be used unless comparative testing is conducted that demonstrated SIM (ASTM D6992) is providing results that are consistent with the "conventional" approach as defined in T 925.

Step 4: Obtain the creep stiffness, $J_{EOC}$ or $J_{DL}$, at the desired time from the creep stiffness curve as shown in Figure C.2.1-1.

C.6 Evaluation of Product Lines

Appendix B Section B.5 shall be used as the basis to apply creep strain data to product lines.

References


This appendix provides guidance regarding the performance of long-term product specific durability testing that may be conducted for product acceptance in lieu of the durability index testing as described in “Determination of Long-term Geosynthetic Strength for Initial Product Acceptance” as provided in this protocol. The procedures that follow are required if it is desired to use a value of RFD less than the default minimum of 1.3, or to determine RFD for environments that are defined as aggressive.

This appendix has been developed to address polypropylene (PP), polyethylene (PE or HDPE), and polyester (PET) geosynthetics. While the chemical and biological durability procedures and criteria provided herein may also be applicable to other polymers (for example, hydrolysis testing as described herein is likely applicable to polyamide and PVA geosynthetics), additional investigation will be required to establish a detailed protocol and acceptance criteria for these other polymers. These other polymers may be considered for evaluation using this protocol once modifications to the chemical/biological durability aspects of this protocol have been developed and are agreed upon by the approval authority (e.g., WSDOT).

The product specific durability studies for the determination of RFD should be conducted in, or if necessary extrapolated to, the chemical/biological environment anticipated in the reinforced backfill. The anticipated temperature of the environment is also a key variable in assessing the durability of a given product, as temperature can have an exponential effect on the rate of product property change. For the purposes of these guidelines, the effective design temperature is defined as the temperature that is halfway between the average yearly air temperature and the normal daily air temperature for the highest month at the wall site. Higher design temperatures may need to be considered for structures with southern exposures. The effective design temperature will be assumed to be 20°C (68°F), if the design temperature is not specifically identified in the contract specifications. Therefore, determine RFD at a temperature of 20°C (68°F) as a minimum. Determination of RFD at higher temperatures is optional.

Standards are currently not available for determining the effect of chemical/biological activity on long-term geosynthetic reinforcement strength. However, long-term product specific durability testing can be conducted in a manner that is likely to produce safe results. Geosynthetic durability can be evaluated using either retrieval and testing of geosynthetics in actual installations, or through long-term accelerated laboratory testing. Use of field retrieval data from actual installations requires that the baseline, in terms of tensile strength before and immediately after installation, and possibly other properties, be known with certainty, and that the observation period be of sufficient length to permit extrapolation to the desired design life. The field retrieval approach is in general fraught with practical and technical difficulties (see Allen and Elias, 1996, Elias, 2001a, and Elias, 2001b).

Furthermore, long periods of time may be needed for polyolefins to establish trends that can be extrapolated due to the presence of antioxidants, as no loss in strength will be observed until the antioxidants are used up. Elias (2001b) suggests that 30 years of in-service time may be required to obtain adequate observational data for polyolefins, and even PET products may require 20 years of in-service observations or more to accomplish this. Because of the very long observation periods required, long-term laboratory durability testing is the more practical approach to dealing with the durability issue. An overview of an appropriate laboratory testing approach for each geosynthetic polymer type is provided.
For polyolefin products in which the fibers/ribs do not exhibit micro-cracks or crazes as manufactured, long-term chemical durability testing may consist of elevated temperature oven aging tests to evaluate potential for oxidation effects (FHWA, 1997; Elias, et al., 1997; Salman, et al., 1998; Elias, et al., 1999). A magnification of x2,000 to x3,000 may be needed to observe whether or not micro-cracks or crazes are present (Salman, et al., 1997). If micro-cracks or crazes are present, elevated temperatures may significantly affect the molecular structure of polyolefins in the vicinity of the micro-cracks/crazes, making extrapolation of elevated temperature oxidation behavior to the behavior of the as-manufactured product at ambient temperatures very difficult (Salman, et al., 1998). For polyolefins in which the fibers/ribs exhibit micro-cracks or crazes, a means other than elevated temperature may be needed to accelerate oxidation behavior. Salman, et al. (1998) suggest that elevated oxygen concentration and pressure at ambient temperature may prove to be an effective accelerator in lieu of elevated temperature, especially for geosynthetic products in which the fibers/ribs exhibit micro-cracks or crazes as manufactured. In addition, Schröder et al (2002) have performed extensive testing and development of a protocol utilizing elevated oxygen pressure to address this issue. However, long-term validation of the protocol and final development of the protocol as a test standard are yet to be performed.

If oven aging tests are conducted, a forced air oven is recommended to keep the environment inside the oven as uniform as possible during the entire test duration, considering the long durations that are likely to be required. Temperature uniformity inside the oven should be maintained at ± 1%. An oven with horizontal air flow is recommended. Specimens should be placed in the oven parallel to the air flow and spaced no closer together than 13 mm (0.5 in.) apart (Allen and Elias, 1996; Elias, et al., 1999). The specimens should not be framed to prevent shrinkage, if shrinkage occurs, as doing this will create load in the specimen, making the resulting data difficult to interpret. Note that oxidation testing using forced air ovens will produce conservative estimates of long-term product strength due to the rapid air circulation and the relatively high oxygen content in the oven relative to the oxygen content in the ground. If the geosynthetic supplier wishes to submit data at a lower oxygen content than that of air (approximately 21% O₃), use of such data for approval of a given product may require that the structure be declared experimental, requiring that instrumentation be placed in the ground to verify the actual oxygen content in the structure backfill. Alternatively, the geosynthetic supplier could submit data from previously constructed structures with similar backfill in which the actual oxygen content in the structure backfill was measured.

For polyesters, long-term chemical durability testing should consist of elevated temperature immersion tests to evaluate potential for hydrolysis effects. A reactor similar to that illustrated in Figure 1 is recommended for incubating the geosynthetic specimens. A description of the test protocol is provided by Elias, et al. (1999). The reactor should be capable of maintaining temperature uniformity (± 1%) and stability during long-term use. A minimum solid/liquid ratio of 1:40 should be used to size the reactor and to determine the maximum number of specimens that can be placed in the reactor. Measures should be taken to minimize possibilities for oxidation during the long-term incubation (e.g., replace any air inside the reactor with nitrogen, use de-aired water, etc.). Specimens should be suspended in the solution on a hanger made of a material that will not react with or contaminate the immersion fluid and specimens (e.g., Teflon, stainless steel, etc.). The specimens should not be framed to prevent shrinkage, if shrinkage occurs so that an unknown amount of tension is not placed on the specimens. Specimens should each be separated by a distance of at least 13 mm (0.5 in.). The solution should be intensively stirred to ensure solution uniformity. For coated polyester products, the immersion tests should be conducted without the coating or the coated specimen ends should not be recoated (i.e., the ends of the core polymer should be left exposed to the immersion liquid). Elevated temperatures should be used to accelerate the degradation process, which allows the data to be extrapolated to the desired design life. Hydrolysis data should be submitted for the product at a pH of approximately 7 (i.e., neutral conditions -
distilled water), at a pH of 9 or more, and at a pH of 4 or less to facilitate the determination of RF.D. RF.D should be determined at a pH of 7 and at an alkaline pH (i.e., a pH of 9) as a minimum. If very acidic soils are anticipated (i.e., a pH near the bottom limit of pH = 4 for conditions defined as nonaggressive), RF.D should be determined at a pH of 4 as well.

