

Materials Manual

M 46-01.29

January 2018

Americans with Disabilities Act (ADA) Information

English

Title VI Notice to Public It is the Washington State Department of Transportation's (WSDOT) policy to assure that no person shall, on the grounds of race, color, national origin or sex, as provided by Title VI of the Civil Rights Act of 1964, be excluded from participation in, be denied the benefits of, or be otherwise discriminated against under any of its federally funded programs and activities. Any person who believes his/her Title VI protection has been violated, may file a complaint with WSDOT's Office of Equal Opportunity (OEO). For additional information regarding Title VI complaint procedures and/or information regarding our non-discrimination obligations, please contact OEO's Title VI Coordinator at 360-705-7090.

Americans with Disabilities Act (ADA) Information This material can be made available in an alternate format by emailing the Office of Equal Opportunity at wsdotada@wsdot.wa.gov or by calling toll free, 855-362-4ADA(4232). Persons who are deaf or hard of hearing may make a request by calling the Washington State Relay at 711.

Spanish

Notificación de Titulo VI al Público Es la póliza de el Departamento de Transportes del Estado de Washington de asegurar que ninguna persona sea excluida de participación o sea negado los beneficios, o sea discriminado bajo cualquiera de sus programas y actividades financiado con fondos federales sobre la base de raza, color, origen nacional o sexo, como proveído por el Título VI de el Acto de Derechos Civiles de 1964. Cualquier persona que cree que sus protecciones de Titulo VI han sido violadas, puede hacer una queja con la Oficina de Igualdad de Oportunidades (OEO). Para información adicional con respecto a procedimientos de quejas de Titulo VI y/o información con respecto a nuestras obligaciones sin discriminación, por favor de comunicarse con el Coordinador de Titulo VI de la Oficina de Igualdad de Oportunidades (OEO) 360-705-7090.

Información del Acta Americans with Disabilities Act (ADA)

Este material es disponible en un formato alternative. Envie su petición por correo electrónico al equipo de Oficina de Igualdad de Oportunidades (OEO) en wsdotada@wsdot.wa.gov o llamando gratis, 855-362-4ADA (4232). Personas sordas o con problemas de audición pueden solicitar llamando el relé de estado de Washington al 711. For public events (like Open Houses, etc) English Accommodation requests for people with disabilities can be made by contacting the WSDOT Diversity/ADA Affairs team at wsdotada@wsdot.wa.gov or by calling toll-free, 855-362-4ADA (4232). Persons who are deaf or hard of hearing may make a request by calling the Washington State Relay at 711. Spanish Solicitudes de alojamiento para personas con discapacidad pueden hacer comunicándose con el equipo de Asuntos de diversidad/ADA WSDOT en wsdotada@wsdot.wa.gov o llamando al número gratuito, 855-362-4ADA (4232). Personas sordas o con problemas de audición pueden solicitar llamando el relé del estado de Washington al 711.

To get the latest information on WSDOT publications, sign up for individual email updates at www.wsdot.wa.gov/publications/manuals.

Washington State Department of Transportation Engineering and Regional Operations State Materials Laboratory PO Box 47365 Olympia, WA 98504-7365 The *Materials Manual* continues to use AASHTO, ASTM, WAQTC, and WSDOT test methods. The strategic direction 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 is using English units. The manual reflects the Quality System concerns of an AASHTO accredited organization and is 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 when a consensus standard (AASHTO, ASTM, or WAQTC) has been adapted to an FOP. The FOP provides the essential performance elements for the field technician.

When not specified by the test procedure, test reports will be generated through the Materials Testing System (MATS) or by the use of forms approved by the State Materials Engineer.

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 email at mawdslr@wsdot.wa.gov; 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 U.S. Postal Service mail.

This manual is updated as needed. To order printed copies, go to: www.wsdot.wa.gov/publications/manuals/orderinformation.htm

To view electronic copies, go to:

www.wsdot.wa.gov/publications/manuals/m46-01.htm

Kurt R. Williams, P.E. State Materials Engineer

				Aggregate
Procedure	<u> </u>	Field	In	1.93.13.11
Number	Owner	Use	Manual	Test Method
T 2	WAQTC	✓	✓	FOP for AASHTO T 2, Sampling of Aggregates
T 11	AASHTO			Materials Finer Than 0.075 mm (No. 200) Sieve in Mineral Aggregates by Washing
T 19	AASHTO	✓	✓	Bulk Density ("Unit Weight") and Voids in Aggregate (Rodding Procedure Only) (Checklist Only)
T 21	AASHTO			Organic Impurities in Fine Aggregates for Concrete
T 27	AASHTO			Sieve Analysis of Fine and Coarse Aggregates
T 27_T 11	WAQTC	✓	✓	FOP for AASHTO T 27_T 11, Sieve Analysis of Fine and Coarse Aggregates
T 37	AASHTO			Sieve Analysis of Mineral Filler
R 76	AASHTO			Reducing Samples of Aggregate to Testing Size
R 76	WAQTC	✓	✓	FOP for AASHTO R 76, Reducing Samples of Aggregate to Testing Size
T 84	AASHTO			Specific Gravity and Absorption of Fine Aggregates
T 85	AASHTO			Specific Gravity and Absorption of Coarse Aggregate
T 85	WAQTC	✓	✓	FOP for AASHTO T 85, Specific Gravity and Absorption of Coarse Aggregate
T 96	AASHTO			Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine
T 112	AASHTO		✓	Clay Lumps and Friable Particles in Aggregate
T 113	WSDOT		✓	Method of Test for Determination of Degradation Value
T 123	WSDOT	✓	✓	Method of Test for Bark Mulch
T 125	WSDOT		✓	Determination of Fiber Length Percentages in Wood Strand Mulch
T 126	WSDOT		✓	Determination of Fiber Length Percentages in Hydraulically-Applied Erosion Control Products
SOP 128	WSDOT	✓	✓	Sampling for Aggregate Source Approval
T 176	AASHT0			Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test
T 176	WAQTC	✓	✓	FOP for AASHTO T 176, Plastic Fines in Graded Aggregates and Soils by the Use of the Sand Equivalent Test
T 255	AASHTO			Total Evaporable Moisture Content of Aggregate by Drying
T 255	WAQTC	✓	✓	FOP for AASHTO T 255, Total Evaporable Moisture Content of Aggregate by Drying
T 288	AASHTO		✓	Determining Minimum Laboratory Soil Resistivity (Checklist Only)
T 289	AASHTO			Determining pH of Soil for Use in Corrosion
T 304	WSDOT	✓	✓	FOP for AASHTO for Uncompacted Void Content of Fine Aggregate
T 335	AASHTO			Determining the Percentage of Fracture in Coarse Aggregate
T 335	WAQTC	✓	✓	FOP for AASHTO T 335, Determining the Percentage of Fracture in Coarse Aggregate
T 417	WSDOT		✓	Method of Test for Determining Minimum Resistivity and pH of Soil and Water
T 716	WSDOT	✓	✓	Method of Random Sampling for Locations of Testing and Sampling Sites
				

				Bituminous Cement
Procedure		Field	ln	
Number	Owner	Use M	lanual	
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 44	AASHTO			Solubility of Bituminous Materials
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 59	AASHTO			Emulsified Asphalts
R 66	WAQTC	✓	✓	FOP for AASHTO R 66, Sampling Asphalt Materials
E 70	ASTM			pH of Aqueous Solutions With the Glass Electrode
T 72	AASHTO			Saybolt Viscosity
T 228	AASHTO			Specific Gravity of Semi-Solid Bituminous Material
T 240	AASHTO			Effect of Heat and Air on a Moving Film of Asphalt Binder (Rolling Thin-Film Oven Test)
T 301	AASHTO			Elastic Recovery Test of Asphalt Materials by Means of a Ductilometer
T 313	AASHTO			Determining the Flexural Creep Stiffness of Asphalt Binder Using the Bending Beam Rheometer (BBR)
T 315	AASHTO			Determining the Rheological Properties of Asphalt Binder Using a Dynamic Shear Rheometer (DSR)
T 316	AASHTO			Viscosity Determination of Asphalt Binder Using Rotational Viscometer
SOP 318	WSDOT		✓	Standard Operating Procedure for Melting of Flexible Bituminous Pavement Marker Adhesive for Evaluation
T 426	WSDOT		✓	Pull-Off Test for Hot Melt Traffic Button Adhesive
D 3111	ASTM			Standard Test Method for Flexibility Determination of Hot-Melt Adhesives by Mandrel Bend Test Method

				Hot Mix Asphalt
Procedure		Field		Total Markhard
Number	Owner	Use	Manual	
R 30	AASHTO			Practice for Short and Long Term Aging of Hot Mix Asphalt (HMA)
T 30	AASHTO			Mechanical Analysis of Extracted Aggregate
T 30	WAQTC	✓	√	FOP for AASHTO T 30, Mechanical Analysis of Extracted Aggregate
R 47	AASHTO			Standard Recommended Practice for Reducing Samples of Hot Mix Asphalt (HMA) to Testing Size
R 47	WAQTC	✓	✓	FOP for AASHTO R 47, Reducing Samples of Hot Mix Asphalt (HMA) to Testing Size
R 79	AASHTO			Vacuum Drying Compacted Asphalt Specimens
T 166	AASHTO			Bulk Specific Gravity of Compacted Asphalt Mixtures Using Saturated Surface-Dry Specimens
T 166	WAQTC	✓	✓	FOP for AASHTO T 166, Bulk Specific Gravity (Gmb) of Compacted Hot Mix Asphalt (HMA) Using Saturated Surface Dry Specimens
T 168	AASHTO			Sampling Bituminous Paving Mixtures
T 168	WAQTC	✓	✓	FOP for AASHTO T 168, Sampling of Hot Mix Asphalt Paving Mixtures
T 209	AASHTO			Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt (HMA)
T 209	WAQTC	✓	✓	FOP for AASHTO T 209, Theoretical Maximum Specific Gravity (Gmm) and Density of Hot Mix Asphalt (HMA) Paving Mixtures
T 269	AASHTO			Percent Air Void in Compacted Dense and Open Asphalt Mixtures
T 308	AASHTO			Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method
T 308	WAQTC	✓	✓	FOP for AASHTO T 308, Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method
T 312	AASHTO			Preparing and Determining the Density of Asphalt Mixture Specimens by Means of the Superpave Gyratory Compactor
T 312	WSDOT	✓	✓	FOP for AASHTO T 312, Asphalt Mixture Specimens by Means of the Superpave Gyratory Compactor
T 324	AASHTO		✓	Standard Method of Test for Hamburg Wheel-Track Testing of Compacted Hot Mix Asphalt (HMA)
T 329	AASHTO			Moisture Content of Asphalt Mixtures by Oven Method
T 329	WAQTC	✓	✓	FOP for AASHTO T 329, Moisture Content of Asphalt Mixtures by Oven Method
T 331	WSDOT		✓	Bulk Specific Gravity (G _{mb}) and Density of Compacted Hot Mix Asphalt (HMA) Using Automatic Vacuum Sealing Method
T 355	WAQTC	✓	✓	FOP for AASHTO T 355, In-Place Density of Asphalt Mixtures by Nuclear Method
T 716	WSDOT	✓	✓	Method of Random Sampling for Locations of Testing and Sampling Sites
T 718	WSDOT		✓	Method of Test for Determining Stripping of Hot Mix Asphalt
T 720	WSDOT		✓	Method of Test for Thickness Measurement of Hot Mix Asphalt (HMA) Cores
SOP 723	WSDOT		✓	Standard Operating Procedure for Submitting Hot Mix Asphalt (HMA) Mix Designs for Verification
T 724	WSDOT	✓	✓	Method of Preparation of Aggregate for Hot Mix Asphalt (HMA) Mix Designs
T 726	WSDOT	✓	✓	Mixing Procedure for Hot Mix Asphalt (HMA)
			-	

				Hot Mix Asphalt
Procedure Number	Owner	Field	In Manual	Test Method
SOP 728	WSDOT	√	√	Standard Operating Procedure for Determining the Ignition Furnace Calibration Factor (IFCF) for Hot Mix Asphalt (HMA)
SOP 729	WSDOT	✓	✓	Standard Operating Procedure for Determination of the Moving Average of Theoretical Maximum Density (TMD) for HMA
SOP 730	WSDOT	✓	✓	Standard Operating Procedure for Correlation of Nuclear Gauge Densities With Hot Mix Asphalt (HMA) Cores
SOP 731	WSDOT	✓	✓	Standard Operating Procedure for Determining Volumetric Properties of Hot Mix Asphalt
SOP 732	WSDOT	✓	✓	Standard Operating Procedure for Volumetric Design for Hot-Mix Asphalt (HMA)
SOP 733	WSDOT	✓	✓	Standard Operating Procedure for Determination of Pavement Density Differentials Using the Nuclear Density Gauge
SOP 734	WSDOT	✓	✓	Standard Operating Procedure for Sampling Hot Mix Asphalt After Compaction (Obtaining Cores)
SOP 735	WSDOT	✓	✓	Standard Operating Procedure for Longitudinal Joint Density
SOP 736	WSDOT		✓	In-Place Density of Bituminous Mixes Using Cores
SOP 737			✓	Procedure for the Forensic Testing of HMA Field Cores
D 6931	ASTM		✓	Standard Test Method for Indirect Tensile (IDT) Strength of Bituminous Mixtures

				Cement
Procedure)	Field	In	
Number	Owner	Use	Manua	
T 105	AASHTO			Chemical Analysis of Hydraulic Cement
T 106	AASHTO			Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)
T 106	WSDOT	✓	✓	FOP for AASHTO for Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)
T 107	AASHTO			Autoclave Expansion of Portland Cement
T 129	AASHTO			Normal Consistency of Hydraulic Cement
T 131	AASHTO			Time of Setting of Hydraulic Cement by Vicat Needle
T 133	AASHTO			Density of Hydraulic Cement
T 137	AASHTO			Air Content of Hydraulic Cement Mortar
T 153	AASHTO			Fineness of Hydraulic Cement by Air Permeability Apparatus
T 162	AASHTO			Mechanical Mixing of Hydraulic Cement Pastes and Mortars of Plastic Consistency
T 260	AASHTO			Sampling and Testing for Chloride Ion in Concrete and Concrete Raw Materials
T 303	AASHTO			Accelerated Detection of Potentially Deleterious Expansion of Mortar Bars Due to Alkali-Silica Reaction
T 313	WSDOT		✓	Method of Test for Cement-Latex Compatibility
T 314	WSDOT		✓	Method of Test for Photovolt Reflectance
T 413	WSDOT		✓	Method of Test for Evaluating Waterproofing Effectiveness of Membrane and Membrane-Pavement Systems
D 562	ASTM			Standard Test Method for Consistency of Paints Measuring Krebs Unit (KU) Viscosity Using a Stormer-Type Viscometer
T 813	WSDOT	✓	✓	Field Method of Fabrication of 2 in (50 mm) Cube Specimens for Compressive Strength Testing of Grouts and Mortars
T 814	WSDOT		✓	Method of Test for Water Retention Efficiency of Liquid Membrane- Forming Compounds and Impermeable Sheet Materials for Curing Concrete
C 939	WSDOT	✓	✓	FOP for ASTM for Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)

				Chemical
Procedure Number	Owner	Field Use	In Manual	Test Method
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 420	WSDOT	✓	✓	Test Method for Determining the Maturity of Compost (Solvita Test)
C 881	ASTM			Standard Specification for Epoxy-Resin-Base Bonding Systems for Concrete
C 882	ASTM		✓	Bond Strength (Diagonal Shear) (Checklist Only)
C 1218	ASTM			Standard Test Method for Water-Soluble Chloride in Mortar and Concrete
D 1429	ASTM			Standard Test Methods for Specific Gravity of Water and Brine
D 1475	ASTM			Test Method for Consistency of Paints Test Method for Density of Paint,
D 2628/ M 220	ASTM		✓	Test for High and Low Temperature Recovery of Elastomeric Joint Seals for Concrete Pavements
D 4758	ASTM			Test Method for Nonvolatile Contents of Latexes
D 5329	ASTM			Standard Test Methods for Sealants and Fillers, Hot-Applied, for Joints and Cracks in Asphaltic and Portland Cement Concrete Pavements
D 7091	ASTM	✓	✓	Nondestructive Measurement of Thickness of Nonmagnetic Coatings on a Ferrous Base (Checklist Only)

				Concrete
Procedure		Field	In	Took Makka d
Number	Owner		Manual	
TM 2	WAQTC	√	√	FOP for WAQTC TM 2, Sampling Freshly Mixed Concrete
T 22	AASHTO			Compressive Strength of Cylindrical Concrete Specimens
T 22	WSDOT	√	√	FOP for AASHTO for Compressive Strength of Cylindrical Concrete Specimens
T 23	AASHTO			Making and Curing Concrete Test Specimens in the Field
T 23	WAQTC	✓	✓	FOP for AASHTO T 23, Making and Curing Concrete Test Specimens in the Field
T 24	AASHTO			Obtaining and Testing Drilled Cores and Sawed Beams of Concrete
R 39	AASHTO			Making and Curing Concrete Test Specimens in the Laboratory
T 106	AASHTO			Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)
T 106	WSDOT	✓	✓	FOP for AASHTO for Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)
T 119	AASHTO			Slump of Hydraulic Cement Concrete
T 119	WAQTC	✓	✓	FOP for AASHTO T 119, Slump of Hydraulic Cement Concrete
T 121	WAQTC	✓	✓	FOP for AASHTO T 121, Density (Unit Weight), Yield, and Air Content (Gravimetric) of Concrete
C 140	ASTM			Standard Test Methods for Sampling and Testing Concrete Masonry Units and Related Units
T 141	AASHTO			Sampling Freshly Mixed Concrete
T 152	AASHTO			Air Content of Freshly Mixed Concrete by the Pressure Method
T 152	WAQTC	✓	✓	FOP for AASHTO T 152, Air Content of Freshly Mixed Concrete by the Pressure Method
T 196	AASHTO	-	√	Air Content of Concrete (Volumetric Method) (Checklist Only)
T 197	AASHTO			Time of Setting of Concrete Mixtures by Penetration Resistance
T 198	AASHTO			Splitting Tensile Strength of Cylindrical Concrete Specimens
T 231	AASHTO			Capping Cylindrical Concrete Specimens
T 231	WSDOT	√	√	FOP for AASHTO for Capping Cylindrical Concrete Specimens
T 260	AASHTO			Sampling and Testing for Chloride Ion in Concrete and Concrete Raw Materials
T 277	AASHTO			Electrical Indication of Concrete's Ability to Resist Chloride Ion Penetration
T 309	AASHTO			Temperature of Freshly Mixed Portland Cement Concrete
T 309	WAQTC	✓	✓	FOP for AASHTO T 309, Temperature of Freshly Mixed Portland Cement Concrete
C 457	ASTM			Standard Test Method for Microscopical Determination of Parameters of the Air-Void System in Hardened Concrete
C 495	ASTM			Test Method for Compressive Strength of Lightweight Insulated Concrete
T 716	WSDOT	√	√	Method of Random Sampling for Locations of Testing and Sampling Sites
T 802	WSDOT	✓	✓	Method of Test for Flexural Strength of Concrete (Using Simple Beam With Center-Point Loading)
C 805	ASTM		,	Test Method for Rebound Number of Hardened Concrete
				The state of the s

				Concrete
Procedure Number	Owner	Field Use	In Manual	Test Method
C 805	WSDOT	✓	✓	Rebound Hammer Determination of Compressive Strength of Hardened Concrete
T 808	WSDOT	✓	✓	Method for Making Flexural Test Beams
T 810	WSDOT	✓	✓	Method of Test for Determination of the Density of Portland Cement Concrete Pavement Cores
T 812	WSDOT	✓	✓	Method of Test for Measuring Length of Drilled Concrete Cores
T 813	WSDOT	✓	✓	Field Method of Fabrication of 2 in (50 mm) Cube Specimens for Compressive Strength Testing of Grouts and Mortars
T 818	WSDOT		✓	Air Content of Freshly Mixed Self-Compacting Concrete by the Pressure Method
T 819	WSDOT		✓	Making and Curing Self-Compacting Concrete Test Specimens in the Field
C 939	ASTM			Standard Test Method for Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)
C 939	WSDOT	✓	✓	FOP for ASTM for Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)
C 1218	ASTM			Standard Test Method for Water-Soluble Chloride in Mortar and Concrete
D 1429	ASTM			Standard Test Methods for Specific Gravity of Water and Brine
C 1611	WSDOT	✓	✓	FOP for ASTM C 1611/C 1611M Standard Test Method for Slump Flow of Self-Consolidating Concrete
C 1621	WSDOT	✓	✓	FOP for ASTM C 1621/C 1621M Standard Test Method for Passing Ability of Self-Consolidating Concrete by J-Ring

				Electrical and Traffic
Procedure Number	Owner	Field Use	In Manual	Test Method
IP 78-16	FHWA			Signal Controller Evaluation
T 257	AASHTO			Instrumental Photometeric Measurements of Retroreflectivie Material and Retroreflective
T 314	WSDOT		✓	Method of Test for Photovolt Reflectance
T 421	WSDOT		✓	Test Method for Traffic Controller Inspection and Test Procedure
T 422	WSDOT		✓	Test Method for Traffic Controller Transient Voltage Test (Spike Test) Procedure
T 423	WSDOT		✓	Test Method for Traffic Controller Conflict Monitoresting
T 424	WSDOT		✓	Test Method for Traffic Controller Power Interruption Test Procedure
T 425	WSDOT		✓	Test Method for Traffic Controller NEM and 170 Type Environmental Chamber Test
T 426	WSDOT		✓	Pull-Off Test for Hot Melt Traffic Button Adhesive
T 427	WSDOT		✓	Test Method for Loop Amplifier Testing Procedure
T 428	WSDOT		✓	Test Method for Traffic Controller Compliance Inspection and Test Procedure
SOP 429	WSDOT		✓	Methods for Determining the Acceptance of Traffic Signal Controller Assembly
DMCT 700	ATSI			Manual on Signal Controller Evaluation
PCMZ 2000	TS			Manual on Signal Controller Evaluation
D 4956	ASTM			Standard Specification for Retroreflective Sheeting for Traffic Control
TS1	NEMA			Signal Controller Evaluation Geotechnical – Soils

				Geotechnical – Soils
Procedure		Field		
Number	Owner	Use	Manual	
R 58	AASHTO			Dry Preparation of Disturbed Soil and Soil Aggregate Samples for Test
R 75	WAQTC	✓	√	FOP for AASHTO R 75, Developing a Family of Curves
T 88	AASHTO			Particle Size Analysis of Soils
T 89	AASHTO		√	Determining the Liquid Limit of Soils
T 90	AASHTO		√	Determining the Plastic Limit and Plasticity Index of Soils (Checklist Only)
T 99	AASHTO		,	Moisture-Density Relations of Soils Using a 5.5 lb (2.5 kg) Rammer and a 12 in (305 mm) Drop
T 99	WAQTC	✓	✓	FOP for AASHTO T 99, Moisture-Density Relations of Soils Using a 5.5 lb (2.5 kg) Rammer and a 12 in (305 mm) Drop
T 100	AASHTO			Specific Gravity of Soil
T 180	AASHTO			Moisture-Density Relations of Soils Using a 10 lb (4.54 kg) Rammer and an 18 in (457 mm) Drop Checklist
T 180	WAQTC	✓	✓	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
T 208	AASHTO			Unconfined Compressive Strength of Cohesive Soil
T 215	AASHTO			Permeability of Granular Soils (Constant Head)
T 216	AASHTO			One-Dimensional Consolidation Properties of Soils
T 236	AASHTO			Direct Shear Test of Soils Under Consolidated Drained Conditions
T 265	AASHTO			Laboratory Determination of Moisture Content of Soils
T 265	WAQTC	✓	✓	FOP for AASHTO T 255, Laboratory Determination of Moisture Content of Soils
T 296	AASHTO			Unconsolidated, Undrained Compressive Strength of Cohesive Soils in Triaxial Compression
T 297	AASHTO			Consolidated, Undrained Triaxial Compressive Test on Cohesive Soils Shear
D 2487	ASTM			Practice for Classification of Soils for Engineering Purposes (Unified Soil Classification System)
D 2488	ASTM			Practice for Description and Identification of Soils (Visual-Manual Procedure)
D 4186	ASTM			Standard Test Method for One-Dimensional Consolidation Properties of Saturated Cohesive Soils Using Controlled-Strain Loading
D 4644	ASTM			Standard Test Method for Slake Durability of Shales and Similar Weak Rocks
T 501	WSDOT		✓	Test Method to Determine Durability of Very Weak Rock
D 5084	ASTM			Standard Test Methods for Measurement of Hydraulic Conductivity of Saturated Porous Materials Using a Flexible Wall Permeameter
D 5311	ASTM			Standard Test Method for Load Controlled Cyclic Triaxial Strength of Soil
D 5731	ASTM			Standard Test Method for Determination of the Point Load Strength Index of Rock and Application to Rock Strength Classifications
D 6467	ASTM			Standard Test Method for Torsional Ring Shear Test to Determine Drained Residual Shear Strength of Cohesive Soils
D 6528	ASTM			Standard Test Method for Consolidated Undrained Direct Simple Shear Testing of Cohesive Soils
D 7012	ASTM		✓	Standard Test Method for Unconfined Compressive Strength of Intact Rock Core Specimens

				Geotextile and Steel
Procedure		Field		
Number	Owner	Use	Manual	Test Method
A 143	ASTM			Standard Practice for Safeguarding Against Embrittlement of Hot-Dip Galvanized Structural Steel Products and Procedure for Detecting Embrittlement
T 244	AASHTO			Mechanical Testing of Steel Products
A 370	ASTM			Standard Test Methods and Definitions for Mechanical Testing of Steel Products
F 606	ASTM			Mechanical Properties: Steel Fasteners
T 914	WSDOT	✓	✓	Practice for Sampling of Geosynthetic Material for Testing
T 915	WSDOT		✓	Practice for Conditioning of Geotextiles for Testing
T 923	WSDOT		✓	Thickness Measurement of Geotextiles
T 925	WSDOT		✓	Standard Practice for Determination of Long-Term Strength for Geosynthetic Reinforcement
T 926	WSDOT		✓	Geogrid Brittleness Test
D 1683	ASTM			Sewen Seams (Geotextiles)
D 4355	ASTM			Standard Test Method for Deterioration of Geotextiles From Exposure to Ultraviolet Light and Water (Xenon-Arc Type Apparatus)
D 4491	ASTM			Water Permeability (Geotextiles)
D 4533	ASTM			Tear Strength (Geotextiles)
D 4354	ASTM		✓	Standard Practice for Sampling of Geosynthetics for Testing
D 4595	ASTM			Wide Width Breaking Load (Geotextiles)
D 4632	ASTM			Grab Breaking Load (Geotextiles)
D 4751	ASTM			Apparent Opening Size (Geotextiles)
D 6241	ASTM			Puncture (Geotextiles)

			Paint
Procedure Number	Owner	Field In Use Manua	Test Method
D 185	ASTM		Standard Test Methods for Coarse Particles in Pigments, Pastes, and Paints
T 314	ASTM		Method of Test for Photovolt Reflectance
D 562	ASTM		Standard Test Method for Consistency of Paints Measuring Krebs Unit (KU) Viscosity Using a Stormer-Type Viscometer
D 1208	ASTM		Method for Determination of Loss on Ignition
D 1210	ASTM		Standard Test Method for Fineness of Dispersion of Pigment-Vehicle Systems by Hegman-Type Gage
D 1475	ASTM		Test Method for Density of Paint and Related Products
D 2244	ASTM		Standard Practice for Calculation of Color Tolerances and Color Differences From Instrumentally Measured Color Coordinates
D 2369	ASTM		Method for Determination of Volatile and Nonvolatile Content (Ordinary Laboratory Oven)
D 2371	ASTM		Standard Test Method for Pigment Content of Solvent-Reducible Paints (Centrifuge)
D 2621	ASTM		Standard Test Method for Infrared Identification of Vehicle Solids From Solvent-Reducible Paints
D 2697	ASTM		Standard Test Method for Volume Nonvolatile Matter in Clear or Pigmented Coatings
3011	FTMS		Method for Determination of Condition in Container
D 3723	ASTM		Standard Test Method for Pigment Content of Water Emulsion Paints by Temperature Ashing
4053	FTMS		Method for Determination of Nonvolatile Vehicle Content
4061	FTMS		Method for Determination of Drying Time (Oil-Based Paints)
4122	FTMS		Method for Determination of Hiding Power (Contrast Ratio)
D 4505	ASTM		Standard Specification for Preformed Plastic Pavement Marking Tape for Extended Service Life Pavement Soils

				Pavement Soils
Procedure Number	Owner	Field Use	In Manual	Test Method
T 242	AASHTO			Frictional Properties of Paved Surfaces Using a Full-Size Tire
T 272	AASHTO			One-Point Method for Determining Maximum Dry Density and Optimum Moisture
T 272	WAQTC	✓	✓	FOP for AASHTO T 272, One-Point Method for Determining Maximum Dry Density and Optimum Moisture
T 307	AASHTO		✓	Determining the Resilient Modulus of Soils and Aggregate Materials
T 310	WAQTC	✓	✓	FOP for AASHTO T 310, In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)
T 606	WSDOT		✓	Method of Test for Compaction Control of Granular Materials
T 610	WSDOT		✓	Method of Test for the Capillary Rise of Soils
SOP 615	WSDOT	✓	✓	Determination of the % Compaction for Embankment & Untreated Surfacing Materials Using the Nuclear Moisture-Density Gauge
T 807	WSDOT	✓	✓	Method of Operation of California Profilograph and Evaluation of Profiles
D 4694	ASTM			Test Method for Deflections With Falling-eight Type Impulse Load Device

	Standard Practice						
Procedure Number	Owner	Field In Use Manua	I Test Method				
QC 1	WSDOT	✓	Standard Practice for Cement Producers/Importers/Distributors That Certify Portland Cement and Blended Hydraulic Cement				
QC 2	WSDOT	✓	Standard Practice for Asphalt Suppliers That Certify Performance Graded and Emulsified Asphalts				
QC 3	WSDOT	✓	Quality System Laboratory Review				
QC 4	WSDOT	✓	Standard Practice for Fly Ash Producers/Importers/Distributors That Certify Fly Ash				
QC 5	WSDOT	✓	Standard Practice for Ground Granulated Blast-Furnace Slag Producers/ Importers/Distributors That Certify Ground Granulated Blast-Furnace Slag				
QC 6	WSDOT	✓	Annual Prestressed Plant Review and Approval Process				
QC 7	WSDOT	✓	Annual Precast Plant Review and Approval Process				
QC 8	WSDOT	✓	Standard Practice for Approval of Hot Mix Asphalt Mix Designs for the Qualified Products List				

				Numerical Order
Procedure		Field	In .	-
Number	Owner	Use	Manual	
QC 1	WSDOT		√	Standard Practice for Cement Producers/Importers/Distributors That Certify Portland Cement and Blended Hydraulic Cement
QC 2	WSDOT		✓	Standard Practice for Asphalt Suppliers That Certify Performance Graded and Emulsified Asphalts
QC 3	WSDOT		✓	Quality System Laboratory Review
QC 4	WSDOT		✓	Standard Practice for Fly Ash Producers/Importers/Distributors That Certify Fly Ash
QC 5	WSDOT		✓	Standard Practice for Ground Granulated Blast-Furnace Slag Producers/ Importers/Distributors That Certify Ground Granulated Blast-Furnace Slag
QC 6	WSDOT		√	Annual Prestressed Plant Review and Approval Process
QC 7	WSDOT		√	Annual Precast Plant Review and Approval Process
QC 8	WSDOT		✓	Standard Practice for Approval of Hot Mix Asphalt Mix Designs for the Qualified Products List
TS1	NEMA			Signal Controller Evaluation Geotechnical – Soils
T 2	WAQTC	√	√	FOP for AASHTO T 2, Sampling of Aggregates
TM 2	WAQTC	✓	√	FOP for WAQTC TM 2, Sampling Freshly Mixed Concrete
T 11	AASHTO			Materials Finer Than 0.075 mm (No. 200) Sieve in Mineral Aggregates by Washing
E 18	ASTM			Standard Test Methods for Rockwell Hardness of Metallic Materials
T 19	AASHTO	✓	✓	Bulk Density ("Unit Weight") and Voids in Aggregate (Rodding Procedure Only) (Checklist Only)
T 21	AASHTO			Organic Impurities in Fine Aggregates for Concrete
T 22	AASHTO			Compressive Strength of Cylindrical Concrete Specimens
T 22	WSDOT	✓	✓	FOP for AASHTO for Compressive Strength of Cylindrical Concrete Specimens
T 23	AASHTO			Making and Curing Concrete Test Specimens in the Field
T 23	WAQTC	✓	✓	FOP for AASHTO T 23, Making and Curing Concrete Test Specimens in the Field
T 24	AASHTO			Obtaining and Testing Drilled Cores and Sawed Beams of Concrete
T 27	AASHTO			Sieve Analysis of Fine and Coarse Aggregates
T 27_T 11	WAQTC	✓	✓	FOP for AASHTO T 27_T 11, Sieve Analysis of Fine and Coarse Aggregates
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
R 30	AASHTO			Practice for Short and Long Term Aging of Hot Mix Asphalt (HMA)
T 30	AASHTO			Mechanical Analysis of Extracted Aggregate
T 30	WAQTC	✓	✓	FOP for AASHTO T 30, Mechanical Analysis of Extracted Aggregate
T 37	AASHTO			Sieve Analysis of Mineral Filler
R 39	AASHTO			Making and curing Concrete Test Specimens in the Laboratory
T 44	AASHTO			Solubility of Bituminous Materials
R 47	AASHTO			Standard Recommended Practice for Reducing Samples of Hot Mix Asphalt (HMA) to Testing Size

				Numerical Order
Procedure Number	Owner	Field Use	In Manual	Test Method
R 47	WAQTC	✓	✓	FOP for AASHTO R 47, Reducing Samples of Hot Mix Asphalt (HMA) to Testing Size
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)
R 58	AASHTO			Dry Preparation of Disturbed Soil and Soil Aggregate Samples for Test
T 59	AASHTO			Emulsified Asphalts
T 65	AASHTO			Mass (Weight) of Coating on Iron and Steel Articles With Zinc or Zinc-Alloy Coatings
R 66	WAQTC	✓	✓	FOP for AASHTO R 66, Sampling Asphalt Materials
E 70	ASTM			pH of Aqueous Solutions With the Glass Electrode
T 72	AASHTO			Saybolt Viscosity
R 75	WAQTC			FOP for AASHTO R 75, Developing a Family of Curves
R 76	AASHTO			Reducing Samples of Aggregate to Testing Size
R 76	WAQTC	✓	✓	FOP for AASHTO R 76, Reducing Samples of Aggregate to Testing Size
IP 78-16	FHWA			Signal Controller Evaluation
R 79	AASHTO			Vacuum Drying Compacted Asphalt Specimens
T 84	AASHTO			Specific Gravity and Absorption of Fine Aggregates
T 85	AASHTO			Specific Gravity and Absorption of Coarse Aggregates
T 85	WAQTC	✓	✓	FOP for AASHTO T 85, Specific Gravity and Absorption of Coarse Aggregate
T 88	AASHTO			Particle Size Analysis of Soils
T 89	AASHTO		✓	Determining the Liquid Limit of Soils
T 90	AASHTO		✓	Determining the Plastic Limit and Plasticity Index of Soils (Checklist Only)
T 96	AASHTO			Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine
T 99	AASHTO			Moisture-Density Relations of Soils Using a 5.5 lb (2.5 kg) Rammer and a 12 in (305 mm) Drop
T 99	WAQTC	✓	✓	FOP for AASHTO T 99, Moisture-Density Relations of Soils Using a 5.5 lb (2.5 kg) Rammer and a 12 in (305 mm) Drop
T 100	AASHTO			Specific Gravity of Soil
T 105	AASHTO			Chemical Analysis of Hydraulic Cement
T 106	AASHTO			Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)
T 106	WSDOT	✓	✓	FOP for AASHTO for Compressive Strength of Hydraulic Cement Mortars (Using 2-in. or (50-mm) Cube Specimens)
T 107	AASHTO			Autoclave Expansion of Hydraulic Cement
T 112	AASHTO		✓	Clay Lumps and Friable Particles in Aggregate
T 113	WSDOT		✓	Method of Test for Determination of Degradation Value
T 119	AASHTO			Slump of Hydraulic Cement Concrete
T 119	WAQTC	✓	✓	FOP for AASHTO T 119, Slump of Hydraulic Cement Concrete