Note that EPA 9090 testing, or the ASTM equivalent (ASTM D5322), is not considered adequate for a laboratory testing program to provide an estimate of RF.D. However, EPA 9090 or ASTM D5322 testing can be used as a first cut screening tool. That is, if any significant degradation of the strength of the product in question is observed for the chemical environment tested, the product would be disqualified for use in that chemical environment unless longer term testing conducted in accordance with this appendix is performed. EPA 9090 testing (or ASTM D5322) could also be used verify the effects of certain environmental variables which are known, based on the literature, to not significantly affect the given material. For example, low or high pH is known, based on the literature, to have little effect on polyolefins. This type of testing could be used to verify that the low or high pH does not affect the tensile strength of a polyolefin product, to allow that product to be used in environments that have a pH outside the range defined as a nonaggressive environment.

The long-term chemical durability testing program should be conducted and interpreted using the following guidelines:

- Incubation temperatures for the testing should be high enough to adequately accelerate the degradation process but below any major transitions in polymer behavior (e.g., glass transition, melting). Maximum recommended test temperatures to avoid major transitions are on the order of 70 to 750°C for polypropylene, high density polyethylene, and polyester, except as discussed above for polyolefin products which have micro-cracks or crazes as manufactured. However, exposure to temperatures near the upper end of these ranges could affect the stress-strain behavior of the material due to loss of molecular orientation, or possibly other effects that are not the result of chemical degradation. Therefore, care needs to be exercised when interpreting results from strength testing after exposure to temperatures near the maximum test temperatures indicated above. In general, if the stiffness of the material after exposure to the environment is significantly different from that of the virgin material, the stress-strain properties, and possibly the strength, of the material may have been affected by the exposure temperature in addition to the chemical environment. If the stiffness has been affected, the cause of the stiffness change should be thoroughly investigated to determine whether or not the change in stiffness is partially or fully due to the effect of temperature, or alternatively not use the data obtained at and above the temperature where the stiffness was affected. It is additionally recommended that the Arrhenius plot of the data be checked for linearity (see the discussion of Arrhenius modeling which follows). As a minimum, two to three data points above and below the suspected transition should be obtained and the plot checked for linearity through the entire range of temperatures, if it is desired to validate the use of data above the suspected transition for Arrhenius modeling and extrapolation purposes.

- A minimum of three to four test temperatures are recommended, typically spaced monotonically at 10°C increments (e.g., see ASTM D3045), except as discussed above for some polyolefin products.

- At the lowest test temperature (e.g., 30 to 50°C), incubation times of 2 to 4 years should be anticipated to get data adequate for long-term extrapolation.

- Enough retrievals (e.g., a minimum of three to four retrievals) should be made at a given test condition to adequately define the property loss as a function of incubation time.

- As a minimum, degradation should be tracked using the tensile strength of the specimens retrieved from the incubation chambers. Full wide width (ASTM D4595 or ASTM D6637) specimens are preferred; however, single rib or yarn specimens can be used.
• It is also recommended that degradation be tracked by chemical means, if possible, as well as through the use of scanning electron microscope (SEM) micrographs to verify the significance of the mechanical property degradation observed.

• The statistical variation of the measured properties after degradation are likely to be greater than what would be observed for the virgin material. This may require that the number of specimens per retrieval be greater than what the property variation for the virgin material would indicate.

• Extrapolation of chemical durability data for polymers typically utilizes an Arrhenius approach, though there is evidence which suggests that the Arrhenius model does not always work well for geosynthetics. Assuming Arrhenius modeling is appropriate, the slope of the strength degradation versus time plots (transformed mathematically to be linear through zero, first, or second order Arrhenius equations) can be used to characterize the degradation behavior as a function of temperature, allowing the slope at the desired design temperature to be estimated through the Arrhenius extrapolation. The equation order which best fits the data should be used (see Salman, et. al., 1998, for details). The strength retained at a given time at the design temperature can be calculated directly from the linear equation with the extrapolated slope. Note that Arrhenius modeling could also be conducted as a function of reactant (oxygen) concentration and pressure instead of temperature (Shelton and Bright, 1993; Salman, et. al., 1998). The extrapolation concept would be similar to that used for temperature. See Shelton and Bright (1993), Salman, et. al. (1997), and Salman, et. al. (1998), Elias, et al. (1999), Elias (2001a) for guidance on Arrhenius modeling techniques as applied to geosynthetics. Also note that since the extrapolation is being conducted over several log cycles of time, uncertainty in the data should be considered when determining the retained strength at the design life and design environment.

• For polyolefin oxidation, Arrhenius modeling will likely need to be conducted in two steps, as there are two main phases in the oxidation process for polyolefins: 1) the induction phase, where antioxidant consumption is the primary activity and little, if any, product strength loss occurs, and 2) the main polymer oxidation phase, where oxidative degradation of the polyolefin occurs, resulting in strength loss, and can generally be described by the kinetics of a Basic Auto-oxidation Scheme (Salman, et. al., 1997; Elias, et al., 1999). An Arrhenius model for the first phase should be developed so that the induction period $t_{ind}$ at the design temperature or reactant (i.e., oxygen) concentration can be estimated. A second Arrhenius model should then be developed using only the data after the induction period, and time in this case would begin at the end of the induction period at each temperature or reactant concentration tested. This second Arrhenius model is then extrapolated to the design temperature or reactant concentration to estimate the strength loss anticipated at the desired design life minus the induction period. Analysis of the remaining antioxidant content provides an additional method of measuring the duration of Step 1, particularly at lower temperatures and long durations, since changes in the antioxidant content take place ahead of the reduction in strength. Note that if the estimated induction period at the design environment is greater than the desired design life, this second phase Arrhenius modeling is unnecessary.

• Once the tensile strength at a given design life and design temperature has been estimated from the test data, determine $RF_D$ as follows:

$$RF_D = \frac{T_{tot}}{T_D}$$  \hspace{1cm} (D-1)
where, $T_{\text{lot}}$ is the average lot specific ultimate tensile strength for the lot of material used for the durability testing, and $T_D$ is the extrapolated (i.e., to the required design life) lot specific tensile strength after degradation based on the laboratory aging tests. In no case should $R_{FD}$ be less than 1.1.

Biological degradation has not proved a serious factor in the service life of geosynthetics. This is because the high molecular weight polyethylene, polyester, polypropylene and polyamide used are not easily broken down by bacteria and fungi. The high tensile strength of soil reinforcements prevents them from damage by roots of burrowing animals such as rabbits. For this reason it is not in general necessary to apply consider biological degradation in defining $R_{FD}$. However, the possibility of biological degradation should be reviewed if new polymers other than those described are used. Biological durability, if specifically requested by the approval authority, should be evaluated based on ASTM D3083-89, except the test should be modified to use ASTM D4595 or ASTM D6637 as the tensile test method. If any significant tensile strength loss is observed, as determined using ASTM D4595 or ASTM D6637, additional longer term testing should be performed before the product is further considered for use in reinforcement applications (see Bright, 1993).

If, as an alternative to long-term laboratory testing, the geosynthetic manufacturer prefers to submit data from long-term field retrievals, the following requirements must be met:

- A minimum of three retrievals must be obtained over a minimum time period adequate to permit extrapolation to the required design life (i.e., the first retrieval is taken right after installation, the second retrieval is taken at some time during the middle of this period, and the third retrieval is taken at the end of the minimum study period).
- The retrieved samples shall be evaluated both physically and chemically to determine changes in strength, fiber/yarn/rib appearance, and polymer chemistry. Lot specific baseline data for the exhumed geosynthetic regarding strength and polymer properties must be available for comparison. The soil environment in which the geosynthetic was placed must also be well defined in terms chemistry, moisture, temperature, gradation, and approximate geosynthetic stress level. Enough specimens for each retrieval must be taken to account for statistical variance in the properties measured. See Elias (2001a, 2001b) for a more detailed description of the procedures required to obtain the needed information to determine $R_{FD}$ using this approach.
- The polymer and physical characteristics of the exhumed material must meet the requirements for use in determining $R_{FD}$ for “similar” products as described later in this appendix.

**Note 1:** Chemical degradation of geosynthetics is a result of both environmental factors and polymer compositional factors. For a given polymer type, one can expect the greatest amount of chemical degradation to occur, in general, with polymers which have low molecular weights, low percent crystallinity, low density, and low draw ratios (Elias, 1990). Polymer additives also influence the degradation rate. Regarding environmental factors, one can expect the greatest amount of degradation to occur, in general, at relatively high temperatures, in moderate to high moisture conditions, in soils which are active chemically (especially in terms of pH and certain types of ions present), and with the geosynthetic under stress (Elias, 1990). Thickness of the polymer fibers may also have a strong influence on the degradation rate, as degradation mechanisms are dependent on diffusion processes or exposure and removal of surface material (Wrigley, 1987). Key chemical degradation mechanisms in typical soil environments include oxidation, hydrolysis, and environmental stress cracking (ESC).
The oxidation reaction can either be initiated by ultraviolet (UV) radiation or thermal energy. The rate of oxidation will be governed by chemical rate kinetics and by the rate of diffusion of the oxygen, the antioxidants and the degradation products. Diffusion (or migration) is frequently the controlling factor. Where the geosynthetic is buried, thermally activated oxidation is of most interest. UV activated oxidation is of importance only where the geosynthetic is exposed to light, such as at the wall face. In general, the wall face is covered from light with a permanent facing. Of the polymers used in geosynthetics, relatively speaking, polypropylene (PP) is potentially the most susceptible to oxidation, followed by high density polyethylene (HDPE) and polyester (PET) which have a relatively low susceptibility. Though in actuality a rather complex reaction, conceptually, polyolefin (i.e., PP and HDPE) oxidation is the reaction of free radicals within the polymer with oxygen, resulting in breakdown and/or crosslinking of the molecular chains and embrittlement of the polymer.

Antioxidants are typically added to the polymer (sometimes multiple types are added to increase effectiveness) to prevent oxidation during processing and use. Broad classes of antioxidants often used in geosynthetics include phenolic stabilizers and hindered amine light stabilizers (HALS). As the antioxidants are used up, resistance of the polymer to oxidation will decrease. The rate of polymer oxidation is dependent on how much and what type of antioxidant is present initially, at what rate it is used up, and how well it is distributed within the polymer (Van Zanten, 1986). Environmental factors which affect the rate of oxidation include temperature, oxygen concentration which in soil can vary from 21% in gravels at shallow depth to on the order of 1% in fine-grained soils at deeper depths (Yanful, 1993; Yanful, et. al., 1993), and the presence of transition metal ions such as iron (most common) or copper which act as a catalyst and accelerate the oxidation reaction. Thermal oxidation at typical in-soil temperatures appears to be quite slow.

Of the polymers mentioned above, only PET is potentially susceptible to hydrolysis. Hydrolysis occurs when water molecules react with the polymer molecules, resulting in chain scission, reduced molecular weight, and strength loss. Hydrolysis is simply the very slow inverse reaction of the synthesis of PET when water is present. The specifics of the reaction vary depending on the pH of the liquid. This results in high pH (alkaline) hydrolysis being relatively rapid, whereas neutral or low pH conditions can result in a slow hydrolysis rate. The rate of hydrolysis is also highly temperature dependent and can become relatively rapid at high temperatures in the vicinity of the glass transition temperature or above for the polymer, which is on the order of 70°C to 80°C (160°F to 180°F). The polymer does not need to be submerged for hydrolysis to occur, as hydrolysis can occur in moderate to high humidity conditions, though the reaction rate becomes slower as the humidity decreases (McMahon, et. al., 1959).

Hydrolysis appears to be the result of both a surface erosional phenomenon as well as a diffusional process of water to the polymer fiber core. These two phenomena have given rise to the terms “outer” and “inner” hydrolysis. Outer, or surficial, hydrolysis is dominant in high pH conditions and is characterized by loss in fiber cross-sectional area with minimal reduction in the molecular weight of the polymer that remains (Anderson, et. al., 1992; Jailloux, et. al., 1992). Inner, or diffusional controlled hydrolysis is dominant in neutral and acidic conditions and is characterized by significant losses in molecular weight of the polymer with minimal surficial erosion or damage (Anderson, et. al., 1992; Jailloux, et. al., 1992).

Rupture of a polymer when under stress is either ductile or brittle in nature. The ductile failure mode occurs when stresses are high enough to cause tie molecules to stretch out, lamellae to separate and start unfolding, resulting in fracture of the spherulites and plastic flow of the molecular structure (Lustiger, 1983). When failure occurs in a brittle manner, stress levels are usually lower, allowing sufficient time for tie molecules to slowly disentangle themselves from adjacent spherulites, initiating crack formation followed by slow crack growth (Bright, 1993).
ESC is the result of an accelerated crack initiation and growth process occurring when a polymer is subjected concurrently to a particular chemical environment and long-term stress. This accelerated crack initiation and growth process can result in premature brittle failure. ESC results in molecular chain disentanglement rather than chain breakage or chemical change.

Evaluation of ESC has been focused on polyethylene due to its use in various critical applications (e.g., telephone transmission cables, natural gas pipe) and the potential sensitivity of some polyethylenes to this phenomenon. The literature indicates that other polymers used for geosynthetics may also experience some sensitivity to this phenomenon (Bright, 1993).

The results of previous studies show that polyethylene resistance to ESC can be improved by increasing its average molecular weight, decreasing its molecular weight distribution, increasing its crystalline content, reducing the crystallite and/or spherulite size, increasing the degree of orientation, and using copolymerization (Wrigley, 1987). Therefore, the potential for ESC in a given polymer can be controlled.

ESC is closely related to the more general phenomenon of stress cracking. The difference between the two is that the chemical present for ESC accelerates the chain disentanglement process, whereas in stress cracking no accelerating chemical is present. Chemicals identified in the literature that can accelerate the stress cracking process include water, weak acids and bases, alcohols, metallic soaps, and solvents (Wrigley, 1987). Water, and to some extent weak acids and bases, are “chemicals” which need to be considered for ESC in typical in-soil environments.

An extensive long-term laboratory study of oxidation and hydrolysis as applied to geosynthetic reinforcement has been carried out (Elias, et al., 1999). Field studies have been carried out to evaluate many of the longer-term installations in which some baseline data was available (Elias, 2001b). In most cases, degradation has been minimal, if it is even significant enough to be measurable. These laboratory studies appear to corroborate the lack of degradation that has been observed in the field (Elias, 2001b). The protocols for oxidation and hydrolysis evaluation provided in this appendix, as well as the durability evaluation criteria provided elsewhere in Standard Practice T925, have been developed based on results from the laboratory and field studies conducted by Elias, et al. (1999) and Elias (2001b).

Use of Durability Data from “Similar” Products

Long-term chemical/biological durability data obtained from tests performed on older product lines, or other products within the same product line, may be applied to new product lines, or a similar product within the same product line, if one or both of the following conditions are met:

- The chemical and physical characteristics of tested products and proposed products are shown to be similar. Research data, though not necessarily developed by the product manufacturer, should be provided which shows that the minor differences between the tested and the untested products will result in equal or greater chemical/biological degradation resistance for the untested products.
- A limited testing program is conducted on the new or similar product in question and compared with the results of the previously conducted full testing program.