				Numerical Order
Procedure Number	Owner	Field Use	In Manual	Test Method
T 121	WAQTC	✓	✓	FOP for AASHTO T 121, Density (Unit Weight), Yield, and Air Content (Gravimetric) of Concrete
T 123	WSDOT	✓	✓	Method of Test for Bark Mulch
T 125	WSDOT		✓	Determination of Fiber Length Percentages in Wood Strand Mulch
T 126	WSDOT		✓	Determination of Fiber Length Percentages in Hydraulically-Applied Erosion Control Products
T 127	WSDOT		✓	Preparation of Leachate Sample for Testing Toxicity of HECP Effluent
SOP 128	WSDOT	✓	✓	Sampling for Aggregate Source Approval
T 129	AASHTO			Normal Consistency of Hydraulic Cement
T 131	AASHTO			Time of Setting of Hydraulic Cement by Vicat Needle
T 133	AASHTO			Density of Hydraulic Cement
T 137	AASHTO			Air Content of Hydraulic Cement Mortar
C 140	ASTM			Standard Test Methods for Sampling and Testing Concrete Masonry Units and Related Units
T 141	AASHTO			Sampling Freshly Mixed Concrete
A 143	ASTM			Standard Practice for Safeguarding Against Embrittlement of Hot-Dip Galvanized Structural Steel Products and Procedure for Detecting Embrittlement
T 152	AASHTO		,	Air Content of Freshly Mixed Concrete by the Pressure Method
T 152	WAQTC	✓	✓	FOP for AASHTO T 152, Air Content of Freshly Mixed Concrete by the Pressure Method
T 153	AASHTO			Fineness of Hydraulic Cement by Air Permeability Apparatus
T 162	AASHTO			Mechanical Mixing of Hydraulic Cement Pastes and Mortars of Plastic Consistency
T 166	AASHTO			Bulk Specific Gravity of Compacted Hot Mix Asphalt (HMA) Using Saturated Surface-Dry Specimens
T 166	WAQTC	✓	✓	FOP for AASHTO T 166, for Bulk Specific Gravity of Compacted Hot Mix Asphalt (HMA) Using Saturated Surface Dry Specimens
T 168	AASHTO			Sampling Bituminous Paving Mixtures
T 168	WAQTC	✓	✓	FOP for AASHTO T 168, Sampling of Hot Mix Asphalt Paving Mixtures
T 176	AASHTO			Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test
T 176	WAQTC	✓		FOP for AASHTO T 176, Plastic Fines in Graded Aggregates and Soils by the Use of the Sand Equivalent Test
T 180	AASHTO			Moisture-Density Relations of Soils Using a 10 lb (4.54 kg) Rammer and an 18 in (457 mm) Drop
T 180	WAQTC	✓	✓	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
D 185	ASTM			Standard Test Methods for Coarse Particles in Pigments, Pastes, and Paints
T 196	AASHTO		✓	Air Content of Concrete (Volumetric Method) (Checklist Only)
T 197	AASHTO			Time of Setting of Concrete Mixtures by Penetration Resistance
T 198	AASHTO			Splitting Tensile Strength of Cylindrical Concrete Specimens
T 208	AASHTO			Unconfined Compressive Strength of Cohesive Soil

				Numerical Order
Procedure Number	Owner	Field Use	In Manual	Test Method
T 209	AASHTO			Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt (HMA)
T 209	WAQTC	✓	✓	FOP for AASHTO T 209, Theoretical Maximum Specific Gravity (Gmm) and Density of Hot Mix Asphalt (HMA) Paving Mixtures
T 215	AASHTO			Permeability of Granular Soils (Constant Head)
T 216	AASHTO			One-Dimensional Consolidation Properties of Soils
T 228	AASHTO			Specific Gravity of Semi-Solid Bituminous Material
T 231	AASHTO			Capping Cylindrical Concrete Specimens
T 231	WSDOT	✓	✓	FOP for AASHTO for Capping Cylindrical Concrete Specimens
T 236	AASHTO			Direct Shear test of Soils Under Consolidated Drained Conditions
T 240	AASHTO			Effect of Heat and Air on a Moving Film of Asphalt Binder (Rolling Thin-Film Oven Test)
T 242	AASHTO			Frictional Properties of Paved Surfaces Using a Full-Size Tire
T 244	AASHTO			Mechanical Testing of Steel Products
T 255	AASHTO			Total Evaporable Moisture Content of Aggregate by Drying
T 255	WAQTC	✓	✓	FOP for AASHTO T 255, Total Evaporable Moisture Content of Aggregate by Drying
T 257	AASHTO			Instrumental Photometeric Measurements of Retroreflectivie Material and Retroreflective
T 260	AASHTO			Sampling and Testing for Chloride Ion in Concrete and Concrete Raw Materials
T 265	AASHTO		-	Laboratory Determination of Moisture Content of Soils
T 265	WAQTC	✓	✓	FOP for AASHTO T 255, Laboratory Determination of Moisture Content of Soils
T 267	AASHTO			Determination of Organic Content in Soils by Loss on Ignition
T 269	AASHTO			Percent Air Void in Compacted Dense and Open Asphalt Mixtures
T 272	AASHTO			One-Point Method for Determining Maximum Dry Density and Optimum Moisture
T 272	WAQTC	✓	✓	FOP for AASHTO T 272, One-Point Method for Determining Maximum Dry Density and Optimum Moisture
T 277	AASHTO			Electrical Indication of Concrete's Ability to Resist Chloride Ion Penetration
T 288	AASHTO		√	Determining Minimum Laboratory Soil Resistivity (Checklist Only)
T 289	AASHTO			Determining pH of Soil for Use in Corrosion
T 296	AASHTO			Unconsolidated, Undrained Compressive Strength of Cohesive Soils in Triaxial Compression
T 297	AASHTO			Consolidated, Undrained Triaxial Compressive Test on Cohesive Soils Shear
T 301	AASHTO			Elastic Recovery Test of Asphalt Materials by Means of a Ductilometer
T 303	AASHTO			Accelerated Detection of Potentially Deleterious Expansion of Mortar Bars Due to Alkali-Silica Reaction
T 304	WSDOT	√	√	FOP for AASHTO for Uncompacted Void Content of Fine Aggregate
T 307	AASHTO		✓	Determining the Resilient Modulus of Soils and Aggregate Materials
T 308	AASHTO			Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method

				Numerical Order
Procedure Number	Owner	Field Use	In Manual	Test Method
T 308	WAQTC	✓	✓	FOP for AASHTO T 308, Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method
T 309	AASHTO			Temperature of Freshly Mixed Hydraulic Cement Concrete
T 309	WAQTC	✓	✓	FOP for AASHTO T309, Temperature of Freshly Mixed Portland Cement Concrete
T 310	WAQTC	✓	✓	FOP for AASHTO T 310, In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)
T 312	AASHTO			Preparing and Determining the Density of Asphalt Mixture Specimens by Means of the Superpave Gyratory Compactor
T 312	WAQTC	✓	✓	FOP for AASHTO T 312, Asphalt Mixture Specimens by Means of the Superpave Gyratory Compactor
T 313	AASHTO			Determining the Flexural Creep Stiffness of Asphalt Binder Using the Bending Beam Rheometer (BBR)
T 313	WSDOT		✓	Method of Test for Cement-Latex Compatibility
T 314	WSDOT		✓	Method of Test for Photovolt Reflectance
T 315	AASHTO			Determining the Rheological Properties of Asphalt Binder Using a Dynamic Shear Rheometer (DSR)
T 316	AASHTO			Viscosity Determination of Asphalt Binder Using Rotational Viscometer
SOP 318	WSDOT		✓	Standard Operating Procedure for Melting of Flexible Bituminous Pavement Marker Adhesive for Evaluation
T 324	AASHTO		✓	Standard Method of Test for Hamburg Wheel-Track Testing of Compacted Hot Mix Asphalt (HMA)
T 329	AASHTO			Moisture Content of Asphalt Mixtures by Oven Method
T 329	WAQTC	✓	✓	FOP for AASHTO T 329, Moisture Content of Asphalt Mixture by Oven Method
T 331	WSDOT		✓	Bulk Specific Gravity (G _{mb}) and Density of Compacted Hot Mix Asphalt (HMA) Using Automatic Vacuum Sealing Method
T 335	AASHTO			Determining the Percentage of Fracture in Coarse Aggregate
T 335	WAQTC	✓	✓	FOP for AASHTO T 335, Determining the Percentage of Fracture in Coarse Aggregate
T 355	WAQTC	✓	✓	FOP for AASHTO T 355, In-Place Density of Asphalt Mixtures by Nuclear Method
A 370	ASTM			Standard Test Methods and Definitions for Mechanical Testing of Steel Products
T 413	WSDOT	✓	✓	Method of Test for Evaluating Waterproofing Efectiveness of Membrane and Membrane-Pavement Systems
T 417	WSDOT		✓	Method of Test for Determining Minimum Resistivily and pH of Soil and Water
T 420	WSDOT	✓	✓	Test Method for Determining the Maturity of Compost (Solvita Test)
T 421	WSDOT		✓	Test Method for Traffic Controller Inspection and Test Procedure
T 422	WSDOT		✓	Test Method for Traffic Controller Transient Voltage Test (Spike Test) Procedure
T 423	WSDOT		✓	Test Method for Traffic Controller Conflict Monitoresting
T 424	WSDOT		✓	Test Method for Traffic Controller Power Interruption Test Procedure

				Numerical Order
Procedure		Field	In Manual	Toot Mathad
Number	Owner	Use		
T 425	WSDOT		√	Test Method for Traffic Controller NEM and 170 Type Environmental Chamber Test
T 426	WSDOT		✓	Pull-Off Test for Hot Melt Traffic Button Adhesive
T 427	WSDOT		✓	Test Method for Loop Amplifier Testing Procedure
T 428	WSDOT		✓	Test Method for Traffic Controller Compliance Inspection and Test Procedure
SOP 429	WSDOT		✓	Methods for Determining the Acceptance of Traffic Signal Controller Assembly
T 432	WSDOT		✓	Flexibility Test for Hot-Melt Adhesives
C 457	ASTM			Standard Test Method for Microscopical Determination of Parameters of the Air-Void System in Hardened Concrete
C 495	ASTM			Test Method for Compressive Strength of Lightweight Insulated Concrete
T 501	WSDOT		✓	Test Method to Determine Durability of Very Weak Rock
D 562	ASTM			Standard Test Method for Consistency of Paints Measuring Krebs Unit (KU) Viscosity Using a Stormer-Type Viscometer
F 606	ASTM			Test Methods for Determining the Mechanical Properties of Externally and Internally Threaded Fasteners, Washers, Direct Tension Indicators, and Rivets
T 606	WSDOT		✓	Method of Test for Compaction Control of Granular Materials
T 610	WSDOT		✓	Method of Test for the Capillary Rise of Soils
SOP 615	WSDOT	✓	✓	Determination of the % Compaction for Embankment and Untreated Surfacing Materials Using the Nuclear Moisture-Density Gauge
DMCT 700	ATSI			Manual on Signal Controller Evaluation
T 716	WSDOT	✓	✓	Method of Random Sampling for Locations of Testing and Sampling Sites
T 718	WSDOT		✓	Method of Test for Determining Stripping of Hot Mix Asphalt
T 720	WSDOT		✓	Method of Test for Thickness Measurement of Hot Mix Asphalt (HMA) Cores
SOP 723	WSDOT		✓	Standard Operating Procedure for Submitting Hot Mix Asphalt (HMA) Mix Designs for Verification
T 724	WSDOT	✓	✓	Method of Preparation of Aggregate for Hot Mix Asphalt (HMA) Mix Designs
T 726	WSDOT	✓	✓	Mixing Procedure for Hot Mix Asphalt (HMA)
SOP 728	WSDOT	✓	✓	Standard Operating Procedure for Determining the Ignition Furnace Calibration Factor (IFCF) for Hot Mix Asphalt (HMA)
SOP 729	WSDOT	✓	✓	Standard Operating Procedure for Determination of the Moving Average of Theoretical Maximum Density (TMD) for HMA
SOP 730	WSDOT	✓	✓	Standard Operating Procedure for Correlation of Nuclear Gauge Densities With Hot Mix Asphalt (HMA) Cores
SOP 731	WSDOT	✓	✓	Standard Operating Procedure for Determining Volumetric Properties of Hot Mix Asphalt
SOP 732	WSDOT	✓	✓	Standard Operating Procedure for Volumetric Design for Hot-Mix Asphalt (HMA)
SOP 733	WSDOT	✓	✓	Standard Operating Procedure for Determination of Pavement Density Differentials Using the Nuclear Density Gauge

				Numerical Order
Procedure	0	Field	In	Took Makka d
Number	Owner		Manual	
SOP 734	WSDOT	√		Standard Operating Procedure for Sampling Hot Mix Asphalt After Compaction (Obtaining Cores)
SOP 735	WSDOT	✓	✓	Standard Operating Procedure for Longitudinal Joint Density
SOP 736	WSDOT		✓	In-Place Density of Bituminous Mixes Using Cores
SOP 737			✓	Procedure for the Forensic Testing of HMA Field Cores
T 802	WSDOT	✓	✓	Method of Test for Flexural Strength of Concrete (Using Simple Beam With Center-Point Loading)
C 805	ASTM			Test Method for Rebound Number of Hardened Concrete
C 805	WSDOT	✓		Rebound Hammer Determination of Compressive Strength of Hardened Concrete
T 807	WSDOT	✓	√	Method of Operation of California Profilograph and Evaluation of Profiles
T 808 T	WSDOT	✓	✓	Method for Making Flexural Test Beams
T 810	WSDOT	✓		Method of Test for Determination of the Density of Portland Cement Concrete Pavement Cores
T 812	WSDOT	✓	✓	Method of Test for Measuring Length of Drilled Concrete Cores
T 813	WSDOT	✓		Field Method of Fabrication of 2 in (50 mm) Cube Specimens for Compressive Strength Testing of Grouts and Mortars
T 814	WSDOT			Method of Test for Water Retention Efficiency of Liquid Membrane- Forming Compounds and Impermeable Sheet Materials for Curing Concrete
T 818	WSDOT			Air Content of Freshly Mixed Self-Compacting Concrete by the Pressure Method
T 819	WSDOT			Making and Curing Self-Compacting Concrete Test Specimens in the Field
C 881	ASTM			Standard Specification for Epoxy-Resin-Base Bonding Systems for Concrete
C 882	ASTM		✓	Bond Strength (Diagonal Shear) (Checklist Only)
T 914	WSDOT	✓	✓	Practice for Sampling of Geosynthetic Material for Testing
T 915	WSDOT		✓	Practice for Conditioning of Geotextiles for Testing
T 923	WSDOT		✓	Thickness Measurement of Geotextiles
T 925	WSDOT			Standard Practice for Determination of Long-Term Strength for Geosynthetic Reinforcement
T 926	WSDOT		✓	Geogrid Brittleness Test
C 939	ASTM			Standard Test Method for Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)
C 939	WSDOT	✓		FOP for ASTM for Flow of Grout for Preplaced-Aggregate Concrete (Flow Cone Method)
D 1208	ASTM			Test Methods for Common Properties of Certain Pigments (Loss on Ignition)
D 1210	ASTM			Standard Test Method for Fineness of Dispersion of Pigment-Vehicle Systems by Hegman-Type Gage
C 1218	ASTM			Standard Test Method for Water-Soluble Chloride in Mortar and Concrete
D 1429	ASTM			Standard Test Methods for Specific Gravity of Water and Brine
C 1437	ASTM			Standard Test Method for Flow of Hydraulic Cement Mortar

				Numerical Order
Procedure		Field		— (11 (1)
Number	Owner	Use	Manual	
D 1475	ASTM			Test Method for Consistency of Paints Test Method for Density of Paint, Varnish, Lacquer, and Related Products
C 1611	WSDOT	✓	✓	FOP for ASTM C 1611/C 1611M Standard Test Method for Slump Flow of Self-Consolidating Concrete
C 1621	WSDOT	✓	✓	FOP for ASTM C 1621/C 1621M Standard Test Method for Passing Ability of Self-Consolidating Concrete by J-Ring
D 1683	ASTM			Standard Test Method for Failure in Sewn Seams of Woven Apparel Fabrics
PCMZ 2000	TS			Manual on Signal Controller Evaluation
D 2240	ASTM			Standard Test Method for Rubber Property – Durometer Hardness
D 2244	ASTM			Standard Practice for Calculation of Color Tolerances and Color Differences From Instrumentally Measured Color Coordinates
D 2369	ASTM			Test Method for Volatile Content of Coatings (Ordinary Laboratory Oven)
D 2371	ASTM			Standard Test Method for Pigment Content of Solvent-Reducible Paints (Centrifuge)
D 2487	ASTM			Practice for Classification of Soils for Engineering Purposes (Unified Soil Classification System)
D 2488	ASTM			Practice for Description and Identification of Soils (Visual-Manual Procedure)
D 2621	ASTM			Standard Test Method for Infrared Identification of Vehicle Solids From Solvent-Reducible Paints
D 2628/ M 220	ASTM	✓	✓	Test for High and Low Temperature Recovery of Elastomeric Joint Seals for Concrete Pavements
D 2697	ASTM			Standard Test Method for Volume Nonvolatile Matter in Clear or Pigmented Coatings
3011	FTMS			Method for Determination of Condition in Container
D 3111	ASTM			Standard Test Method for Flexibility Determination of Hot-Melt Adhesives by Mandrel Bend Test Method
D 3723	ASTM			Standard Test Method for Pigment Content of Water Emulsion Paints by Temperature Ashing
4053	FTMS			Method for Determination of Nonvolatile Vehicle Content
4061	FTMS			Method for Determination of Drying Time (Oil-Based Paints)
4122	FTMS			Method for Determination of Hiding Power (Contrast Ratio)
D 4186	ASTM			Standard Test Method for One-Dimensional Consolidation Properties of Saturated Cohesive Soils Using Controlled-Strain Loading
D 4354	ASTM		√	Standard Practice for Sampling of Geosynthetics for Testing
D 4355	ASTM			Standard Test Method for Deterioration of Geotextiles From Exposure to Ultraviolet Light and Water (Xenon-Arc Type Apparatus)
D 4491	ASTM			Standard Test Methods for Water Permeability of Geotextiles by Permittivity
D 4505	ASTM			Standard Specification for Preformed Plastic Pavement Marking Tape for Extended Service Life
D 4533	ASTM			Standard Test Method for Trapezoid Tearing Strength of Geotextiles
D 4595	ASTM			Standard Test Method for Tensile Properties of Geotextiles by the Wide-Width Strip Method

				Numerical Order
Procedure		Field		_ ,
Number	Owner	Use	Manual	Test Method
D 4632	ASTM			Standard Test Method for Grab Breaking Load and Elongation of Geotextiles
D 4644	ASTM			Standard Test Method for Slake Durability of Shales and Similar Weak Rocks
D 4694	ASTM			Test Method for Deflections With Falling-Eight Type Impulse Load Device
D 4751	ASTM			Test Method for Determining Apparent Opening Size of a Geotextile
D 4758	ASTM			Test Method for Nonvolatile Contents of Latexes
D 4956	ASTM			Standard Specification for Retroreflective Sheeting for Traffic Control
D 5084	ASTM			Standard Test Methods for Measurement of Hydraulic Conductivity of Saturated Porous Materials Using a Flexible Wall Permeameter
D 5311	ASTM			Standard Test Method for Load Controlled Cyclic Triaxial Strength of Soil
D 5329	ASTM			Standard Test Methods for Sealants and Fillers, Hot-Applied, for Joints and Cracks in Asphaltic and Portland Cement Concrete Pavements
D 5731	ASTM			Standard Test Method for Determination of the Point Load Strength Index of Rock and Application to Rock Strength Classifications
D 6241	ASTM			Puncture (Geotextiles)
D 6467	ASTM			Standard Test Method for Torsional Ring Shear Test to Determine Drained Residual Shear Strength of Cohesive Soils
D 6528	ASTM			Standard Test Method for Consolidated Undrained Direct Simple Shear Testing of Cohesive Soils
D 6931	ASTM		✓	Standard Test Method for Indirect Tensile (IDT) Strength of Bituminous Mixtures
D 7012	ASTM		✓	Standard Test Method for Unconfined Compressive Strength of Intact Rock Core Specimens
D 7091	ASTM	✓	✓	Nondestructive Measurement of Thickness of Nonmagnetic Coatings on a Ferrous Base (Checklist Only)

WSDOT Standard Practice QC 1

Standard Practice for Approval of Portland Cement and/or Blended Hydraulic Cement Producers/Suppliers

1. Scope

This standard specifies requirements for all producers/suppliers of portland cement and/or blended hydraulic 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:
 - M-85 Standard Specifications for Portland Cement
 - M-240 Standard Specifications for Blended Hydraulic Cement
 - R-18 Establishing and Implementing a Quality System for Construction Materials Testing Laboratories
- 2.2 ASTM Standards
 - C-150 Standard Specification for Portland Cement
 - C-595 Standard Specification for Blended Hydraulic Cement
- 2.3 Agency's Standard Specifications

3. Terminology

- 3.1 AASHTO American Association of State Highway and Transportation Officials
- 3.2 ASTM American Society of Testing and Materials
- 3.3 CCRL Cement and Concrete Reference Laboratory
- 3.4 NIST National Institute of Standards and Technology
- 3.5 WSDOT Washington State Department of Transportation
- 3.6 Producer A production facility that has the capacity for producing and/or grinding portland cement and/or blended hydraulic cement meeting the requirements of the *Standard Specifications* Section 9-01.
- 3.7 Supplier A company that supplies portland cement and/or blended hydraulic cement that meets the requirements of *Standard Specifications* Section 9-01.

- 3.8 Specification Compliance Testing Complete testing in accordance with the specification requirements for the material identified.
- 3.9 Quality Management Plan The producer/supplier plan to ensure that the portland cement and/or blended hydraulic cement meets the specification requirements through systematic program of sampling, testing, and inspection.
- 3.10 Quality Control Testing Testing performed per the producer/supplier quality management plan to evaluate the production process.
- 3.11 CAP Cement Acceptance Program
- 3.12 Cement Mill Test Report A document provided by the producer showing the physical and chemical test results with specification limits for each property tested.
- 3.13 Cement Certificate of Analysis A document provided by the supplier showing the physical and chemical test results with specification limits for the properties tested on each shipment of imported portland cement or imported blended hydraulic cement.
- 3.14 Portland Cement portland cement meeting the requirements of *Standard Specifications* Section 9-01.2(1).
- 3.15 Blended Hydraulic Cement blended hydraulic cement meeting the requirements of *Standard Specifications* Section 9-01.2(4).
- 3.16 Negative Report a document provided to the agency when portland cement and/or blended hydraulic cement was not produced or shipped during a given month.

4. Significance and Use

This is accomplished by a system that evaluates quality control and specification compliance tests performed by the producers and suppliers according to their quality management plan. Products determined to meet the requirements of this standard are eligible for listing on the WSDOT Qualified Products List (QPL).

5. Laboratory and Tester Requirements

The producers/suppliers testing laboratory used to conduct specification compliance testing for the quality management program shall be AASHTO accredited by January 1, 2016. Only laboratories that are participants in the CCRL on-site inspection and proficiency sample program and are accredited from the AASHTO Accreditation Program (AAP) are recognized as approved laboratories for this program. The testing laboratory must maintain AASHTO accreditation while providing materials to WSDOT.

6. Qualification of Producers/Suppliers

- 6.1 Producers/Suppliers shall submit a written request to WSDOT for acceptance into the CAP and provide the following:
 - A copy of the producer/supplier Quality Management Plan meeting the requirements of Section 7 of OC 1.
 - A copy of the producer/supplier testing laboratory's AASHTO accreditation. One representative 10 pound sample for each type of portland cement and/or blended hydraulic cement along with the corresponding "Cement Mill Test Report" or the "Cement Certificate of Analysis". Samples shall be taken in accordance with AASHTO T 127.

- A copy of the Material Safety Data Sheet (MSDS) or Safety Data Sheet (SDS) as applicable for each sample submitted.
- Cement Mill Test Reports or the Cement Certificate of Analysis from the previous three (3) months from the production facility.
- 6.2 WSDOT will evaluate the submittal and may test the samples provided in accordance with Section 9 of QC 1. WSDOT will notify prospective producers/suppliers in writing after completion of the evaluation. All determinations of approval or rejection by WSDOT shall be final.
- 6.3 The producer/supplier shall allow WSDOT to visit and observe the quality control activities and provide samples to WSDOT upon request.

7. Producers/Suppliers Quality Management Plan

- 7.1 The quality management plan as a minimum shall identify the following:
 - Facility type
 - · Facility address.
 - Name, email address, and telephone number of the contact person responsible for the quality control of the facility.
 - List each quality control test method to be performed on each type of portland cement or blended hydraulic cement.
 - Name and address of the AAP testing laboratory performing specification compliance testing.
 - Declaration stating that if a test result indicates a lot of portland cement or blended hydraulic cement is not in compliance with the WSDOT specifications, the facility shall immediately notify WSDOT of the lot in question.
 - Description of the method and frequency of sampling, quality control testing, and specification compliance testing.
 - Type of portland cement and/or blended hydraulic cement to be provided to WSDOT.
 - A statement of compliance with Section 5.
- 7.2 A new quality management plan shall be required whenever changes occur that cause the existing quality management plan to become inaccurate or invalid.

8. Documentation Requirements

- 8.1 Each producer/supplier shall certify conformance to *Standard Specifications* for physical and chemical requirements of AASHTO M-85, AASHTO M-240, ASTM C-150 or ASTM C-595 by means of a "Cement Mill Test Report" or "Cement Certificate of Analysis".
- 8.2 A "Cement Mill Test Report" shall be provided monthly by the cement producer to WSDOT on a continuous basis for AASHTO M-85, AASHTO M-240, ASTM C-150 or ASTM C-595 cement production.

Cement mill test reports shall be in English and include the following information:

- Name of producer
- Specific type of cement in accordance with Standard Specifications Section 9-01
- Unique identification number traceable to the date of production
- · Production date

8.3 A "Cement Certificate of Analysis" shall be provided by the cement supplier to WSDOT whenever a new shipment of AASHTO M-85, AASHTO M-240, ASTM C-150 or ASTM C-595 imported cement is received for distribution.

Cement certificate of analysis shall be in English and include the following information:

- Name of supplier
- Specific type of cement in accordance with *Standard Specifications* Section 9-01
- Unique identification number traceable to each shipment
- Certification date
- 8.4 Separate sequences of Cement Mill Test Reports or Cement Certificate of Analyses shall be provided for each individual production facility and a unique lot number traceable to a production run on cement shall identify each report.
- 8.5 "Cement Mill Test Report" or "Cement Certificate of Analysis" shall show the test results and the applicable specifications of AASHTO M-85, AASHTO M-240, ASTM C-150 or ASTM C-595 for each component or property tested and shall show the test requirements specified by WSDOT.
- 8.6 When a production facility does not produce cement in a given month, or no shipments are received by a supplier, the producer/supplier shall notify the Agency with a Negative Report for each month of no production or shipment.
- 8.7 Cement Mill Test Report, Cement Certificate of Analysis and negative reports shall be emailed to the CAP program at following email address: capprogram@wsdot.wa.gov.
- 8.8 The producer/supplier shall notify WSDOT at the email address noted above of any temporary stops in production (greater than one month) or permanent stops in production.

9. Quarterly Split Sample

- 9.1 Cement producers/suppliers shall, on a quarterly basis, provide a split sample of each type of portland cement or blended hydraulic cement being produced
- 9.2 For the purpose of this standard, quarters are defined as; January through March, April through June, July through September, October through December.
- 9.3 Split samples shall be taken from production or shipment in accordance with the producer/supplier's quality management plan.
- 9.4 The production sample shall be split into two portions (approximately 10 pounds each) for each type of cement being produced. One portion shall be retained by the producer/ supplier and one portion shall be sent to WSDOT CAP.
- 9.5 The producer/supplier testing laboratory shall conduct chemical and physical testing on their portion.
- 9.6 The sample submitted to WSDOT will include the "Cement Mill Test Report" or "Cement Certificate of Analysis" for the lot number that is traceable to this production run or lot of cement. WSDOT may elect to test the sample.

9.7 Samples and accompanying documentation shall be sent to:

WSDOT State Materials Laboratory ATTN: Cement Acceptance Program 1655 S. Second Ave SW Tumwater, WA 98512-6951

9.8 The producer/supplier shall email CAP at the email address noted in Section 8.7 if no cement was produced and no sample will be submitted.

10. Comparison of Quarterly Split Sample Test Results

- 10.1 Results of the split sample testing must conform to the applicable AASHTO or ASTM specification requirements.
- 10.2 If any discrepancy is identified between the producer/suppliers and WSDOT's test results the producer/supplier shall prepare a response to WSDOT, within 30 days of being notified of discrepancy.
- 10.3 The response shall identify the cause of the discrepancy and describe any corrective action taken.

11. Revocation Of Qualification

- 11.1 A Producer/Supplier may have its qualification status revoked and be removed from the Qualified Products List if found in nonconformance with the *Standard Specifications* or this Standard Practice. Causes for removal from the QPL may include, but are not limited to, the following:
 - Failure to comply with requirements of Standard Practice QC 1.
 - Failing test results on production, shipment or project samples.
 - Failure to notify WSDOT of changes in product formulation.

Prior to removing a producer/supplier from the Qualified Products List (QPL), WSDOT will take appropriate measures to confirm the validity of the information and will confer with the producer/supplier.

12. Requalification

- 12.1 Once a product has been removed from the QPL, the producer/supplier may request reinstatement by providing the following written information to WSDOT:
 - The root cause and corrective action taken to prevent future reoccurrences of the problem that caused the removal from the QPL.
 - Updated Quality Management Plan showing compliance with QC 1.
 - Other information and test data as determined by WSDOT.

Provided there is a satisfactory resolution of the initial problem, at WSDOT's discretion the product may either be reinstated into the QPL, or the producer/supplier may be required to reapply to the QPL. All costs of the QPL process shall be borne by the producer/supplier.

WSDOT Standard Practice QC 2

Standard Practice for Asphalt Suppliers That Certify Performance Graded and Emulsified 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) and emulsified asphalts. 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:
 - M 320 Specifications for Performance-Graded Asphalt Binder
 - R 29 Standard 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
 - R 5 Selection and use of emulsified asphalts
 - T 59 Standard Method of Test for Emulsified Asphalts
- 2.2 ASTM Standards
 - D 8 Definitions of Terms Relating to Materials for Roads and Pavements
 - D 3665 Random Sampling of Construction Materials
- 2.3 WSDOT Standards and Documents

Current WSDOT Standard Specifications

Current WSDOT Construction Manual

Appropriate State Specifications

Current WSDOT Qualified Products List

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 or emulsified asphalt for the purposes of meeting the requirements of a PGAB or emulsified asphalt.
- 3.5 PGAB Performance Graded Asphalt Binder
- 3.6 Supplier A supplier shall be defined as one who produces the final PGAB or emulsified asphalt product, or who makes, blends, modifies, or alters properties of the PGAB or emulsified asphalt. This process can occur at the refinery, terminal, the HMA Plant, or in a transport vehicle.

If no modifications are made to the PGAB or emulsified asphalt after its initial production at the refinery, the refinery shall be the supplier and must provide the certification. If material is purchased with the intent to resell with or without modification, the reseller shall then be considered the supplier.

If any modifications are made to the PGAB or emulsified asphalt 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 or emulsified asphalt from different sources is made at the HMA Plant or by the supplier of HMA or emulsified asphalt, then the supplier shall provide the certification for the PGAB or emulsified asphalt.

- 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 for the material identified.
- 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
- 3.11 Emulsified asphalt-An emulsion of asphalt cement and water which contains a small amount of an emulsifying agent. Emulsified asphalt droplets may be of either the anionic or cationic type, depending upon the emulsifying agent.
- 3.12 QPL-Washington State Department of Transportation, Qualified Products List.
 - **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 the disruption of PGAB and emulsified asphalt shipments. This is accomplished by a certification system that evaluates quality control, on-site assessments, 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 isimportant to the establishment of a valid certification program.

6. Laboratory and Tester Requirements

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

- 6.1 Laboratory facilities shall adequately house and allow proper operation of all required equipment in accordance with the applicable test procedures.
- 6.2 The laboratory shall use personnel qualified in accordance with the appropriate sections of AASHTO R-18.
- 6.3 The laboratory shall use testing equipment that has been calibrated/standardized/checked to meet the requirements of each test procedure in accordance with the appropriate sections of AASHTO R-18.
- 6.4 Documentation of personnel qualifications and the equipment calibration/standardization/ check records shall be maintained.
- 6.5 The Agency at their discretion may review the laboratory facility, testing equipment, personnel performing the testing, and review all qualification and calibration and verification testing.

7. Supplier Requirements

- 7.1 The Supplier shall submit a written request to the Agency for authorization to supply PGAB or emulsified asphalts. The request shall include copies of their preliminary test reports for the proposed PGAB or emulsified asphalts with the appropriate documentation. If requested by the Agency, a sample of the PGAB or emulsified asphalt shall be provided to the Agency for testing.
 - **Note:** Suppliers currently on the Qualified Products List shall be exempt from submitting a written request for those products they are already approved. WSDOT may request preliminary test reports and a sample 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 testing.
- 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 A new Quality Control Plan shall be required whenever changes occur that cause the existing Quality Control Plan to become inaccurate or invalid.
- 7.6 The Supplier shall establish a continuing test record for each test required on each PGAB or emulsified asphalts.

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 or emulsified asphalt at the facility.
 - 8.1.4 Name the laboratory performing quality control tests on the PGAB or emulsified asphalt 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 or emulsified asphalt 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 type and grade, (3) cease shipment until the material meets specification compliance, (4) notify the Agency prior to resuming shipment.
- 8.3 The Supplier's quality control plan shall describe the method and frequency for, sampling, specification compliance testing and quality control testing.
 - 8.3.1 Specification Compliance Testing shall be performed on an adequate amount of material to ensure specification compliance. The amount of material shall be agreed upon by the supplier and the Agency and included in the Quality Control Plan.
 - **Note:** Due to the various operations and manufacturing processes, each supplier will be treated individually.
 - 8.3.2 With the exception of the 24 Hour Storage Stability test, the Supplier of Emulsified Asphalt shall provide test results for each production batch of CRS-2P showing the product meets WSDOT *Standard Specification* 9-02.1(6)A upon or prior of delivery to the project. The 24 Hour Storage Stability test results shall be provided to the State Materials Laboratory in Tumwater within 48 hours of completion of the production batch.
 - 8.3.3 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 upon request.
- 8.5 The Supplier's Quality Control Plan shall include a procedure, which must 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, acceptance sampling, or verification sampling and testing in accordance with the Agency standards.

10. Requirements For Shipping PGAB or Emulsified Asphalt 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 type and grade of material, (3) the quantity of material shipped, (4) the date of shipment, (5) a statement certifying the material meets specification requirements (6) a statement certifying that the transport vehicle was inspected before loading and was found acceptable for the material shipped, and (7) shipments of CRS-2P shall include test results per section 8.3.2.

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 or emulsified asphalt revoked by the Agency if it is found not to conform to the specifications and standards as established under this standard. This will include being removed from the *Qualified Products List* (QPL)
- 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 or emulsified asphalt 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 or emulsified asphalt as required under Section 9.2 when a test result indicates that the PGAB or emulsified asphalt is not in compliance with the Agency specifications.
- 12.3 A Supplier that has been decertified may seek reinstatement by demonstrating conformance to Agency certification criteria. Reinstatement will also include reapplication to the *Qualified Products List*.

Quality System Laboratory Review

1. Scope

This standard specifies requirements and procedures for the review of WSDOT Regional Materials Laboratory and for Private Laboratories by the Quality Systems Laboratory Review Team. The on-site laboratory review shall include the following elements:

- Review of the testing facility.
- Review of the equipment calibration/verification records.
- Review of the testing technician's training records.
- Physical inspection of the equipment used to perform tests.
- Observation of technician performing the test procedure.
- Review of test reports and calculations.

2. Referenced Documents

- 2.1 AASHTO Standards
 - R 18 Establishing and Implementing a Quality System for Construction Materials Testing Laboratories
 - R 61 Establishing Requirements for and Performing Equipment Calibrations, Standardizations, and Checks

2.2 WSDOT Standards

Materials Manual M 46-01

Construction Manual M 41-01

Standard Specifications for Road, Bridge, and Municipal Construction M 41-10

3. Terminology

- 3.1 AASHTO American Association of State Highway and Transportation Officials
- 3.2 ASTM American Society for Testing and Materials
- 3.3 Calibration A process that establishes the relationship (traceability) between the results of a measurement instrument, measurement system, or material measure and the corresponding values assigned to a reference standard (Note 1).
 - **Note 1:** The definition for calibration and the following definitions for check, standardization, traceability, uncertainty, and verification of calibration are based on the definitions in R 61.
- 3.4 Check A specific type of inspection and/or measurement performed on equipment and materials to indicate compliance or otherwise with stated criteria.

- 3.5 Standardization A process that determines (1) the correction to be applied to the result of a measuring instrument, measuring system, material measure, or reference material when its values are compared to the values realized by standards; or (2) the adjustment to be applied to a piece of equipment when its performance is compared with that of an accepted standard or process.
- 3.6 WSDOT Washington State Department of Transportation

4. Significance and Use

4.1 This standard specifies procedures for reviewing laboratories for the purpose of determining the capability of the facility and its personnel to perform the necessary acceptance testing for WSDOT.

5. Laboratory Requirements

- 5.1 Facility and Equipment
 - 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 temperature and humidity of the laboratory shall meet the requirements of all test procedures performed in the laboratory.
 - 5.1.3 The testing areas shall be clean and free of clutter.
 - 5.1.4 The laboratory shall use testing equipment that meets the requirements of each test procedure.
 - 5.1.5 Testing equipment for private laboratories and the State Materials Laboratory shall be calibrated/standardized/checked in accordance with the test procedure, appropriate sections of AASHTO R 18 and AASHTO R 61. WSDOT region and field laboratories testing equipment shall be calibrated/standardized/checked in accordance with the test procedure and Section 9-5 of the *Construction Manual* M 41-01.
 - 5.1.6 Documentation of equipment calibration/standardization/check shall be maintained and available on-site during laboratory review.
 - 5.1.7 Safety equipment will be available and maintained in proper working order.
- 5.2 Tester Training and Records
 - 5.2.1 The laboratory shall use personnel qualified in accordance with the appropriate sections of AASHTO R 18. WSDOT region and field laboratory personnel shall be qualified in accordance with Section 9-5 of the *Construction Manual M* 41-01.
 - 5.2.2 The laboratory shall maintain records of training for each tester.
 - 5.2.3 A tester's competency for performing a test procedure shall be evaluated using a checklist relating to the test procedure. The checklist shall be filed in the tester's training record.
 - *Note:* Private laboratories may use test procedure checklists from the *Materials Manual*, or may develop their own checklists similar to those found in the *Materials Manual*.
 - 5.2.4 Testers for private laboratories shall be reviewed for qualification at the frequency stated in the *Laboratory Quality Systems Manual* (LQSM).