For polyolefins, similarity could be judged based on molecular weight and structure of the main polymer (i.e., is the polymer branched or crosslinked, is it a homopolymer or a blend, percent crystallinity, etc.), percentage of material reprocessed, tenacity of the fibers and processing history, and polymer additives used (i.e., type and quantity of antioxidants or other additives used). For polyesters and polyamides, similarity could be judged based on molecular weight or intrinsic viscosity of the main polymer, carboxyl end group content, percent crystallinity, or other molecular
structure variables, tenacity of the fibers and processing history, percentage of material reprocessed or recycled, and polymer additives used (e.g., pigments, etc.). The untested products should also have a similar macrostructure (i.e., woven, nonwoven, extruded grid, needlepunched, yarn structure, etc.) and fiber dimensions (e.g., thickness) relative to the tested products. It should be noted that percent crystallinity is not a controlled property and there is presently no indication of what an acceptable value for percent crystallinity should be.

For chemical durability evaluation, the limited testing program could consist of laboratory aging tests with a 1,000 to 2,000 hour incubation period in the same environment used for the full testing program conducted previously, conducted at a temperature near but slightly below any major property transitions. These limited durability test results must show that the durability performance of the new or similar product is equal to or better than the performance of the product previously tested. If so, the results from the full testing program on the older or similar product could be used for the new/similar product. If not, then a full testing and evaluation program for the new product should be conducted.

References


Elias, V., 2001a, Corrosion/Degradation of Soil Reinforcements for Mechanically Stabilized Earth Walls and Reinforced Soil Slopes, FHWA-NHI-00-044, Federal Highway Administration, Washington, D.C.


**WSDOT Test Method T926**

**Geogrid Brittleness Test**

**Purpose and Scope**

The purpose of this test to determine whether or not the product is likely to exhibit brittle behavior (rapid crack growth) when bent. This test is applicable to geogrids made of High Density Polyethylene (HDPE), polypropylene (PP) fiberglass, or other relatively stiff geogrid materials.

**Equipment**

The testing device shall have two planes tangent to a 0.5±0.025 inch radius curve fixed at a 90±2 degree angle, such as shown in Figure 1.

![Figure 1. Geogrid bend test setup.](image)

**Sampling and Specimens**

Sampling shall be in conformance to WSDOT Test Method 914.

Six specimens shall be taken at random locations throughout the sample. The specimens shall be one rib wide, with a minimum length of 12 inches, and approximately 2 inches of the cross-ribs shall remain attached to the rib being tested. No specimen should be located closer than 6 inches from the edge of the sample.

**Procedure**

The laboratory shall be 70±4 degrees F. Samples shall be allowed to remain at the laboratory temperature for a minimum of 1 hour before testing.

Place the geogrid rib on the bending template as shown in Figure 1. Apply the top cover plate on the geogrid, and apply uniform pressure on the plate and the geogrid specimen. Once the upper plate is in position and fixed, place the vertical cover plate over the geogrid, applying uniform pressure, causing the geogrid rib to conform to the 0.5 inch radius in approximately 3 seconds. Hold the geogrid in place for a minimum of 10±1 seconds.
If any geogrid specimen breaks or otherwise exhibits cracking the full width of the rib or the cracking extends into the cross-rib at the bend at any time during the 10 second hold period, the geogrid sample has failed the test.

Report

Identify the number of specimens that failed, if any, in the sample. Close-up photographs of any failed specimen shall be taken and maintained at the testing lab until the contract is completed.
Performance Exam Checklist

Test Method T926
Geogrid Britteness Test

Participant Name __________________________ Exam Date __________

Procedure Element Yes No
1. The tester has a copy of the current procedure on hand? □ □
2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present? □ □
3. Was the sample taken per WSDOT TM 914? □ □
4. Were six specimens properly prepared? □ □
5. Were the specimens allowed to remain at 70±4 degrees F for 1 hour min.? □ □
6. Were the specimens placed on the template with top plate on top? □ □
7. Were the specimens bent over the 0.5 radius using a vertical plate and uniform pressure applied? □ □
8. Geogrid held in place for 10±1 second? □ □
9. Geogrid inspected for cracking? □ □
10. Number of specimens failing reported with photos? □ □
13. All calculations performed correctly? □ □

First attempt: Pass □ Fail □ Second attempt: Pass □ Fail □

Signature of Examiner ____________________________

Comments:
________________________________________________________________________
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**WSDOT Test Method for ASTM C 939**

**Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)**

This standard is issued under the fixed designation C 939; 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 (e) indicates an editorial change since the last revision or reapproval. *This specification has been approved for use by agencies of the Department of Defense.*

1. **SCOPE**

1.1 This test method covers a procedure, used both in the laboratory and in the field, for determining the time of efflux of a specified volume of fluid hydraulic cement grout through a standardized flow cone and used for preplaced-aggregate (PA) concrete; however, the test method may also be used for other fluid grouts.

1.2 It is for use with neat grout and with grouts containing fine aggregate all passing a No. 8 (2.36-mm) sieve.

1.3 This test method is intended for use with grout having an efflux time of 35 s or less.

1.4 When efflux time exceeds 35 s, flowability is better determined by flow table, found in Test Method C 109, using 5 drops in 3 s.

1.5 The values stated in SI units are to be regarded as the standard.

1.6 *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:*

C 109/C109M Test Method for Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or 50-mm Cube Specimens)

C 938 Practice for Proportioning Grout Mixtures for Preplaced-Aggregate Concrete

3. **SUMMARY OF TEST METHOD**

3.1 The time of efflux of a specified volume of grout from a standardized flow cone is measured.

4. **SIGNIFICANCE AND USE**

4.1 This test method is applicable to the determination of the fluidity of various fluid grout mixtures.

5. **INTERFERENCES**

5.1 The presence of solid particles retained on the No. 8 (2.36-mm) sieve or lumps of unmixed material in the grout may cause the grout to flow unevenly through the discharge tube of the flow cone or stop the flow completely. Uneven flow will result in slower transit of the grout, thereby indicating a false consistency.

1This Test Method is based on ASTM C 939-97.
6. **APPARATUS**

6.1 *Flow Cone*, with dimensions as shown in Figure 1. The discharge tube shall be stainless steel. The body can be stainless steel, cast aluminum, or other essentially noncorroding metal.

*Note 1:* Cones with high-density polyethylene bodies are acceptable for field use in situations where precision as described in this test method is not required.

6.2 *Receiving Container*, capacity 2000 mL, minimum.

6.3 *Ring Stand* or other device, capable of supporting the flow cone in a vertical, steady position over the receiving container.

6.4 *Level*, carpenter’s or similar.

6.5 *Stop Watch*, least reading of not more than 0.2 s.

6.6 *Grout Mixer*, conforming to Practice C 938.

7. **TEST SAMPLE**

7.1 The grout test sample shall be in excess of 1725 mL and shall be representative of the grout in the mixer.

7.2 When sampling and testing is being done for the purpose of proportioning or comparing mixes or for qualifying materials, the temperature of the dry materials and mixing water shall be such that the temperature of the freshly mixed grout is 73.4 ± 3°F (23 ± 1.7°C), unless otherwise specified.

8. **CALIBRATION OF APPARATUS**

8.1 Mount the flow cone firmly in such a manner that it is free of vibration. Level the top to assure verticality. Close the outlet of the discharge tube with a finger or a stopper. Introduce 1725 ± 5 mL of water into the cone. Adjust the point gage to indicate the level of the water surface. Then allow the water to drain.