5.3 Manuals and Records

- 5.3.1 Private laboratories shall have an up-to-date LQSM meeting the requirements of AASHTO R 18 and approved by the State Materials Engineer.
- 5.3.2 All private laboratories shall have an up-to-date copy of the LQSM on-site and available to all testers.
- 5.3.3 Each tester must have access to the most current copy of the AASHTO, ASTM, and *Materials Manual*. WSDOT testers must have access to the most current copy of the *Construction Manual* M 41-01.
- 5.3.4 If an earlier version of the *Materials Manual* or *Construction Manual* M 41-01 is required by contract, the laboratory shall maintain an unaltered version of the required manual.
- 5.3.5 A file of MSDS sheets must be maintained in the laboratory and must be available to all testers.
- 5.3.6 Test records are required to contain sufficient information to permit verification of any test report (original observations, calculations, derived data, and identification of personnel involved in the sampling and testing).
- 5.3.7 Amendments to reports must be made in the manner stated in the LQSM.
- 5.3.8 The laboratory shall define the process used to ensure testers are performing the correct testing procedure according to the clients' contractual requirements (i.e., AASHTO, ASTM, or WSDOT test procedure as required by the contract).
- 5.3.9 Test reports are required to contain the following information:
 - Name and address of the testing laboratory.
 - Name and address of the client or identification of the project.
 - Date of receipt of the test sample.
 - Date of test performance.
 - Identification of the standard test method used and notation of all know deviations from the test method.
 - Test results and specification of the material.
 - Name of tester performing the test.
 - Date report was issued.
 - Name of person accepting technical responsibility for test report.

6. Sampling

- 6.1 Test samples required for observation of test procedures shall be obtained by:
 - T 2 WSDOT FOP for AASHTO for Soils and Aggregate
 - T 168 WSDOT FOP for WAQTC for Hot Mix Asphalt
 - TM 2 WSDOT FOP for WAQTC for Concrete

7. Sample Preparation Requirements

7.1 Prior to the performance portion of the laboratory review, for the testing being performed, samples are required to be prepared as shown in Table 1.

Test Procedure	Test	Required Preparation
Aggregate Tests		
FOP for AASHTO T 335	Fracture	Material washed, graded, and ready for counting fracture.
FOP for WAQTC T 27/T 11	Sieve Analysis of Fine and Coarse Aggregates	Split or quarter proper amount of the original sample and dry to constant weight.
		Have a split of the original sample that has been washed and dried, ready for sieving.
		3. Retain all weights in order to do calculations.
FOP for AASHTO T 176	Sand Equivalent Test	Have a sample (approximately 1000 g) of #4 minus material prepared for the moisture conditioning process (do not moisten).
		Have two properly prepared tins ready for introduction into the SE tube.
FOP for AASHTO T 248	Reducing Sample	30 lbs dry material.
FOP for AASHTO T 304	Uncompacted Voids	Have sample washed and dried.
	·	2. Sample separated into individual size fractions.
Concrete Tests	ı	
FOP for AASHTO T 106	Compressive Strength	Three mortar cubes.
FOP for AASHTO T 22	Compressive Strength	Two cylinders.
FOP for AASHTO T 231	Capping Cylinder	Have capping sulfur compound heated and ready for capping.
		2. Have two cylinders available for capping (can be the cylinders for T 22).
WSDOT T 810	Density of Pavement Core	Have a drilled pavement core available.
WSDOT T 812	Length of Drilled PCC Core	May use the core from T 810.
Soils Tests		
WSDOT T 417*	Resistivity and pH	Prepare a 100 g sample of natural #8 minus material for the pH test.
		2. Prepare the soil/water slurry a minimum of 1 hour prior to test review.
		3. Prepare a sample of #8 minus material that is four times the volume of the soil box for the resistivity test.
		4. Add 10 percent by weight of water to the sample and allow it to stand a minimum of 12 hours in a waterproof container.
AASHTO T 84*	Specific Gravity and Absorption Fine Agg.	Prepare sample to step 6.1.2 of the procedure.
AASHTO T 85*	Specific Gravity and Absorption Coarse Agg.	Prepare sample to step 8.2 of the procedure.
AASHTO T 87*	Dry Preparation of Disturbed Soil and Soil Aggregate Samples for Test	500 g of soil aggregate air dried.
AASHTO T 88*	Particle Size Analysis	No preparation.

Sample Preparation Requirements *Table 1*

Test Procedure	Test	Required Preparation
AASHTO T 100*	Specific Gravity Soils	No preparation.
AASHTO T 255	Moisture Content	No preparation.
AASHTO T 265	Moisture Content	No preparation.
FOP for AASHTO T 99/T 180	Proctor	Prepare five representative samples of #4 or ¾" material at approximately 2 percent moisture already added to each sample starting at approximately 4 percent below optimum moisture of the material. Store in sealed containers.
WSDOT T 606	Maximum Density Curve	Dry and split a sample of material into coarse and fine material.
		2. Prepare fine material for Test 1.
		Prepare coarse material for either Test 2, Procedure 1 or Test 2, Procedure 2.
Hot Mix Asphalt Tests	(Have HMA samples ready on	the first day of review.)
WSDOT T 712*	Reducing Sample	An adequate amount of HMA to perform all the testing required. Heat sample and have it ready to reduce. Required to split material from sample for T 308, T 312, T 329, T 209.
FOP for AASHTO T 166*	Bulk Specific Gravity	A room temperature compacted sample must be provided for this test. A gyratory sample or a core sample will suffice.
WSDOT SOP 724*	Preparation of Aggregates	Representative aggregate from stockpiles used in JMF, dried to a constant weight.
WSDOT SOP 726*	Mixing Procedure HMA	Binder used in JMF mix design heated to mixing temperature as recommended by binder supplier (typically one quart container).
		Aggregate representative of JMF sample size based on class of HMA heated to mixing temperature as recommended by binder supplier.

^{*}WSDOT Laboratories only unless review of a private laboratory is requested by the project office.

Sample Preparation Requirements Table 1 (continued)

8. Performance of Test Procedure

- 8.1 All technicians must be current in their qualifications.
- 8.2 The laboratory review team will evaluate the technician's testing proficiency using an approved WSDOT checklist.
- 8.3 All equipment, used during the evaluation of the technician's proficiency, must be operational and have a current calibration sticker on the equipment.
- 8.4 When the test is complete, the reviewer will go over the checklist with the tester and point out any deficiencies that occurred during the performance of the test procedure.

9. Termination of Review

- 9.1 A laboratory review team member may choose to terminate the review of a procedure for the following reasons:
 - 9.1.1 Equipment is non-operational or the wrong equipment is being used.
 - 9.1.2 Tester is not qualified in the test procedure being reviewed.
 - 9.1.3 Tester makes multiple major errors in the performance of the test.
- 9.2 The review of the laboratory may be terminated by the WSDOT Quality Systems Manager for the following reasons:
 - 9.2.1 Facility is not adequate for the test procedures being reviewed.
 - 9.2.2 Two or more testers fail during the proficiency portion of the review.
 - 9.2.3 Documentation of qualification of testers or calibration of equipment is not available for review when team arrives.

10. Failure of Review

- 10.1 Rescheduling a review will require the following wait periods:
 - First Failure Minimum of one week wait to reschedule.
 - Second Failure Minimum of one month wait to reschedule.
 - Third Failure Minimum of one month wait and submittal of corrective action documentation. The documents submitted must state the concerns of the review team and the corrective action taken to solve the problem.

11. Laboratory Review Team Report

- 11.1 The Laboratory Review Team will review the facility, equipment, records, and testers compliance with the established requirements.
- 11.2 The evaluation report will be prepared and sent to the laboratory within 30 days of the completion of the review.
- 11.3 Any items that did not meet the requirements of Section 5 will be written up as "Issues."
 - 11.3.1 Issues resolved during the review shall be noted as "Issue Resolved No Response" necessary. If a "Resolved No Response Required" issue reoccurs in subsequent evaluations, the issue will be escalated to a "Response Required Issue."
 - 11.3.2 Issues that were not able to be resolved during the review will be noted as "Response Required Issue."
- 11.4 During the review, members of the team may make suggestions for improvements to the performance of the test procedure or operation of equipment. These are suggestions only and will be noted in the report as "Observations." These do not require a response.

12. Response to Report

- 12.1 Once the evaluation report has been received, the laboratory will have 90 days to respond in writing to all "Issues" labeled "Response Required."
- 12.2 The response must be a detailed explanation stating how the laboratory has resolved the issue and what measures they have taken to prevent this issue from reoccurring in the future.

13. Approval of Laboratory

- 13.1 If the laboratory review report had no issues or the issues are minor and resolved at the time of the review, the laboratory may be approved to perform acceptance, Independent Assurance, or dispute resolution testing.
- 13.2 If the laboratory review contained Response Required Issues, the laboratory may receive a conditional approval until the deficiencies are corrected or the review team may recommend that the laboratory be disapproved for all testing until the deficiencies are corrected to the satisfaction of the WSDOT Quality System Manager.

14. Suspension of Laboratory Approval

- 14.1 Laboratory approvals are subject to satisfactory results from WSDOT evaluations, including Independent Assurance evaluations. If WSDOT determines an Approved Laboratory no longer meets the approval requirements a Notification of Pending Suspension will be sent to the laboratory stating the reason for the suspension.
 - 14.1.1 The following conditions may result in suspension of a laboratory's approval status:
 - a. Failure to supply required information in a timely manner
 - b. Failure to correct deficiencies in a timely manner
 - c. Unsatisfactory performance report by the Independent Assurance Inspector
 - d. Changing the laboratory's physical location without notification to the WSDOT Quality Systems Manager
 - f. Delays in reporting the test data to WSDOT
 - g. Incomplete or inaccurate reporting
 - h. Using unqualified technicians to perform testing
 - i. Using equipment that is not calibrated, standardized or checked in accordance with AASHTO R 18
 - 14.1.2 The laboratory will be given one week to respond to the pending suspension notice with a Letter of Correction, detailing how the suspension issue has been corrected and what measures have been enacted to prevent the issue from reoccurring. The State Materials Engineer will review the Letter of Correction and determine if the corrections are adequate or if a suspension is still required and the duration of the suspension.
 - 14.1.3 A suspended laboratory must resolve all issues to the WSDOT's satisfaction and obtain reinstatement of qualification, prior to being allowed to test materials for a WSDOT project,
- 14.2 Should an approved laboratory be accused of falsifying test data or records the laboratory's approval will be suspended until the charge can be investigated. If found the approved laboratory is found to have falsified test data or records the laboratory will be disqualified from testing for a WSDOT project for a minimum of one year and be subject to further investigation and penalty under state and federal law.

Standard Practice for Fly Ash Producers/Importers/Distributors That Certify Fly Ash

1. Scope

This standard specifies requirements and procedures for a certification system that shall be applicable to all Producers/Importers/Distributors of Fly Ash.

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
 - M 295 Standard Specifications for Coal Fly Ash and Raw or Calcined Natural Pozzolan for Use in Concrete
 - R 18 Establishing and Implementing a Quality System for Construction Materials Testing Laboratories
- 2.2 ASTM Standards
 - C 618 Standard Specifications for Coal Fly Ash and Raw or Calcined Natural Pozzolan for Use in Concrete
- 2.3 Agency's Standard Specifications

3. Terminology

- 3.1 AASHTO American Association of State Highway and Transportation Officials
- 3.2 ASTM American Society of Testing and Materials
- 3.3 CCRL Cement and Concrete Reference Laboratory
- 3.4 NIST National Institute of Standards and Technology
- 3.5 Import/Distribution Facility A facility that receives finished fly ash products for distribution.
- 3.6 Production Facility A facility that has the capacity for producing fly ash.
- 3.7 Supplier A supplier stores and then delivers fly ash produced by another entity to a concrete plant or another supplier.
- 3.8 Supplier Certification Certification of fly ash provided by the supplier or importer using representative test results obtained in accordance with an agency approved QC plan and approved testing lab.

3.9 Agency – State highway agency or other agency responsible for the final acceptance of fly ash. Samples and documentation shall be sent to:

WSDOT State Materials Laboratory Attn: Cement Acceptance Program Director PO Box 47365 Olympia, WA 98504-47365

- 3.10 Specification Compliance Testing Complete testing in accordance with the specification requirements.
- 3.11 Quality Control Testing The quality control testing shall be described in the Production/Import/Distribution Facility's quality control plan. The Production/Import/Distribution Facility's quality control plan must be approved by the Agency.
- 3.12 CAP Cement Acceptance Program
- 3.13 Mill Test Report A document provided to the Agency on a monthly basis by a fly ash producer for fly ash that is actually produced at a U.S. or Canadian production facility. This document will list the actual chemical and physical test results of the product sample along with the appropriate AASHTO or ASTM specification limits.
- 3.14 Certificate of Analysis A document provided to the Agency on a per shipload basis by a fly ash importer/distributor. This document shall represent a specific shipload of imported fly ash. This document will list the actual chemical and physical test results of the product sample along with the appropriate AASHTO or ASTM specification limits.

4. Significance and Use

4.1 This standard specifies procedures for accepting fly ash. This is accomplished by a certification system that evaluates quality control and specification compliance tests performed by the Production/Import/Distribution 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/standardized/checked 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/standardization/checked records shall be maintained.
 - 5.1.5 The agency at their discretion may review the laboratory in accordance with WSDOT QC 3.
 - 5.1.6 The laboratory must participate in the NIST's CCRL proficiency sample program.

6. Production/Import/Distribution Facility Qualification

- 6.1 The Production/Import/Distribution Facility shall submit a written request for acceptance into the Cement Acceptance Program to the Agency along with a copy of the Production/Import/Distribution Facility's Quality Control Plan.
- 6.2 The Production/Import/Distribution Facility shall submit one sample with its "Mill Test Report" or "Certificate of Analysis" for the initial lot for each class of fly ash 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 either AASHTO M 295 or ASTM C 618.
- 6.4 The Production/Import/Distribution Facility shall allow the Agency to visit and observe the quality control activities and obtain samples for testing.

7. Production/Import/Distribution 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 class of fly ash.
 - 7.1.5 Name of the laboratory performing quality control tests on the fly ash if independent of the Production/Import/Distribution Facility.
 - 7.1.6 Declaration stating that if a test result indicates that a lot of fly ash 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 Class of fly ash the Production/Import/Distribution 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/Import/Distribution Facility shall document its conformance to the Agency's Standard Specifications and both physical and chemical requirements of AASHTO M 295 or ASTM C 618 by means of either a "Mill Test Report" or "Certificate of Analysis" that certifies the sample test results.
- 8.2 "Mill Test Reports" of all fly ash shall be submitted by the producer on a monthly basis to the Agency. Negative reports (i.e., reports indicating no production for the month) are required to insure that a continuous flow of documentation is maintained.
- 8.3 "Certificates of Analysis" shall be provided by the importer/distributor to the Agency whenever a new shipment of imported fly ash is received for distribution.

- 8.4 Separate sequences of "Mill Test Reports" shall be provided for each individual production facility and a unique lot number traceable to a production run shall be included in each report.
- 8.5 "Mill Test Reports" and "Certificates of Analysis" shall show the applicable test results and the applicable specifications for each component or property tested and shall show the test requirements specified by the Agency.

9. Agency Requirements

- 9.1 The Agency will review the Production/Import/Distribution Facility's quality control plan listed in Section 6 and respond to the Production Facility 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 Fly Ash to Projects

- 10.1 The Production/Import/Distribution Facility's quality control plan as approved by the Agency (see Section 9) shall be implemented.
- 10.2 Each shipment shall identify the applicable "Mill Test Report" or "Certificate of Analysis." 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" or "Certificate of Analysis."

11. Quarterly Split Sample Testing

- 11.1 Production/Import/Distribution Facilities, on a quarterly basis, shall split a production sample into two portions (10 pounds each) for each class of fly ash being produced.
- 11.2 For the purpose of this standard, quarters are defined as January through March, April through June, July through September, and October through December.
- 11.3 All fly ash test samples required by this standard shall be obtained as provided in the applicable standard specification or the Production Facility's quality control plan.
- 11.4 The Production/Import/Distribution Facility or an independent test facility meeting the requirements specified in Section 5 shall conduct chemical and physical testing on one portion.
- 11.5 The other portion, along with accompanying chemical and physical analysis, shall be submitted to the Agency. The sample will include the "Mill Test Report" or "Certificate of Analysis" for the lot number that is traceable to the production run of fly ash.
- 11.6 The Production/Import/Distribution Facility shall submit a letter in lieu of split sample(s) indicating the class(es) of fly ash (if any) for which they were accepted under this program that were not produced during the quarter.

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 The results of split sample tests must conform to the applicable AASHTO or ASTM specification requirements.

- 12.3 If any discrepancies or problems are identified between the Production/Import/Distribution Facility's test results and the Agency's test results the Production/Import/Distribution Facility shall respond to the Agency 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 properly transferred to the report?
 - c. Were the calculations leading to the test result correct?
 - d. Did the equipment used to perform the test meet specification requirements?
 - e. Did the test procedures conform to specification requirements?
 - f. Was corrective action taken to repair or replace defective equipment?
 - g. Was the technician instructed of the correct procedure?
- 12.4 The Production/Import/Distribution Facility shall prepare a response to the Agency, summarizing the results of the investigation, identifying the cause, 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/Import/Distribution 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/Import/Distribution 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 fly ash was not in compliance with the specifications as required under Section 8.1.
 - 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/Import/Distribution 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 The Import/Distribution Facility shall submit two random samples for retest.
 - 13.2.3.3 If two of three successive samples show nonconformance, the Agency will revoke certification status.
- 13.3 A Production/Import/Distribution Facility that has had its certification status revoked may seek reinstatement by demonstrating conformance to the qualification criteria shown in Section 7.

Standard Practice for Ground Granulated Blast-Furnace Slag Producers/Importers/Distributors That Certify Ground Granulated Blast-Furnace Slag

1. Scope

This standard specifies requirements and procedures for a certification system that shall be applicable to all Producers/Importers/Distributors of Ground Granulated Blast-Furnace Slag.

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
 - M 302 Standard Specifications for Ground Granulated Blast-Furnace Slag for Use in Concrete and Mortars
 - R 18 Establishing and Implementing a Quality System for Construction Materials Testing Laboratories
- 2.2 ASTM Standards
 - C 989 Standard Specifications for Ground Granulated Blast-Furnace Slag for Use in Concrete and Mortars
- 2.3 Agency's Standard Specifications

3. Terminology

- 3.1 AASHTO American Association of State Highway and Transportation Officials
- 3.2 ASTM American Society of Testing and Materials
- 3.3 CCRL Cement and Concrete Reference Laboratory
- 3.4 NIST National Institute of Standards and Technology
- 3.5 Import/Distribution Facility A facility that receives finished ground granulated blast-furnace slag for distribution.
- 3.6 Production Facility A facility that has the capacity for producing and/or grinding ground granulated blast-furnace slag.
- 3.7 Supplier A supplier stores and then delivers ground granulated blast-furnace slag produced by another entity to a concrete plant or another supplier.

- 3.8 Supplier Certification Certification of ground granulated blast-furnace slag provided by the supplier or importer using representative test results obtained in accordance with an agency approved QC plan and approved testing lab.
- 3.9 Agency State highway agency or other agency responsible for the final acceptance of ground granulated blast-furnace slag. Samples and documentation shall be sent to:

WSDOT State Materials Laboratory Attn: Cement Acceptance Program Director PO Box 47365 Olympia, WA 98504-47365

- 3.10 Specification Compliance Testing Complete testing in accordance with the specification requirements.
- 3.11 Quality Control Testing The quality control testing shall be described in the Production/Import/Distribution Facility's quality control plan. The Production/Import/Distribution Facility's quality control plan must be approved by the Agency.
- 3.12 CAP Cement Acceptance Program
- 3.13 Mill Test Report A document provided to the Agency on a monthly basis by a ground granulated blast-furnace slag producer that is actually produced at a U.S. or Canadian production facility. This document will list the actual chemical and physical test results of the product sample along with the appropriate AASHTO or ASTM specification limits.
- 3.14 Certificate of Analysis A document provided to the Agency on a per shipload basis by a ground granulated blast-furnace slag importer/distributor for imported ground granulated blast-furnace slag. This document shall represent a specific shipload of imported ground granulated blast-furnace slag. This document will list the actual chemical and physical test results of the product sample along with the appropriate AASHTO or ASTM specification limits.

4. Significance and Use

4.1 This standard specifies procedures for accepting ground granulated blast-furnace slag. This is accomplished by a certification system that evaluates quality control and specification compliance tests performed by the Production/Import/Distribution 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/standardized/checked 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/standardization/checked records shall be maintained.
- 5.1.5 The agency at their discretion may review the laboratory in accordance with WSDOT OC 3.
- 5.1.6 The laboratory must participate in the NIST's CCRL proficiency sample program.

6. Production/Import/Distribution Facility Qualification

- 6.1 The Production/Import/Distribution Facility shall submit a written request for acceptance into the Cement Acceptance Program to the Agency along with a copy of the Production/Import/Distribution Facility's Quality Control Plan.
- 6.2 The Production/Import/Distribution Facility shall submit one sample with its "Mill Test Report" or "Certificate of Analysis" for the initial lot for each grade of ground granulated blast-furnace slag 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 either AASHTO M 302 or ASTM C 989.
- 6.4 The Production/Import/Distribution Facility shall allow the Agency to visit and observe the quality control activities and obtain samples for testing.

7. Production/Import/Distribution 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 grade of ground granulated blast-furnace slag.
 - 7.1.5 Name of the laboratory performing quality control tests on the ground granulated blast-furnace slag if independent of the Production/Import/Distribution Facility.
 - 7.1.6 Declaration stating that if a test result indicates that a lot of ground granulated blast-furnace slag 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 ground granulated blast-furnace slag the Production/Import/Distribution 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/Import/Distribution Facility shall document its conformance to the Agency's Standard Specifications and both physical and chemical requirements of AASHTO M 302 or ASTM C 989 by means of either, a "Mill Test Report" or "Certificate of Analysis" that certifies the sample test results.
- 8.2 "Mill Test Reports" of all ground granulated blast-furnace slag shall be submitted by the producer on a monthly basis to the Agency. Negative reports (i.e., reports indicating no production for the month) are required to insure that a continuous flow of documentation is maintained.
- 8.3 "Certificates of Analysis" shall be provided by the importer/distributor to the Agency whenever a new shipment of imported ground granulated blast-furnace slag is received for distribution.
- 8.4 Separate sequences of "Mill Test Reports" shall be provided for each individual production facility and a unique lot number traceable to a production run shall be included in each report.
- 8.5 "Mill Test Reports" and "Certificates of Analysis" shall show the applicable test results and the applicable specifications of AASHTO M 302 or ASTM C 989 for each component or property tested and shall show the test requirements specified by the Agency.

9. Agency Requirements

- 9.1 The Agency will review the Production/Import/Distribution Facility's quality control plan listed in Section 6 and respond to the Production Facility 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 Ground Granulated Blast-Furnace Slag to Projects

- 10.1 The Production/Import/Distribution Facility's quality control plan as approved by the Agency (see Section 9) shall be implemented.
- 10.2 Each shipment shall identify the applicable "Mill Test Report" or "Certificate of Analysis." 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" or "Certificate of Analysis."

11. Quarterly Split Sample Testing

- 11.1 Production/Import/Distribution Facilities, on a quarterly basis, shall split a production sample into two portions (10 pounds each) for each type of ground granulated blast-furnace slag being produced.
- 11.2 For the purpose of this standard, quarters are defined as January through March, April through June, July through September, and October through December.
- 11.3 All ground granulated blast-furnace slag test samples required by this standard shall be obtained as provided in the applicable standard specification or the Production Facility's quality control plan.

- 11.4 The Production/Import/Distribution Facility or an independent test facility meeting the requirements specified in Section 5 shall conduct chemical and physical testing on one portion.
- 11.5 The other portion, along with accompanying chemical and physical analysis, shall be submitted to the Agency. The sample will include the "Mill Test Report" or "Certificate of Analysis" for the lot number that is traceable to the production run of ground granulated blast-furnace slag.
- 11.6 The Production/Import/Distribution Facility shall submit a letter in lieu of split sample(s) indicating the grade(s) of ground granulated blast-furnace slag (if any) for which they were accepted under this program that were not produced during the quarter.

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 The results of split sample tests must conform to the applicable AASHTO or ASTM specification requirements.
- 12.3 If any discrepancies or problems are identified between the Production/Import/Distribution Facility's test results and the Agency's test results the Production/Import/Distribution Facility shall respond to the Agency 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 properly transferred to the report?
 - c. Were the calculations leading to the test result correct?
 - d. Did the equipment used to perform the test meet specification requirements?
 - e. Did the test procedures conform to specification requirements?
 - f. Was corrective action taken to repair or replace defective equipment?
 - g. Was the technician instructed of the correct procedure?
- 12.4 The Production/Import/Distribution Facility shall prepare a response to the Agency, summarizing the results of the investigation, identifying the cause, 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/Import/Distribution 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/Import/Distribution 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 ground granulated blast-furnace slag was not in compliance with the specifications as required under Section 8.1.
- 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/Import/Distribution 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 The Import/Distribution Facility shall submit two random samples for retest.
 - 13.2.3.3 If two of three successive samples show nonconformance, the Agency will revoke certification status.
- 13.3 A Production/Import/Distribution Facility that has had its certification status revoked may seek reinstatement by demonstrating conformance to the qualification criteria shown in Section 7.

Annual Prestressed Plant Review and Approval Process

1. Scope

This standard specifies requirements and procedures for WSDOT annual approval of all manufacturing facilities producing prestressed concrete girders or precast prestressed concrete members in accordance with WSDOT *Standard Specifications* Section 6-02.3(25). Standard Practice QC 6 also applies to precast concrete units that are prestressed as identified in WSDOT *Standard Specifications* Section 6-02.3(28).

2. Referenced Documents

- 2.1 Precast/Prestressed Concrete Institute (PCI) Certification Program
- 2.2 WSDOT Qualified Products List (QPL)
- 2.3 WSDOT Standard Specification

3. Terminology

- 3.1 Plant Manufacturing facility producing prestressed concrete members with single plant location.
- 3.2. NRMCA National Ready Mix Concrete Association
- 3.3 PCI Precast/Prestressed Concrete Institute
- 3.4 RAM Request for Approval of Material (WSDOT Form 350-071) document submitted by the plant, identifying their material sources for WSDOT approval.
- 3.5 Quality Control Quality control inspection and documentation provided by the plant.
- 3.6 QPL WSDOT Qualified Products List
- 3.7 WSDOT Washington State Department of Transportation
- 3.8 WSDOT Annual Approval The approval process defined in WSDOT Standard Practice QC 6.
- 3.9 WSDOT Fabrication Inspector Quality Assurance inspector provided by the WSDOT Headquarters Materials Laboratory Materials and Fabrication Inspection Office.

4. Significance and Use

4.1 This Standard Practice specifies procedures for approving plants on an annual basis and maintaining a plant approval document reviewed annually. Submittal documents pre approve specific documentation identified in this Standard Practice; replacing the requirement for contract specific submittals. Modifications can be made to the plants submittal at any time during the annual approval period.

5. Annual Plant Approval Requirements

- 5.1 Plants shall be initially approved through the QPL or RAM approval process. Plants shall not begin fabricating prestress members prior to receiving WSDOT annual plant approval.
- 5.2 Maintain current PCI certification for the type of prestressed member being manufactured.
- 5.3 Maintain quality control staff meeting the training and certification requirements specified by the PCI plant certification program.
- 5.4 Submit and maintain an annual approval document detailed in Section 6 .Plants must complete and maintain all submittal requirements to remain in active approval status.
- 5.5 Submit annually by December 1st a document identifying the plant has reviewed their annual approval document and identifies whether the plant approval document remains unchanged, or details any anticipated revisions.
- 5.6 Successful completion of initial WSDOT plant approval detailed in Section 7 or successful maintenance of annual approval status detailed in Section 10.

6. Plant Submittal Requirement

- 6.1 Plants shall initially submit a document for annual approval that covers the submittal requirements of this section starting with Section 6.1.1. The document may be submitted by mail or submitted electronically. Submit documents to the WSDOT Fabrication and Coatings Engineer.
 - 6.1.1 Table of Contents
 - 6.1.2 PCI plant certification document. Submit plants current PCI certification letter.
 - 6.1.3 Table of Organization.
 - 6.1.4 RAM documents. Submit RAM documents for the following materials.
 - Concrete Ready Mix Batch Plant (as applicable)
 - Epoxy Coated Reinforcing Steel
 - Fabrication Facilities Manufacturing Welded Embeds and Fabricating Reinforcing Steel
 - · Prestress Strand
 - Reinforcing Steel
 - 6.1.5 Concrete mix designs. Submit mix designs on the latest revision of WSDOT Form 350-040. Mix designs are to be filled out completely. Aggregate, cement, slag, and admixtures must be from WSDOT approved sources. Mix design submittal shall include the following for each mix design;
 - Compressive strength break history. A minimum of 15 sets (2 cylinders per set)
 - Cement mill certification report
 - Chloride Ion test results
 - 6.1.6 Curing procedures. Submit the procedure that will be used to cure prestressed members. Identify whether accelerated curing will be used and detail the procedure for monitoring and documenting curing operations.

- 6.1.7 Fabrication procedures and drawings. This submittal is for standard fabrication procedures and specialized fabrication procedures. Specialized procedures may be added to the annual plant submittal as approved by the WSDOT Bridge and Structures Office, to facilitate fabrication of prestressed members. Examples of specialized procedures are as follows:
 - Tensioning and Detensioning procedures
 - · Hold down devices
 - Other procedures and drawings as determined by the Plant.
- 6.1.8 Weld procedures. Submit weld procedures for welding of embed plates or other structures as applicable to the plants manufacturing process.
- 6.1.9 Repair procedures. Submit repair procedures for anticipated repair scenarios.

 Approved repair procedures can be used during fabrication of prestressed members without further WSDOT engineering approval. Repair procedures must be detailed, including dimensional limits, and specific repair materials identified by material type, and brand name. Submittal shall include catalog cuts for repair materials.
- 6.1.10 Quality control plan. Submit quality control procedures and inspection forms. Inspection forms shall include information for the following.
 - Pre Pour Inspection Report
 - Wet Concrete Testing Report
 - Stressing Record
 - Compressive Strength Testing Report
 - Non Conformance Report
 - Post Pour Inspection Report
- 6.1.11 Ready mix batch plant NRMCA or NRMCA self-certification documentation as applicable. Not required for plants in house batch plant.
- 6.1.12 Problem resolution form. Submit a plant specific problem resolution form. Appendix "B" has an example of the Problem Resolution Form. This form is used to expedite resolution of construction Issues encountered during fabrication of prestressed concrete members.
- 6.1.13 Certificate of Compliance Document: Submit the form that will be used for the Certificate of Compliance document.
- 6.1.14 Final documentation package. Detail or outline the documents that will be provided to the WSDOT Materials and Fabrication Inspector prior to WSDOT final approval of prestressed members. Documents required in the final document package are as listed below.
 - Pre Pour Inspection Report
 - Wet Concrete Testing Report
 - Compressive Strength Testing Report
 - Post Pour Inspection Report
 - Gradation Reports
 - Cure Charts for accelerated curing

- Stressing Records
- Non Conformance Reports
- Problem Resolution documents
- Certificate of Compliance
- Certificate of Materials Origin (for projects with "Buy America" Requirement, WSDOT Form 350-109)
- · Mill certs
 - Cement
 - Epoxy Coated Reinforcing Steel
 - Fly Ash
 - Micro Silica
 - Prestress Strand
 - Reinforcing Steel
 - Slag
 - Steel components not used for Contractors convenience

7. Initial Plant Approval Process

- 7.1 Upon receiving the plants initial submittal, WSDOT will review the contents of the submittal in preparation for WSDOT's initial plant approval meeting with representatives of the plant.
- 7.2 WSDOT will be allowed 90 days for review of the plants initial submittal document. Time for review will be longer if submittals are incomplete.
- 7.3 WSDOT review responsibilities.
 - 7.3.1 WSDOT Fabrication and Coatings Engineer. Overall responsibility for annual approval and submittal review process. Coordinates all annual approval submittal activities. Reviews for acceptance all documentation with the exception of mix designs, specialized fabrication procedures, and repair procedures.
 - 7.3.2 WSDOT HQ Materials Laboratory Structural Materials Testing Engineer. Responsible for review and acceptance of mix designs.
 - 7.3.3 WSDOT Bridge Construction Office. Responsible for review and approval of repair procedures.
 - 7.3.4 WSDOT Bridge and Structures Office. Responsible for approval of specialized fabrication procedures, and review and approval of a welding procedures.
- 7.4 Review process.
 - 7.4.1 The WSDOT Fabrication and Coatings Engineer will review portions of the plant submittal and will send specific sections referenced in Section 7.3 to the respective approving authorities.
 - 7.4.2 Approving authorities will send reviewed documents back to the WSDOT Fabrication and Coatings Engineer.
 - 7.4.3 The WSDOT Fabrication and Coatings Engineer will review the status of the submittals returned from the approving authorities and incorporate the documents into the annual plant approval document.

- 7.4.4 Submittal documents will be signed or stamped "Approved", "Approved as Noted", "Not Approved", or Accepted depending on their review status.
- 7.4.5 RAM documents will be coded with acceptance codes by the WSDOT Fabrication and Coatings Engineer. RAM codes for specific items are referenced in Appendix "C" of this Standard Practice.
- 7.4.6 The WSDOT Fabrication and Coatings Engineer will make an itemized list of review comments and action items and will place them at the front of the annual submittal document returned to the plant at the time of the initial plant approval meeting. If time allows, the Fabrication and Coatings Engineer will work directly with representatives from the plant to address review comments prior to the initial plant approval meeting.
- 7.4.7 The WSDOT Fabrication and Coatings Engineer will schedule the initial plant approval meeting and will send the plant a letter and email notifying them of the date and time WSDOT will be at the plant for the initial plant approval meeting.
- 7.4.8 WSDOT will perform a formal audit of the plants facility, and operating and quality control procedures prior to the initial plant approval meeting. WSDOT will contact the plant and inform them of the date and time a WSDOT inspector will be at the plant for an inspection audit. The audit will follow the outline detailed in Appendix A.
- 7.4.9 WSDOT will provide the plant with an electronic version of the reviewed annual approval document within 15 days following completion of the initial plant approval meeting.

8. Initial Plant Approval Meeting

8.1 Scheduling

8.1.1 An initial plant approval meeting will be scheduled after WSDOT has completed its review of the plants initial submittal. The meeting will be held at the plants physical location. WSDOT will notify the plant of the date and time the meeting will be held.

8.2 Attendees

- 8.2.1 WSDOT attendees will include at a minimum, the WSDOT Fabrication and Coatings Engineer and a supervising inspector from the Materials and Fabrication Inspection Office.
- 8.2.2 Attendees from the plant shall include at a minimum the plant manager, production manager, and quality control manager, or their respective representatives.

8.3 Meeting Agenda

- 8.3.1 The meeting agenda will focus on comments from WSDOT's review of the plants annual approval document submittal, and WSDOT's plant inspection audit completed prior to the meeting.
- 8.3.2 WSDOT will inform the plant of their approval status upon completion of the initial approval meeting. Any deficiencies that would prevent approval will be identified and discussed during the meeting.

9. Initial Plant Approval Status Notification

9.1 Within 30 days following the initial plant approval meeting, the WSDOT Fabrication and Coatings Engineer will send the plant a letter informing the plant of their approval status and the period of effectiveness. Any deficiencies identified during the annual plant review and audit that would prevent annual approval will be identified in the letter.

10. Maintenance of Plant Approval Status After Initial Approval

- 10.1 Annual approval documents will remain in affect indefinitely as long as the document is maintained each calendar year.
- 10.2 The WSDOT Fabrication and Coatings Engineer will send each plant a letter in October requesting a document identifying the plant has reviewed their annual approval document and identifies whether the plant approval documents remain unchanged, or details any anticipated revisions. The letter will also detail any changes to WSDOT's program that would affect the annual approval document. The document and any revisions ready for submittal shall be sent electronically to the WSDOT Fabrication and Coatings Engineer no later than December 1st.
- 10.3 Revisions to mix designs, repair procedures, and specialized fabrication procedures will be processed through the QPL or reviewed during WSDOT project specific work activities. WSDOT will provide plants with a cost estimate for review of submittals being submitted through the QPL process. Submittals reviewed during WSDOT project specific work activities will be added to the annual approval document. Costs associated with approval through project specific work activities will be based on the same process for approvals through the RAM process.
- 10.4 Revisions submitted by the plant will be reviewed as detailed in Section 7.4.1 through 7.4.5.
- 10.5 The WSDOT Fabrication and Coatings Engineer will review annual plant approval documents in December. Review comments will be provided to plants for their action by January 15th. Upon resolution of review comments, the WSDOT Fabrication and Coatings Engineer will document revisions to the annual approval document and will maintain revision control by adding "Approved", "Approved as Noted", "Not Approved", or "Accepted" revisions to the document and providing plants with an electronic version of the plants complete approval document. WSDOT will provide a revision control document at the front of the annual approval document, which details the changes from the previous version.
- 10.6 Onsite inspection audits will be performed by WSDOT when the plant starts its first project each calendar year. Audits will not be performed by WSDOT until there is work taking place. If a calendar year passes without an active project, WSDOT will perform an inspection audit when WSDOT project specific work starts. The audit will follow the outline detailed in Appendix A.