8.2 Before first use of the flow cone with grout and periodically thereafter, check the accuracy of the cone by filling it with water as described in 8.1. After checking or adjusting the point gage, start the stop watch and simultaneously remove the finger. Stop the watch at the first break in the continuous flow of water. The time indicated by the stop watch is the time of efflux of water. If this time is 8.0 ± 0.2 s, the cone may be used for determining the time of efflux of grout.

*Note:* It is imperative that the water be completely still prior to allowing it to flow from the cone, any movement will cause the time of efflux to increase.

9. **PROCEDURE**

9.1 Moisten the inside of the flow cone by filling the cone with water and, 1 min before introducing the grout sample, allow the water to drain from the cone. Close the outlet of the discharge tube with a finger or a stopper. Introduce the grout into the cone until the grout surface rises to contact the point gage, start the stop watch, and simultaneously remove the finger or stopper. Stop the watch at the first break in the continuous flow of grout from the discharge tube, then look into the top of the cone; if the grout has passed sufficiently, such that light is visible through the discharge tube, the time indicated by the stop watch is the time of efflux of the grout. If light is not visible through the discharge tube, then the use of the flow cone is not applicable for grout of this consistency. At least two tests having times of efflux within 1.8 s of their average shall be made for each grout mixture.
9.2 The test for time of efflux shall be made within 1 min of drawing of the grout from the mixer or transmission line. When grout is being placed over a significant period of time, the time of efflux may be determined at selected intervals to demonstrate that the consistency is suitable for the work.

10. REPORT

10.1 Report the following information:

10.1.1 Identification of sample,

10.1.2 Identification of materials in the sample, the proportions, and whether laboratory prepared or taken from the field production mix,

10.1.3 Average time of efflux to nearest 0.2 s and time interval from completion of mixing at which the test was made, and

*Note 2:* Other means of indicating grout level may be used as long as accurate indication of grout level on volume is obtained.

10.1.4 Temperature, ambient and of the sample at the time of test.

11. PRECISION AND BIAS

11.1 Precision — The following within-laboratory, multiple-operator precision applies. The single laboratory standard deviation has been found to be 0.88 s. Therefore, results from two properly conducted tests on the same material should not differ by more than 2.49 s.

11.2 Bias — No statement on bias can be prepared because there are no standard reference materials.

12. KEYWORDS

12.1 flow cone; grout; preplaced—aggregate concrete; time of efflux
### Performance Exam Checklist

**Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)**

**FOP FOR ASTM C 939**

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td></td>
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<tr>
<td>2. All equipment is functioning according to the test procedure,</td>
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<tr>
<td>and if required, has the current calibration/verification tags present?</td>
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<tr>
<td>3. Is the grout that is being tested a “fluid grout?”</td>
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<tr>
<td>4. Will the grout pass through a No. 8 (2.36 mm) sieve?</td>
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<tr>
<td>5. Is the cone set level and vibration free?</td>
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<tr>
<td>6. Is the grout test sample in excess of 1.8 quarts and representative of the</td>
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<tr>
<td>grout being produced?</td>
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<tr>
<td>7. Is the grout being produced at the specified temperature (73.4 \pm 3\ F)?</td>
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<tr>
<td>8. Does the tester have a verified stopwatch capable measuring to a time of 0.2</td>
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<td>sec.?</td>
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<td>9. Was the water calibration performed prior to use and is there a record of the</td>
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<tr>
<td>previous calibrations for this cone?</td>
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<tr>
<td>10. Was adjustment of the level indicator required?</td>
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</tbody>
</table>

**Note:** The calibration with water of a volume of \(1725\ mL \pm 5\ mL\) is to be 8 seconds \(\pm 0.2\) seconds to be considered valid for acceptance.

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>11. Was the cone filled with water a minute prior to introducing grout?</td>
<td></td>
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<tr>
<td>12. Water drained and cone outlet closed with a stopper/finger then grout</td>
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<tr>
<td>introduced into the cone until the grout surface rises to contact the point</td>
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<tr>
<td>gauge?</td>
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<tr>
<td>13. Stopwatch started as stopper/finger is removed and then stopped and</td>
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<tr>
<td>then stopped at the first break in continuous flow is observed?</td>
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<tr>
<td>14. Immediately observe to see if discharge tube is clear and light is visible</td>
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<tr>
<td>through it?</td>
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<tr>
<td>15. Repeat procedure and determine if the second observed flow rate is within</td>
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<tr>
<td>1.8 s of the average of the two flow rates.</td>
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<tr>
<td>Procedure Element</td>
<td>Yes</td>
<td>No</td>
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</tr>
<tr>
<td>16. Record the average time of efflux to the nearest 0.2 seconds</td>
<td>☐</td>
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</tr>
<tr>
<td>17. All calculations performed correctly?</td>
<td>☐</td>
<td>☐</td>
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</tbody>
</table>

First attempt: Pass ☐ Fail ☐
Second attempt: Pass ☐ Fail ☐

Signature of Examiner ________________________________

Comments:

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WSDOT FOP For ASTM D 1186

Nondestructive Measurement of Dry Film Thickness of Nonmagnetic Coatings Applied to a Ferrous Base

1. SCOPE

1.1 These test methods cover the nondestructive measurement of the dry film thickness of nonmagnetic coatings applied over a ferrous base material using commercially available test instruments. The test methods are intended to supplement manufacturers’ instructions for the manual operation of the gages and are not intended to replace them. They cover the use of instruments based on magnetic measuring principles only. Test Method A provides for the measurement of films using mechanical magnetic pull-off gages and Test Method B provides for the measurement of films using magnetic electronic gages.

1.2 These test methods are not applicable to coatings that will be readily deformable under the load of the measuring instruments, as the instrument probe must be placed directly on the coating surface to take a reading.

1.3 The values given in SI units of measurement are to be regarded as the standard. The values 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:

D 609 Practice for Preparation of Cold-Rolled Steel Panels for Testing Paint, Varnish, Conversion Coatings, and Related Coating Products

D 823 Practices for Producing Films of Uniform Thickness of Paint, Varnish, and Related Products on Test Panels

2.2 Steel Structures Painting Council Standard:

SSPC-PA2 Measurement of Dry Paint Thickness with Magnetic Gages

TEST METHOD A—MAGNETIC PULL-OFF GAGES

3. SUMMARY OF TEST METHOD

3.1 Instruments complying with this test method measure thickness by using a spring calibrated to determine the force required to pull a magnet from a ferrous base coated with a nonmagnetic film. The instrument must be placed directly on the coating surface to take a reading.

3.2 The attractive force of the magnet to the substrate varies inversely with the thickness of the applied film. The spring tension required to overcome the attraction of the magnet to the substrate is shown on the instrument scale as the distance (in mils or microns) between the magnet and the substrate.

4. SIGNIFICANCE AND USE

4.1 Many coating properties are markedly affected by the thickness of the dry film such as adhesion, corrosion protection, flexibility, and hardness. To be able to compare results obtained by different operators, it is essential to know film thickness.

1 This FOP is based on ASTM D 1186-01
4.2 Most protective and high performance coatings are applied to meet a requirement or a specification for the dry-film thickness of each coat, or for the complete system, or both. Coatings must be applied within certain minimum and maximum thicknesses to fill their expected function. In addition to potential performance deficiencies, it is uneconomical to apply more material than necessary when coating large areas. This test method is used to measure film thickness of coatings on ferrous metals.

5. APPARATUS

5.1 **Permanent Magnet**, small, either attached directly to a coil spring (“pencil” gage) or to a horizontal lever arm that is attached to a helical spring (“dial-type” gage). Increasing force is applied to the magnet by extending the coil spring in the first case or turning a graduated dial that coils the helical spring in the second. The readings obtained are shown directly on the instrument scale.