11. Annual Maintenance Approval Status Notification

11.1 Upon successful completion of WSDOT's annual plant approval document review and onsite plant inspection audit as applicable, the WSDOT Fabrication and Coatings Engineer will send the plant a letter informing the plant of their approval status and the period of effectiveness. Any deficiencies identified during the annual plant review that would prevent annual approval will be identified in the letter.

Appendix A Precast / Prestress Plant Inspection Audit

Plant:		Date:			
Phone Number:	Conta	ct Person:			
Plant Reviewed by:					
Review Results: Acceptable Unac	cceptable				
Materials					
Concrete Cylinders					
Is cylinder fabrication and testing in acc	cordance with WS	DOT test methods?		Yes	No
Does cylinder storage comply with specifications?				Yes	No
Is cylinder capping acceptable?				Yes	No
Method of capping:	Sulphur	Rubber caps	Other _		
What types of molds are used?	Paper	Plastic	Steel	Secure	
Is cylinder testing machine calibrated?				Yes	No
Comments:					
Cement					
Is cement from an approved source?				Yes	No
Are cement certifications available?				Yes	No
Is cement storage acceptable?				Yes	No
Comments:					
Aggregate					
Has aggregate source been approved by	WSDOT?			Yes	No
Does plant use WSDOT grading?				Yes	No
Is aggregate sampled and tested prior to	use?			Yes	No
Is aggregate storage acceptable?				Yes	No
Comments:					

Reinforcing Steel		
Are mill test certificates available?	Yes	No
Is fabrication acceptable?	Yes	No
Is storage acceptable?	Yes	No
Comments:		
Forms		
Are forms clean, straight and in good condition?	Yes	No
Are forms checked for dimensions prior to use?	Yes	No
Comments:		
Batch Plant		
Does batch plant meet the certification requirements of the WSDOT Std. Spec.	Yes	No
Date of scale calibration:		
Comments:		
Fabrication		
Set up		
Is reinforcing steel placed per contract	Yes	No
Is steel tied according to specifications and held in place during concrete placement?	Yes	No
Is the plant aware tack welding is not permitted?	Yes	No
Is there a Plant QC hold point for inspection prior to setting forms?	Yes	No
Comments:		
Concrete Placement		
Is concrete delivered in a timely manner?	Yes	No
Is plant using approved concrete mix design?	Yes	No
Is required concrete testing being done?	Yes	No
Is there adequate equipment for concrete placement in forms?	Yes	No
Is concrete placed per specifications?	Yes	No
Comments:		

Curing			
Is temperature measuring equipment accepta	able?	Yes	No
Is product protected during curing?		Yes	No
Are test cylinders cured under same condition	ons as product?		
		Yes	No
What type of curing system is used?	Radiant Hot air		
	Convection		_
	Conducted Steam		
	Otner		
Comments:			
Stripping Procedures			
Are concrete cylinders for verification of strip	pping strength representative of the product?	Yes	No
Is required stripping strength being verified	with cylinder breaks prior to stripping?	Yes	No
Comments:			
Inspection			
Does plant inspection staff have a good und	erstanding of their job responsibilities?	Yes	No
Does plant inspection staff have adequate Tr	raining?	Yes	No
Is plant inspection staff familiar with the W	SDOT Annual Approval		
process and procedures?		Yes	No
Are approved shop drawings, plans, and cal	culations available?	Yes	No
Are quality control procedures being follow	ed?	Yes	No
Are quality control reports being filled out F	Properly?	Yes	No
Has the plant quality control department ver and finish are acceptable?	rified product repairs, workmanship,	Yes	No
Comments:			
Product Handling and Storage			
Are products handled and stored properly?		Yes	No
Comments:			

Overall Review Comments

Review Attendees

Name	Job Description	Phone/Email

Appendix B Manufacturers Name Problem Resolution Request Submitted To: WSDOT Bridge Construction Engineer Fax: 360-705-6809/Email: Fax: ____/Email: ____ WSDOT Contract No: Project Name: Company Name: Submitted By (Contact Person): Telephone No: Fax No: Email: Priority: High Medium Low Request Response Time: Description of Problem: **Proposed Resolution:**

Signature

WSDOT Materials Manual M 46-01.27

Name

April 2017

WSDOT Fabrication Inspector's Name and Signature:

Appendix C RAM Acceptance Codes

Products	RAM Code
Concrete Ready Mix Batch Plant	8
Epoxy Coated Reinforcing Steel	2, 5, 6
Fabrication Facilities (Steel Embeds & Fabricating Rebar)	8
Prestress Strand	1, 2, 6
Reinforcing Steel	2, 6

Acceptance Action Codes

- 1) Acceptance based upon 'Satisfactory' Test Report for samples of materials to be incorporated into the project.
- 2) Mfg. Cert. of Compliance for 'Acceptance' prior to use of material.
- 3) Catalog Cuts for 'Acceptance' prior to use of material.
- 4) Not Listed (No relevance to annual submittal process)
- 5) Only Materials Tagged 'Approved for Shipment'
- 6) Submit Certificate of Materials Origin to Project Engineer Office.(Only for projects with "Buy America" requirement.
- 7) Not Listed (No relevance to annual submittal process)
- 8) Source Approved
- 9) Approval Withheld; submit samples for preliminary evaluation
- 10) Approval Withheld
- 11) Miscellaneous Acceptance Criteria

Annual Precast Plant Review and Approval Process

1. Scope

This standard specifies requirements and procedures for WSDOT annual approval of all manufacturing facilities producing precast concrete structures in accordance with WSDOT *Standard Specification* Section 6-02.3(28).

2. Referenced Documents

- 2.1 WSDOT Qualified Products List (QPL)
- 2.2 WSDOT Standard Specification

3. Terminology

- 3.1 Plant Manufacturing facility producing precast concrete structures with single plant location.
- 3.2 NRMCA National Ready Mix Concrete Association
- 3.3 ICBO International Congress Building Officials
- 3.4 ICC-ES International Code Council Evaluation Services
- 3.5 PCI Precast/Prestressed Concrete Institute
- 3.6 NPCA National Precast Concrete Association
- 3.7 RAM Request for Approval of Material (WSDOT Form 350-071) submitted by the plant, identifying their material sources for WSDOT approval.
- 3.8 Quality Control Quality control inspection and documentation provided by the plant.
- 3.9 QPL WSDOT Qualified Products List
- 3.10 WSDOT Washington State Department of Transportation
- 3.11 WSDOT Annual Approval The certification process defined in WSDOT Standard Practice OC 7.
- 3.12 WSDOT Fabrication Inspector Quality Assurance inspector provided by the WSDOT Headquarters Materials Laboratory Materials and Fabrication Inspection Office.

4. Significance and Use

4.1 This Standard Practice specifies procedures for approving plants on an annual basis and maintaining a plant approval document reviewed annually. Submittal documents pre approve specific documentation identified in this Standard Practice; replacing the requirement for contract specific submittals. Modifications can be made to the plants submittal at any time during the annual approval period.

5. Annual Plant Approval Requirements

- 5.1 Plants shall be initially approved through the QPL or RAM approval process. Plants shall not begin fabricating precast members prior to receiving WSDOT annual plant approval.
- 5.2 Maintain current PCI, or NPCA certification, or maintain current status as a recognized fabricator by ICBO or ICC-ES.
- 5.3 Maintain quality control staff meeting the training and certification requirements specified by the certifying organization.
- 5.4 Submit and maintain an annual approval document detailed in Section 6. Plants must complete and maintain all submittal requirements to remain in active approval status.
- 5.5 Submit annually by December 1st a document identifying the plant has reviewed their annual approval document and identifies whether the plant approval document remains unchanged, or details any anticipated revisions.
- 5.6 Successful completion of initial WSDOT plant approval detailed in Section 7 or successful maintenance of annual approval status detailed in Section 10.

6. Plant Submittal Requirement

- 6.1 Plants shall initially submit a document for annual approval that covers the submittal requirements of this section starting with section 6.1.1. The document may be submitted by mail or submitted electronically. Submit documents to the WSDOT Fabrication and Coatings Engineer.
 - 6.1.1 Table of Contents
 - 6.1.2 PCI, NPCA, ICBO, or ICC-ES plant certification document. Submit plants current certification letter.
 - 6.1.3 Table of Organization.
 - 6.1.4 RAM documents. Submit RAM documents for the following materials.
 - Concrete Ready Mix Batch Plant (as applicable)
 - Epoxy Coated Reinforcing Steel
 - Fabrication Facilities Manufacturing Welded Embeds and Fabricating Reinforcing Steel
 - Reinforcing Steel
 - 6.1.5 Concrete mix designs. Submit mix designs on the latest revision of WSDOT Form 350-040. Mix designs are to be filled out completely. Aggregate, cement, slag, and admixtures must be from WSDOT approved sources.

Mix design submittal shall include the following for each mix design;

- Compressive strength break history. A minimum of 15 sets (2 cylinders per set)
- Cement mill certification report
- Chloride Ion test results
- Self-compacting concrete test data
- 6.1.6 Curing procedures. Submit the procedure that will be used to cure precast members. Identify whether accelerated curing will be used and detail the procedure for monitoring and documenting curing operations.

- 6.1.7 Fabrication procedures and drawings. This submittal is for standard fabrication procedures and specialized fabrication procedures. Specialized procedures may be added to the annual plant submittal as approved by the WSDOT Bridge and Structures Office, to facilitate fabrication of precast structures. This is the section to add lower stripping strength approvals per Standard Specification section 6-02.3(28)B.
- 6.1.8 Weld procedures. Submit weld procedures for welding of embed plates or other structures as applicable to the plants manufacturing process.
- 6.1.9 Repair procedures. Submit repair procedures for anticipated repair scenarios. Approved repair procedures can be used during fabrication of precast members without further WSDOT engineering approval. Repair procedures must be detailed, including dimensional limits, and specific repair materials identified by material type, and brand name. Submittal shall include catalog cuts for repair materials.
- 6.1.10 Quality control plan. Submit quality control procedures and inspection forms. Inspection forms shall include information for the following.
 - Pre Pour Inspection Report
 - Wet Concrete Testing Report
 - Compressive Strength Testing Report
 - Non Conformance Report
 - Post Pour Inspection Report
- 6.1.11 Ready mix batch plant NRMCA certification documentation as applicable. Not required for plants in house batch plant.
- 6.1.12 Problem resolution form. Submit a plant specific problem resolution form. Appendix "B" has an example of the Problem Resolution Form. This form is used to expedite resolution of construction Issues encountered during fabrication of precast concrete structures.
- 6.1.13 Certificate of Compliance Document: Submit the form that will be used for the Certificate of Compliance document.
- 6.1.14 Final documentation package. Detail or outline the documents that will be provided to the WSDOT Materials and Fabrication Inspector prior to WSDOT final approval of precast members. Documents required in the final document package are as listed below.
 - Pre Pour Inspection Report
 - Wet Concrete Testing Report
 - Compressive Strength Testing Report
 - Post Pour Inspection Report
 - Gradation Reports
 - Cure Charts for accelerated curing
 - Non Conformance Reports
 - Problem Resolution documents
 - Certificate of Compliance

- Certificate of Materials Origin (for projects with "Buy America" Requirement, WSDOT Form 350-109)
- Mill certs
 - Cement
 - Epoxy Coated Reinforcing Steel
 - Fly Ash
 - Micro Silica
 - Reinforcing Steel
 - Slag
 - Steel components not used for Contractors convenience

7. Initial Plant Approval Process

- 7.1 Upon receiving the plants initial submittal, WSDOT will review the contents of the submittal in preparation for WSDOT's initial plant approval meeting with representatives of the plant.
- 7.2 WSDOT will be allowed 90 days for review of the plants initial submittal document. Time for review will be longer if submittals are incomplete.
- 7.3 WSDOT review responsibilities.
 - 7.3.1 WSDOT Fabrication and Coatings Engineer. Overall responsibility for annual approval and submittal review process. Coordinates all annual approval submittal activities. Reviews for acceptance all documentation with the exception of mix designs, specialized fabrication procedures, and repair procedures.
 - 7.3.2 WSDOT HQ Materials Laboratory Structural Materials Testing Engineer. Responsible for review and acceptance of mix designs.
 - 7.3.3 WSDOT Bridge Construction Office. Responsible for review and approval of repair procedures.
 - 7.3.4 WSDOT Bridge and Structures Office. Responsible for approval of specialized fabrication procedures, and review and approval of a welding procedures.
- 7.4 Review process.
 - 7.4.1 The WSDOT Fabrication and Coatings Engineer will review portions of the plant submittal and will send specific sections referenced in Section 7.3 to the respective approving authorities.
 - 7.4.2 Approving authorities will send reviewed documents back to the WSDOT Fabrication and Coatings Engineer.
 - 7.4.3 The WSDOT Fabrication and Coatings Engineer will review the status of the submittals returned from the approving authorities and incorporate the documents into the annual plant approval document.
 - 7.4.4 Submittal documents will be signed or stamped "Approved", "Approved as Noted", "Not Approved", or "Accepted" depending on their review status.

- 7.4.5 RAM documents will be coded with acceptance codes by the WSDOT Fabrication and Coatings Engineer. RAM codes for specific items are referenced in Appendix "C" of this Standard Practice.
- 7.4.6 The WSDOT Fabrication and Coatings Engineer will make an itemized list of review comments and action items and will place them at the front of the annual submittal document returned to the plant at the time of the initial plant approval meeting. If time allows, the Fabrication and Coatings Engineer will work directly with representatives from the plant to address review comments prior to the initial plant approval meeting.
- 7.4.7 The WSDOT Fabrication and Coatings Engineer will schedule the initial plant approval meeting and will send the plant a letter and email notifying them of the date and time WSDOT will be at the plant for the initial plant approval meeting.
- 7.4.8 WSDOT will perform a formal audit of the plants facility, and operating and quality control procedures prior to the initial plant approval meeting. WSDOT will contact the plant and inform them of the date and time a WSDOT inspector will be at the plant for an inspection audit. The audit will follow the outline detailed in Appendix "A".
- 7.4.9 WSDOT will provide the plant with an electronic version of the reviewed annual approval document within 15 days following completion of the initial plant approval meeting.

8. Initial Plant Approval Meeting

8.1 Scheduling

8.1.1 An initial plant approval meeting will be scheduled after WSDOT has completed its review of the plants initial submittal. The meeting will be held at the plants physical location. WSDOT will notify the plant of the date and time the meeting will be held.

8.2 Attendees

- 8.2.1 WSDOT attendees will include at a minimum, the WSDOT Fabrication and Coatings Engineer and a supervising inspector from the Materials and Fabrication Inspection Office.
- 8.2.2 Attendees from the plant shall include at a minimum the plant manager, production manager, and quality control manager, or their respective representatives.

8.3 Meeting Agenda

- 8.3.1 The meeting agenda will focus on comments from WSDOT's review of the plants annual approval document submittal, and WSDOT's plant inspection audit completed prior to the meeting.
- 8.3.2 WSDOT will inform the plant of their approval status upon completion of the initial approval meeting. Any deficiencies that would prevent approval will be identified and discussed during the meeting.

9. Initial Plant Approval Status Notification

9.1 Within 30 days following the initial plant approval meeting, the WSDOT Fabrication and Coatings Engineer will send the plant a letter informing the plant of their approval status and the period of effectiveness. Any deficiencies identified during the annual plant review audit that would prevent annual approval will be identified in the letter.

10. Maintenance of Plant Approval Status After Initial Approval

- 10.1 Annual approval documents will remain in affect indefinitely as long as the document is maintained each calendar year.
- 10.2 The WSDOT Fabrication and Coatings Engineer will send each plant a letter in October requesting a document identifying the plant has reviewed their annual approval document and identifies whether the plant approval documents remain unchanged, or details any anticipated revisions. The letter will also detail any changes to WSDOT's program that would affect the annual approval document. The document and any revisions ready for submittal shall be sent electronically to the WSDOT Fabrication and Coatings Engineer no later than December 1st.
- 10.3 Revisions to mix designs, repair procedures, and specialized fabrication procedures will be processed through the QPL or reviewed during WSDOT project specific work activities. WSDOT will provide plants with a cost estimate for review of submittals being submitted through the QPL process. Submittals reviewed during WSDOT project specific work activities will be added to the annual approval document. Costs associated with approval through project specific work activities will be based on the same process for approvals through the RAM process.
- 10.4 Revisions submitted by the plant will be reviewed as detailed in Section 7.4.1 through 7.4.5.
- 10.5 The WSDOT Fabrication and Coatings Engineer will review annual plant approval documents in December. Review comments will be provided to plants for their action by January 15th. Upon resolution of review comments, the WSDOT Fabrication and Coatings Engineer will document revisions to the annual approval document and will maintain revision control by adding "Approved", "Approved as Noted", "Not approved", or "Accepted" revisions to the document and providing plants with an electronic version of the plants complete approval document. WSDOT will provide a revision control document at the front of the annual approval document, which details the changes from the previous version.
- 10.6 Onsite inspection audits will be performed by WSDOT when the plant starts its first project each calendar year. Audits will not be performed by WSDOT until there is work taking place. If a calendar year passes without an active project, WSDOT will perform an inspection audit when WSDOT project specific work starts. The audit will follow the outline detailed in Appendix A.

11. Annual Maintenance Approval Status Notification

11.1 Upon successful completion of WSDOT's annual plant approval document review and onsite plant inspection as applicable, the WSDOT Fabrication and Coatings Engineer will send the plant a letter informing the plant of their approval status and the period of effectiveness. Any deficiencies identified during the annual plant review that would prevent annual approval will be identified in the letter.