5.2 **Coating Thickness Standards**, with assigned values traceable to national standards are available from several sources, including most manufacturers of coating thickness gages.

1 Available from SSPC: The Society for Protective Coatings, 40 24th St., Sixth Floor, Pittsburgh, PA 15222–4643 (see www.sspc.org).

6. TEST SPECIMENS

6.1 When this test method is used in the field, the specimen is the coated structure or article on which the thickness is to be evaluated.

6.2 For laboratory use, apply the material to be tested to panels of similar roughness, shape, thickness, composition and magnetic properties on which it is desired to determine the thickness.

NOTE 1—Applicable test panel description and surface preparation methods are given in Practice D 609.

NOTE 2—Coatings should be applied in accordance with Practices D 823 or as agreed upon between the contracting parties.

7. VERIFICATION OF CALIBRATION OF APPARATUS

7.1 Different gage manufacturers follow different methods of calibration adjustment. Verify calibration according to manufacturer’s instructions.

7.2 The section of the type of standards used to verify calibration should be predicated upon which type provides the best and most appropriate calibration considering: type of gage, sample surface geometry, and contract requirements. Appendix X1 provides information helpful to making an informed selection of standards.

7.3 Following the manufacturer’s operating instructions, measure the thickness of a series of calibration standards covering the expected range of coating thickness. To guard against measuring with an inaccurate gage, recheck the gage at regular intervals. That interval should be set by agreement between contracting parties and maintained throughout the control process.

NOTE 3—Generally “Dial-type” instruments can be used in any position, while “pencil-type” instruments may be used in the vertical position only unless they have separate indicators for the horizontal and vertical positions. Follow the manufacturer’s recommendations.

8. PROCEDURE

8.1 Use the instrument only after calibration has been verified in accordance with Section 7.

8.2 Ensure that the coating is dry prior to use of the instrument.
8.3 Inspect the probe tip and surface to be measured to ensure that they are clean. Adherent magnetic filings or other surface contaminants will affect gage readings.

8.4 Take readings in locations free of electrical or magnetic fields. The location should also be free of vibration when using mechanical magnetic pull-off instruments.

8.5 The accuracy of the measurement can be influenced when made within 25 mm (1 in.) of the edge or right angle in the sample.

8.6 Measure the coating, following the manufacturer’s instructions.

8.7 Verify calibration periodically to ensure that the instrument continues to read properly. If the instrument is found to be out of adjustment, remeasure the thicknesses taken since the last satisfactory calibration check was made.

8.8 Take a sufficient number of readings to characterize the surface.

8.8.1 For laboratory measurements, a recommended minimum is three for a 75 by 150-mm (3 by 6-in.) panel and more in proportion to size.

8.8.2 For field measurements, a recommended minimum is five determinations at random for every 10 m² (100 ft²) of surface area. Each of the five determinations should be the mean of three separate gage readings within the area of a 4-cm (1.5-in.) diameter circle.

8.9 Make measurements at least 13 mm (1/2 in.) away from any edge or corner of the specimen. If it is necessary to measure closer than 13 mm (1/2 in.), verify the effect (if any), the edge has on the measurement.

**NOTE 4**—For additional information describing the number of measurements to be taken on large structures, and on non-smooth surfaces, refer to SSPC PA-2.

9. **REPORT**

9.1 Report the following information:

9.1.1 Instrument used, serial number,

9.1.2 Range, and mean of the thickness readings, and

9.1.3 Depending upon the application, record the individual readings as well.

**Report the information on the attached form.**

Material represented by the test specimens when tested under this method and found to meet the specified minimum coating thickness may be accepted. Any specimens which does not meet the minimum coating thickness will not be retested using this test method. Samples of the material will be submitted to either the Eastern Region Consolidated Materials Laboratory or the State Material laboratory for referee testing in accordance with AASHTO T 65.
# Field Report of Thickness of Nonmagnetic Coating on a Ferrous Base

<table>
<thead>
<tr>
<th>Contract</th>
<th>Bid Item No</th>
<th>Item</th>
<th>Specimen No.</th>
<th>Specification</th>
<th>Coating Thickness Required</th>
<th>Surface area of test specimen</th>
<th>Test represents</th>
<th>Instrument Serial No.</th>
<th>Calibration Date</th>
<th>Tested by</th>
<th>Date</th>
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<table>
<thead>
<tr>
<th>Reading No.</th>
<th>Test Location</th>
<th>Reading</th>
<th>Avg Readings</th>
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</table>

**Average**
10. **PRECISION AND BIAS**

10.1 A new round-robin study was performed recently. Data are being analyzed statistically. When completed, the required “Repeatability and Reproducibility” sections of this test method will be written and the round-robin study documented in an ASTM research report.

10.2 *Bias*—The bias for Test Method A of this standard for measuring dry film thickness cannot be determined because each instrument has its own bias.

**TEST METHOD B—ELECTRONIC GAGES**

11. **SUMMARY OF TEST METHOD**

11.1 Instruments complying with this test method measure thicknesses by placing a probe on the coated surface and use electronic circuitry to convert a reference signal into coating thickness.

11.2 Instruments of this type determine, within the probe or the instrument itself, changes in the magnetic flux caused by variations in the distance between the probe and the substrate.

12. **APPARATUS**

12.1 The testing apparatus shall be an electrically operated instrument utilizing a probe that houses a permanent magnet or coil energized by alternating current that is placed directly on the surface. The coating thickness is shown on the instrument’s display.

12.2 Coating thickness standards with assigned values traceable to national standards are available.

13. **TEST SPECIMENS**

13.1 See Section 6.

14. **CALIBRATION OF APPARATUS**

14.1 See Section 7.

15. **PROCEDURE**

15.1 See Section 8. Exclude steps 8.5 and 8.7.

16. **REPORT**

16.1 See Section 9.

17. **PRECISION AND BIAS**

17.1 *Precision*—See Section 10.

17.2 *Bias*—The bias for Test Method B of this standard for measuring dry film thickness cannot be determined because each instrument has its own bias.

18. **KEYWORDS**

18.1 coating thickness; dry film thickness; magnetic gages; nondestructive thickness; paint thickness

**APPENDIX**

X1. **CHARACTERISTICS AFFECTING GAGE READINGS**

X1.1 It is always good practice to ensure the reliability of gage readings by performing a verification test periodically, either before or after critical determinations. This practice ensures that, not only is the gage reading correct, but also that it is correctly calibrated to provide maximum accuracy of readings on the sample. Not all applications require this level of certainty so, while suggested, the inclusion of this practice is up to the contacting individuals to decide on implementation.
X1.2 Certain characteristics of samples may affect the accuracy of the calibrations. These include, but may not be limited to:

- X1.2.1 Surface profile of the substrate (roughness),
- X1.2.2 Surface profile of the coating,
- X1.2.3 Thickness of the substrate,
- X1.2.4 Geography of the sample surface (curves with small radii, small diameters, complex curves, etc.), and
- X1.2.5 Any characteristic that affects the magnetic or eddy current permeability of the substrate or coating, such as residual magnetism, or lack of homogeneity of magnetic characteristics.

X1.3 Calibration done on smooth, polished standards ensure that a gage can be properly calibrated, and that calibration is appropriate for any measurements on samples of the same characteristics, but it may not be the best for measurements of samples that differ from the calibration materials. When possible, verification should be done on samples of known thickness of coating applied to substrates as similar as possible to the sample to be tested.

X1.4 It is not practical to provide known thickness standards for all possible sample configurations. An alternative method is to verify calibration on a bare substrate as similar as possible to the sample, using a nonmagnetic metal foil, plastic shim or film of known thickness to simulate a coating.

X1.5 In using this verification of calibration method, it is necessary to be aware of additional characteristics that can affect the measured values. Plastic or brass shim stock typically has an inherent curve. This curve can act as a leaf spring and cause a magnetic pull-off gage to be “pushed” off the surface prematurely, resulting in an incorrect reading.

X1.6 With some materials and thickness, it is possible that the shim will not lie flat, which will also cause an erroneous reading. Various techniques exist to minimize this effect, such as mounting the shim in a holder that maintains tension on the shim to eliminate the tendency of the shim to curve.

X1.7 Other factors experienced with plastic shims, which are not usually present with painted or plated calibration standards include (but are not limited to):

- X1.7.1 Permanent creases in the shim due to folding,
- X1.7.2 Air entrapment between the shim and substrate,
- X1.7.3 Distortion due to environmental conditions, such as temperature, and
- X1.7.4 Shim thickness inconsistency due to the pressure of the probe tip. This may be a permanent “dimple” in the shim.

X1.8 Even with these factors affecting potential accuracy of plastic shims, in many applications, verification of calibration using plastic shims on the sample to be measured, can be a more appropriate (accurate) calibration than using plated or painted standards.

X1.9 No matter what standards are used, they should be periodically verified to ensure the assigned value is correct. Even metal coated on metal can wear or be damaged to an extent that readings are affected.
Performance Exam Checklist

Nondestructive Measurement of Thickness of Nonmagnetic Coatings on a Ferrous Base
FOP For ASTM D 1186

<table>
<thead>
<tr>
<th>Procedure Element</th>
<th>Test</th>
<th>Retest</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The tester has a copy of the current procedure on hand?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>3. Instrument calibrated in accordance with the manufacturer’s instructions before use employing a suitable thickness standard?</td>
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<tr>
<td>4. Several readings taken and recorded taking into account edge and curvature effects?</td>
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<tr>
<td>5. The average thickness converted to oz. ft² (g/m²) using appropriate conversion factor?</td>
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</tbody>
</table>

First attempt: Pass ☐ Fail ☐  Second attempt: Pass ☐ Fail ☐

Signature of Examiner __________________________________________

Comments:

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ASTM D 2938 (Modified)

Standard Test Method for Unconfined Compressive Strength of Intact Rock Core Specimens

ASTM D 2938 has been adopted by WSDOT with the following changes and/or additions:

The standard preparation of specimens for the above test will be as described below, which replaces Section 4 of ASTM D 2938:

4. Test Specimens
   4.1 Specimen arrives cut to length, ready for testing
   4.2 Specimens are measured for length to the nearest 0.01 in (0.25 mm) at the centers of the faces.
   4.3 Specimens are measured for diameter to the nearest 0.01 in (0.25 mm) by averaging two diameters measured at right angles to each other at about mid height of the specimen.
   4.4 The specimen ends are checked that they are at right angles to the longitudinal axis. The ends are recut if greater than 5 degrees different from a right angle.
   4.5 The specimen shall have a length to diameter ratio of 2.0 to 2.5 and the diameter shall be greater than 1 7/8 inch (48 mm).
ASTM D 4354 (Modified)

*Standard Practice for Sampling of Geosynthetics for Testing*

ASTM D 4354 has been adopted by WSDOT with the following changes and/or additions:

- Geotextiles tested in the Materials Laboratory will be conditioned in accordance with WSDOT Method T 915 in preparation for testing rather than the conditioning prescribed in the ASTM test procedures. This will be identified on all Geotextile Test Reports.
WSDOT FOP for ASTM D 4791

Standard Test Method for
Flat Particles, Elongated Particles, or Flat and Elongated Particles in Coarse Aggregate

1. Scope

1.1 This test method covers the determination of the percentages of flat particles, elongated particles, or flat and elongated particles in coarse aggregates.

1.2 The values stated in inch-pound units are to be regarded as the standard except in regard to sieve size and the size of aggregate, which are given in SI units in accordance with Specification E 11. The SI units in parentheses are for information purposes only.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

Note: WSDOT will be determining flat and elongated particles in accordance with section 8.4.

2. Referenced Documents

2.1 WSDOT Standards:
   T 2 WSDOT FOP for AASHTO for the Sampling of Aggregates
   T 248 WSDOT FOP for AASHTO for Reducing Field Samples of Aggregates to Testing Size
   T 27/11 WAQTC FOP for AASHTO for the Sieve Analysis of Fine & Coarse Aggregates & Materials Finer Than 75 mm (No. 200) in Mineral Aggregates by Washing

3. Terminology

3.1 Definitions:
   3.1.1 flat or elongated particles of aggregate—those particles of aggregate having a ratio of width to thickness or length to width greater than a specified value (see Terminology C 125).
   3.1.2 flat and elongated particles of aggregate—those particles having a ratio of length to thickness greater than a specified value.
   3.1.3 length—maximum dimension of the particle.
   3.1.4 width—maximum dimension in the plane perpendicular to the length
   3.1.5 thickness—maximum dimension perpendicular to the length and width.

4. Summary of Test Method

4.1 Individual particles of aggregate of specific sieve sizes are measured to determine the ratios of width to thickness, length to width, or length to thickness.

1This Test Method is Based on ASTM D 4791-99
5. Significance and Use

5.1 Flat or elongated particles of aggregates, for some construction uses, may interfere with consolidation and result in harsh, difficult to place materials.

5.2 This test method provides a means for checking compliance with specifications that limit such particles or to determine the relative shape characteristics of coarse aggregates.

6. Apparatus

6.1 The apparatus used shall be equipment suitable for testing aggregate particles for compliance with the definitions in 3.1, at the dimensional ratios desired.

6.1.1 Proportional Caliper Device—The proportional caliper devices illustrated in Fig. 1, Fig. 2, and Fig. 3 are examples of devices suitable for this test method. The device illustrated in Fig. 1 and Fig. 2 consists of a base plate with two fixed posts and a swinging arm mounted between them so that the openings between the arms and the posts maintain a constant ratio. The axis position can be adjusted to provide the desired ratio of opening dimensions. Fig. 1 illustrates a device on which ratios of 1:2, 1:3, 1:4, and 1:5 may be set. The device illustrated in Fig. 3 contains several fixed posts and has the capability of measuring various ratios simultaneously.

6.1.1.1 Verification of Ratio—The ratio settings on the proportional caliper device shall be verified by the use of a machined block, micrometer, or other appropriate device.

6.1.2 Balance—The balance or scales used shall be accurate to 0.5 % of the mass of the sample.

FIG. 1 Proportional Caliper
7. Sampling

7.1 Sample the coarse aggregate in accordance with FOP for AASHTO T2 Practice D 75. The mass of the field sample shall be the mass shown in FOP for AASHTO T2 Practice D 75.

7.2 Thoroughly mix the sample and reduce it to an amount suitable for testing using the applicable procedures described in FOP for AASHTO T 248 Practice C 702. The sample for test shall be approximately the mass desired when dry and shall be the end result of the reduction. Reduction to an exact predetermined mass shall not be permitted. The mass of the test sample shall conform to the following:

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<tr>
<th>Nominal Maximum Size*</th>
<th>Minimum Mass of Test Sample, lb (kg.)</th>
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<tr>
<td>3/8 (9.5)</td>
<td>2 (1)</td>
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<td>1/2 (12.5)</td>
<td>4 (2)</td>
</tr>
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<td>3/4 (19)</td>
<td>11 (5)</td>
</tr>
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<td>1 (25.0)</td>
<td>22 (10)</td>
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<td>1 1/2 (37.5)</td>
<td>33 (15)</td>
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<td>2 (50)</td>
<td>44 (20)</td>
</tr>
<tr>
<td>2 1/2 (63)</td>
<td>77 (35)</td>
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<tr>
<td>3 (75)</td>
<td>130 (60)</td>
</tr>
<tr>
<td>3 1/2 (90)</td>
<td>220 (100)</td>
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<tr>
<td>4 (100)</td>
<td>330 (150)</td>
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<tr>
<td>4 1/2 (112)</td>
<td>440 (200)</td>
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<tr>
<td>5 (125)</td>
<td>660 (300)</td>
</tr>
<tr>
<td>6 (150)</td>
<td>1100 (500)</td>
</tr>
</tbody>
</table>

* For aggregate, the nominal maximum size, (NMS) is the largest standard sieve opening listed in the applicable specification, upon which any material is permitted to be retained. For concrete aggregate, NMS is the smallest standard sieve opening through which the entire amount of aggregate is permitted to pass.

Note: For an aggregate specification having a generally unrestrictive gradation (i.e., wide range of permissible upper sizes), where the source consistently fully passes a screen substantially smaller than the maximum specified size, the nominal maximum size, for the purpose of defining sampling and test specimen size requirements may be adjusted to the screen, found by experience to retain no more than 5% of the materials.

8. Procedure

8.1 If determination by mass is required, oven dry the sample to constant mass at a temperature of 230 + 9°F (110 + 5°C). Dry in accordance with FOP for AASHTO T 255. If determination is by particle count, drying is not necessary.

8.2 Sieve the sample to be tested in accordance with FOP for AASHTO T 27/11 Test Method C 136. If the material retained on each required size (3/8 and larger) is more than 5% of the sample, reduce the material in accordance with FOP for AASHTO T 248 until approximately 100 particles are obtained for each required size. Using the material retained on the 9.5 mm (3/8 in.) or 4.75 mm (No. 4), as required by the specification being used, reduce each size-fraction present in the amount of 10% or more of the original sample in accordance with Practice C 702 until approximately 100 particles are obtained for each size-fraction required.
8.3  **Flat Particle Test and Elongated Particle Test**—Test each of the particles in each size fraction, and place in one of three groups: (1) flat, (2) elongated, (3) neither flat nor elongated.

8.3.1  Use the proportional caliper device, positioned at the proper ratio, shown in Fig. 2, as follows:

8.3.1.1  **Flat Particle Test**—Set the larger opening equal to the particle width. The particle is flat if the thickness can be placed in the smaller opening.

8.3.1.2  **Elongated Particle Test**—Set the larger opening equal to the particle length. The particle is elongated if the width can be placed within the smaller opening.

8.3.2  After the particles have been classified into the group described in 8.3, determine the proportion of the sample in each group by either count or by mass, as required.

8.4  **Flat and Elongated Particle Test**—Test each of the particles in each size fraction and place in one of two groups: (1) flat and elongated or (2) not flat and elongated.

8.4.1  Use the proportional caliper device, set at the desired ratio.

8.4.2  **Measurement**:

8.4.2.1  On proportional caliper devices similar to the devices shown in Fig. 1 and Fig. 2, set the larger opening equal to the length of the particle. The particle is flat and elongated if the particle, (biggest to smallest) when oriented to measure its thickness (biggest), can pass completely through the smaller opening of the caliper when it is rotated in any direction.

![FIG. 2 Use of Proportional Caliper](image)

<table>
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<tr>
<td>in.</td>
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</tr>
<tr>
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</tr>
<tr>
<td>3/16</td>
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<tr>
<td>1/4</td>
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<tr>
<td>5/16</td>
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<tr>
<td>3/8</td>
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</table>
FIG. 3 Proportional Caliper

8.4.2.2 On calipers similar to the one described in Fig. 3, set the minimum dimension of the proportional caliper device such that the particle, when oriented to measure its thickness, passes snugly between the post and swing arm. The particle is flat and elongated if the particle, when oriented to measure its length, fails to pass the desired large opening of the proportional caliper device.

8.4.3 After the particles have been classified into the groups described in 8.4, determine the proportion of the sample in each group by count or mass, as required.

Note: WSDOT performs this test by weight.

9. Calculation

9.1 Calculate the percentage of flat and elongated particles to the nearest 1 % for each sieve size than 3/8 in. and larger (9.5 mm), as required.

10. Report

10.1 Include the following information in the report:

10.1.1 Identification of the coarse aggregate tested, and

10.1.2 Grading of the aggregate sample, showing percentage retained on each sieve.

10.1.3 For flat particle tests and elongated particle tests:

10.1.3.1 Number of particles in each sieve size tested;

10.1.3.2 Percentages, calculated by number or by mass, or both, for: (1) flat particles, (2) elongated particles, and (3) total flat particles and elongated particles for each sieve size tested, and

10.1.3.3 The dimensional ratios used in the tests.
10.1.4 For flat and elongated particle tests:

10.1.4.1 Number of particles in each sieve size tested;

10.1.4.2 Percentages, calculated by number or by mass, or both, for flat and elongated particles for each sieve size tested,

10.1.4.3 The dimensional ratio used in the tests, and

10.1.5 When required, weighted average percentages based on the actual or assumed proportions of the various sieve sizes tested. Report the grading used for the weighted average if different from that in 10.1.2.

10.2 Report results using WSDOT form 350-161, or other report approved by the State Materials Engineer.

11. Precision and Bias

11.1 Precision—The precision of this test method is being determined.

11.2 Bias—Since there is no accepted reference material suitable for determining the bias for this test method, no statement on bias is being made.

12. Keywords

12.1 aggregates; coarse aggregates; particle shape
Performance Exam Checklist  
FLAT AND ELONGATED PARTICLES IN COARSE AGGREGATE  
FOP FOR ASTM D 4791

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### Procedure Element

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<td>2. All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?</td>
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<tr>
<td>3. Field sample obtained per AASHTO T-2?</td>
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<td>4. Sample thoroughly mixed prior to reducing to testing size?</td>
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<tr>
<td>5. Sample reduced to testing size per AASHTO T-248?</td>
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<td>6. Mass of the test sample conforms to the table in Section 7.2, ASTM D-4791?</td>
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### PROCEDURE

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<tr>
<td>2. Sample sieved per AASHTO T27/T11?</td>
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</tr>
<tr>
<td>3. Proportional caliper device positioned at proper ratio?</td>
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<tr>
<td>4. Each size fraction 3/8 inch and larger retaining more than 5% of the original sample reduced per AASHTO T-248 until approximately 100 particles are obtained for each size fraction required?</td>
<td></td>
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</tr>
<tr>
<td>5. Each particle of each size fraction tested for FLAT and ELONGATED using the proportional caliper device put in the appropriate group classification? (Flat &amp; Elongated or Not flat &amp; Elongated)</td>
<td></td>
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</tr>
<tr>
<td>6. Proportion of the sample of each sieve size determined by Mass?</td>
<td></td>
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<tr>
<td>7. Percent of Flat and Elongated particles figured to the nearest 1% for each sieve size?</td>
<td></td>
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<tr>
<td>8. Record number of particles in each sieve size tested?</td>
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<tr>
<td>9. Record percentages calculated by Mass?</td>
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<td>10. All calculations performed correctly?</td>
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First attempt:  Pass [ ] Fail [ ]  
Second attempt: Pass [ ] Fail [ ]

Signature of Examiner __________________________________________
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