Appendix A Precast / Prestress Plant Inspection Audit

Plant:		Date:			
Phone Number:	Conta	ct Person:			
Plant Reviewed by:					
Review Results: Acceptable Unacce	eptable				
Materials					
Concrete Cylinders					
Is cylinder fabrication and testing in accor	dance with WSI	OOT test methods?		Yes	No
Does cylinder storage comply with specific	cations?			Yes	No
Is cylinder capping acceptable?				Yes	No
Method of capping:	Sulphur	Rubber caps	Other _		
What types of molds are used?	Paper	Plastic	Steel	S	ecurer
Is cylinder testing machine calibrated?				Yes	No
Comments:					
Cement					
Is cement from an approved source?				Yes	No
Are cement certifications available?				Yes	No
Is cement storage acceptable?				Yes	No
Comments:					
Aggregate					
Has aggregate source been approved by W	SDOT?			Yes	No
Does plant use WSDOT grading?				Yes	No
Is aggregate sampled and tested prior to us	se?			Yes	No
Is aggregate storage acceptable?				Yes	No
Comments:					
Reinforcing Steel					
Are mill test certificates available?				Yes	No
Is fabrication acceptable?				Yes	No

Is storage acceptable?	Yes	No
Comments:		
Forms		
Are forms clean, straight and in good condition?	Yes	No
Are forms checked for dimensions prior to use?	Yes	No
Comments:		
Batch Plant		
Does batch plant meet the certification requirements of the WSDOT Std. Spec.	Yes	No
Date of scale calibration:		
Comments:		
Fabrication		
Set Up		
Is reinforcing steel placed per contract	Yes	No
Is steel tied according to specifications and held in place during concrete placement?	Yes	No
Is the plant aware tack welding is not permitted?	Yes	No
Is there a Plant QC hold point for inspection prior to setting forms?	Yes	No
Comments:		
Concrete Placement		
Is concrete delivered in a timely manner?	Yes	No
Is plant using approved concrete mix design?	Yes	No
Is required concrete testing being done?	Yes	No
Is there adequate equipment for concrete placement in forms?	Yes	No
Is concrete placed per specifications?	Yes	No
Comments:		
Curing		
Is temperature measuring equipment acceptable?	Yes	No
Is product protected during curing?	Yes	No
Are test cylinders cured under same conditions as product?	Yes	No
What type of curing system is used? Radiant Hot air		

	Convection		
	Conducted Steam		
Comments:			
Stripping Procedures			
Are concrete cylinders for verification of stripping	strength representative of the product?	Yes	No
Is required stripping strength being verified with	cylinder breaks prior to stripping?	Yes	No
Comments:			
Inspection			
Does plant inspection staff have a good understa	nding of their job responsibilities?	Yes	No
Does plant inspection staff have adequate Training	ng?	Yes	No
Is plant inspection staff familiar with the WSDO and procedures?	T Annual Approval process	Yes	No
Are approved shop drawings, plans, and calculat	ions available?	Yes	No
Are quality control procedures being followed?		Yes	No
Are quality control reports being filled out Prope	erly?	Yes	No
Has the plant quality control department verified and finish are acceptable?	product repairs, workmanship,	Yes	No
Comments:			
Product Handling and Storage			
Are products handled and stored properly?		Yes	No
Comments:			

Overall Review Comments

Review Attendees

Name	Job Description	Phone/Email

Appendix B Manufacturers Name Problem Resolution Request			
Date:			
Submitted To:			
WSDOT Bridge Construction Engineer	Fax: 360-705-6809/Email:		
Contractor			
	_/Email:		
WSDOT Contract No:			
Submitted By (Contact Person):			
Telephone No:	Fax No:		
Email:			
Priority: High Medium Low Request Response Time:			
Description of Problem:			
Proposed Resolution:			
WSDOT Fabrication Inspector's Name as	nd Signature: Signature		

Appendix C AM Acceptance Codes

Products	RAM Code
Concrete Ready Mix Batch Plant	8
Epoxy Coated Reinforcing Steel	2, 5, 6
Fabrication Facilities (Steel Embeds & Fabricating Rebar)	8
Reinforcing Steel	2, 6

Acceptance Action Codes

- 1) Acceptance based upon 'Satisfactory' Test Report for samples of materials to be incorporated into the project.
- 2) Mfg. Cert. of Compliance for 'Acceptance' prior to use of material.
- 3) Catalog Cuts for 'Acceptance' prior to use of material.
- 4) Not Listed (No relevance to annual submittal process)
- 5) Only Materials Tagged 'Approved for Shipment'
- 6) Submit Certificate of Materials Origin to Project Engineer Office. (Only for projects with "Buy America" requirement.
- 7) Not Listed (No relevance to annual submittal process)
- 8) Source Approved
- 9) Approval Withheld; submit samples for preliminary evaluation
- 10) Approval Withheld
- 11) Miscellaneous Acceptance Criteria

WSDOT Standard Practice for HMA Mix Designs QC 8 Standard Practice for Development of Hot Mix Asphalt Mix Designs

1. Scope

- 1.1. This standard specifies requirements and procedures for approval Hot Mix Asphalt mix designs for the Qualified Products List.
- 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. WSDOT Standards
 - 2.1.1. Standard Specifications for Road, Bridge, and Municipal Construction M 41-10

3. Terminology

- 1.1. AASHTO American Association of State Highway and Transportation Officials
- 1.2. Contractor/Producer The Contractor, Producer or production facility that has the capacity for producing HMA meeting WSDOT *Standard Specifications*.
- 1.3. ASA Aggregate Source Approval
- 1.4. ASTM American Society of Testing and Materials
- 1.5. HMA Hot Mix Asphalt
- 1.6. PG Performance Graded asphalt binder
- 1.7. QPL Qualified Products List
- 1.8. State Materials Laboratory 1655 S. 2nd Avenue SW, Tumwater, WA 98512-6951
- 1.9. WSDOT Washington State Department of Transportation.

4. Significance and Use

4.1. This standard specifies procedures for designing, submitting, evaluating and approving HMA mix designs for inclusion to the QPL.

5. Mix Design Development

5.1. The Contractor/Producer or designee shall develop a HMA mix design in accordance with Section 5-04.2(1) of the *Standard Specifications*. The HMA mix design aggregate structure, asphalt binder content, anti-stripping additive, rutting susceptibility and indirect tensile strength shall be determined in accordance with WSDOT SOP 732, FOP for AASHTO T 324 and WSDOT FOP for ASTM D 6931 and meet the requirements of Sections 9-03.8(2) and 9-03.8(6) of the *Standard Specifications*.

6. Submission to the WSDOT Qualified Products List

- 6.1. Once the HMA mix design has been developed, the Contractor/Producer shall contact the QPL Engineer (www.wsdot.wa.gov/Business/MaterialsLab/QPL.htm) or 360-709-5442 to initiate the HMA mix design submittal process.
- 6.2. To initiate the mix design submittal process the Contractor/Producer shall provide the following:
 - Company contact and billing information
 - A completed copy of WSDOT Form 350-042EF. A completed QPL Application
 - ASA Report for the aggregate source(s)
 - QPL Contractor/Producer Product Information page(s) for the PG asphalt binder and the anti-stripping additive
- 6.3. The QPL Engineer will provide the following to the Contractor/Producer:
 - QPL evaluation tracking number
 - Initial letter detailing mix design evaluation
 - Cost sheet for mix design evaluation detailing submittal requirements and associated charges
- 6.4. After payment is received for the mix design evaluation the QPL Engineer shall provide:
 - Assigned delivery date of materials and documentation to State Materials Laboratory
 - Estimated date of completion
 - Final letter indicating QPL status
- 6.5. A priority queue will be established by the State Materials Laboratory for HMA mix design evaluations.
- 6.6. Preference will be given to mix designs submitted for WSDOT contracts.
 - 6.6.1. HMA mix design evaluation for WSDOT contracts shall be completed within 25 calendar days of acceptance by the State Materials Laboratory. Acceptance will be determined when all required documentation, materials and payment have been received at the State Materials Laboratory.
 - 6.6.2. HMA mix design evaluations submitted that are not for WSDOT contracts will be completed within approximately 40 calendar days of acceptance by the State Materials Laboratory.
 - 6.6.3. The State Materials Laboratory reserves the right to limit the number of HMA mix design evaluations accepted that are not for WSDOT contracts at any given time. Workload and staffing will dictate the number of HMA mix designs accepted at one time

7. Mix Design Evaluation

- 7.1. The HMA mix design submitted by the Contractor/Producer will be evaluated by the State Materials Laboratory in accordance with Section 9-03.8(2) and 9-03.8(6) of the *Standard Specifications*. All communication from the State Materials Laboratory will be to the Contractor's/Producer's contact as specified on WSDOT Form 350-042EF.
- 7.2. HMA mix designs will be placed on the QPL provided they meet the requirements of Section 9-03.8(2) and 9-03.8(6) of the *Standard Specifications*.
 - 7.2.1. Voids in Mineral Aggregate (VMA) must be within 1.0% of the minimum specification in accordance with Section 9-03.8(2) of the *Standard Specifications* for the class of HMA evaluated.
 - 7.2.2. % Gmm at N design must be within 1.5% of the specification in Section 9-03.8(2) of the *Standard Specifications* for the class of HMA evaluated.
 - 7.2.3. Voids Filled with Asphalt (VFA) in Section 9-03.8(2) will not be part of the mix design evaluation.
- 7.3. A mix design that fails to meet the requirements listed in Section 7.2, 7.2.1 and 7.2.2 will not be accepted or placed on the QPL.
- 7.4. Adjustments to mix designs will not be allowed once they have been evaluated.
- 7.5. The Contractor/Producer will be issued a QPL mix design record providing the mix design is in compliance with Section 9 of this Standard Practice.
- 7.6. The QPL listing for HMA mix designs will show the following information:
 - · Company name
 - HMA Class
 - Aggregate Source(s)
 - PG Grade
 - PG Supplier

Anti-stripping additive brand and quantity (if applicable)

8. Referencing Mix Designs From The QPL

- 8.1. Requests for reference HMA mix designs for non WSDOT projects will be completed on WSDOT Form 350-041EF and emailed to BituminousMaterials@wsdot.wa.gov.
- 8.2. Reference HMA mix design reports will be issued for new mix designs on active and awarded WSDOT contracts once accepted and placed on the QPL.
- 8.3. Reference HMA mix design reports will be issued for current mix designs on active and awarded WSDOT contracts provided the HMA production history is in compliance with *Standard Specifications* Section 5-04.3(11)D.

9. Removal From The QPL

- 9.1. HMA mix designs will be automatically removed from the QPL in accordance with *Standard Specifications* Section 5-04.2(1).
- 9.2. HMA mix designs may be removed from the QPL if found in nonconformance with the *Standard Specifications* or this Standard Practice. Causes for removal from the QPL may include, but are not limited to the following:
 - Failure to comply with requirements of Standard Practice QC 8.
 - HMA mix designs that are out of compliance in accordance with Section 5-04.3(11)F of the *Standard Specifications*.
 - Failure to notify WSDOT of changes in HMA production.
 - Removal at the request of the Contractor/Producer

10. Ignition Furnace Calibration Factor (IFCF) Samples

- 10.2. Each HMA mix design submitted for evaluation will have 12 IFCF samples produced for WSDOT as part of the QPL evaluation process.
- 10.3. The Contractor/Producer may elect to have 4 IFCF samples produced as part of the QPL evaluation process.

WSDOT Errata to FOP for AASHTO T 2

Sampling of Aggregates

WAQTC FOP for AASHTO T 2 has been adopted by WSDOT with the following changes:

Procedure - General

TABLE 1 Recommended Sample Sizes - Shall conform to the following table and note.

Nominal Max (m		Minimum Mass lb (kg)	
US No. 4	(4.75)	5	(2)
1/4	(6.3)	10	(4)
3/8	(9.5)	10	(4)
1/2	(12.5)	20	(8)
5/8	(16.0)	20	(8)
3/4	(19.0)	30	(12)
1	(25.0)	55	(25)
11/4	(31.5)	70	(30)
1½	(37.5)	80	(36)
2	(50)	90	(40)
2½	(63)	110	(50)
3	(75)	140	(60)
3½	(90)	180	(80)

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 percent of the materials.

Procedure - Specific Situations

Roadways

Method A (Berm or Windrow) – Method not recognized by WSDOT.

Method B (In-Place) – Method not recognized by WSDOT.

Sampling of Aggregates T 2

FOP for AASHTO T 2

Sampling of Aggregates

Scope

This procedure covers sampling of coarse, fine, or a combination of coarse and fine aggregates (CA and FA) in accordance with AASHTO T 2-91. Sampling from conveyor belts, transport units, roadways, and stockpiles is covered.

Apparatus

- Shovels or scoops, or both
- · Sampling tubes of acceptable dimensions
- Mechanical sampling systems: normally a permanently attached device that allows a sample container to pass perpendicularly through the entire stream of material or diverts the entire stream of material into the container by manual, hydraulic, or pneumatic operation
- · Belt template
- · Sampling containers

Procedure - General

Sampling is as important as testing. The technician shall use every precaution to obtain samples that are representative of the material. Determine the time or location for sampling in a random manner.

- 1. Wherever samples are taken, obtain multiple increments of approximately equal size.
- 2. Mix the increments thoroughly to form a field sample that meets or exceeds the minimum mass recommended in Table 1.

Table 1 Recommended Sample Sizes

Nominal Maxim	um Size* mm (in)	Minimum Mass g (lb)	
90	(3½)	175,000	(385)
75	(3)	150,000	(330)
63	(2½)	125,000	(275)
50	(2)	100,000	(220)
37.5	(1½)	75,000	(165)
25.0	(1)	50,000	(110)
19.0	(3/4)	25,000	(55)
12.5	(1/2)	15,000	(35)
9.5	(3/8)	10,000	(25)
4.75	(No. 4)	10,000	(25)
2.36	(No. 8)	10,000	(25)

^{*}One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size. Maximum size is one size larger than nominal maximum size.

Note 1: Sample size is based upon the test(s) required. As a general rule, the field sample size should be such that, when split twice will provide a testing sample of proper size. For example, the sample size may be four times that shown in Table 2 of the FOP for AASHTO T 27/T 11, if that mass is more appropriate.

Procedure - Specific Situations

Conveyor Belts

Avoid sampling at the beginning or end of the aggregate run due to the potential for segregation. Be careful when sampling in the rain. Make sure to capture fines that may stick to the belt or that the rain tends to wash away.

Method A (From the Belt)

- 1. Stop the belt.
- 2. Set the sampling template in place on the belt, avoiding intrusion by adjacent material.
- 3. Remove the material from inside the template, including all fines.
- 4. Obtain at least three approximately equal increments.
- 5. Combine the increments to form a single sample.

Method B (From the Belt Discharge)

- 1. Pass a sampling device through the full stream of the material as it runs off the end of the conveyor belt. The sampling device may be manually, semi-automatic or automatically powered.
- 2. The sampling device shall pass through the stream at least twice, once in each direction, without overfilling while maintaining a constant speed during the sampling process.
- 3. When emptying the sampling device into the container, include all fines.
- 4. Combine the increments to form a single sample.

Transport Units

- 1. Visually divide the unit into four quadrants.
- 2. Identify one sampling location in each quadrant.
- 3. Dig down and remove approximately 0.3 m (1 ft) of material to avoid surface segregation. Obtain each increment from below this level.
- 4. Combine the increments to form a single sample.

Roadways

Method A (Berm or Windrow)

- 1. Obtain sample before spreading.
- 2. Take the increments from at least three random locations along the fully-formed windrow or berm. Do not take the increments from the beginning or the end of the windrow or berm.
- 3. Obtain full cross-section samples of approximately equal size at each location. Take care to exclude the underlying material.
- 4. Combine the increments to form a single sample.

Sampling of Aggregates T 2

Note 2: Obtaining samples from berms or windrows may yield extra-large samples and may not be the preferred sampling location.

Method B (In-Place)

- 1. Obtain sample after spreading and before compaction.
- 2. Take the increments from at least three random locations.
- 3. Obtain full-depth increments of approximately equal size from each location. Take care to exclude the underlying material.
- 4. Combine the increments to form a single sample.

Stockpiles

Method A- Loader sampling

- 1. Direct the loader operator to enter the stockpile with the bucket at least 150 mm (6 in) above ground level without contaminating the stockpile.
- 2. Discard the first bucketful.
- 3. Have the loader re-enter the stockpile and obtain a full loader bucket of the material, tilt the bucket back and up.
- 4. Form a small sampling pile at the base of the stockpile by gently rolling the material out of the bucket with the bucket just high enough to permit free-flow of the material. (Repeat as necessary.)
- 5. Create a flat surface by having the loader back drag the small pile.
- 6. Visually divide the flat surface into four quadrants.
- 7. Collect an increment from each quadrant by fully inserting the shovel into the flat pile as vertically as possible, take care to exclude the underlying material, roll back the shovel and lift the material slowly out of the pile to avoid material rolling off the shovel.

Method B - Stockpile Face Sampling

- 1. Create horizontal surfaces with vertical faces in the top, middle, and bottom third of the stockpile with a shovel or loader.
- 2. Prevent continued sloughing by shoving a flat board against the vertical face. Sloughed material will be discarded to create the horizontal surface.
- 3. Obtain sample from the horizontal surface as close to the intersection as possible of the horizontal and vertical faces.
- 4. Obtain at least one increment of equal size from each of the top, middle, and bottom thirds of the pile.
- 5. Combine the increments to form a single sample.

Method C - Alternate Tube Method (Fine Aggregate)

- 1. Remove the outer layer that may have become segregated.
- 2. Using a sampling tube, obtain one increment of equal size from a minimum of five random locations on the pile.
- 3. Combine the increments to form a single sample.

Note 3: Obtaining samples at stockpiles should be avoided whenever possible due to problems involved in obtaining a representative gradation of material.

Report

- On forms approved by the agency
- Date
- Time
- Sample ID
- Location
- Quantity represented

Sampling of Aggregates T 2

Performance Exam Checklist FOP for AASHTO T 2 Sampling of Aggregates

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
Con	veyor Belts – Method A (From the Belt)		
1.	Belt stopped?		
2.	Sampling template set on belt, avoiding intrusion of adjacent material?		
3.	Sample, including all fines, scooped off?		
4.	Samples taken in at least three approximately equal increments?		
Con	veyor Belts – Method B (From the Belt Discharge)		
5.	Sampling device passed through full stream of material twice (once in each direction) as it runs off end of belt?		
Trar	nsport Units		
6.	Unit divided into four quadrants?		
7.	Increment obtained from each quadrant, 0.3 m (1 ft) below surface?		
8.	Increments combined to make up the sample?		
Roa	dways Method A (Berm or Windrow)		
9.	Sample taken prior to spreading?		
10.	Full depth of material taken?		
11.	Underlying material excluded?		
12.	Samples taken in at least three approximately equal increments?		
Roa	dways Method B (In-place)		
13.	Sample taken after spreading?		
14.	Full depth of material taken?		
15.	Underlying material excluded?		
16.	Samples taken in at least three approximately equal increments?		

Proc	edure Element	Trial 1	Trial 2
Stoc	kpile Method A– (Loader sampling)		
17.	Loader operator directed to enter the stockpile with the bucket at least 150 mm (6 in) above ground level without contaminating the stockpile?		
18.	First bucketful discarded?		
19.	The loader re-entered the stockpile and obtained a full loader bucket of the material with the bucket tilted back and up?		
20.	A small sampling pile formed at the base of the stockpile by gently rolling the material out of the bucket with the bucket just high enough to permit free-flow of the material?		
21.	A flat surface created by the loader back dragging the small pile?		
22.	Increment sampled from each quadrant by fully inserting the shovel into the flat pile as vertically as possible, care taken to exclude the underlying material?		
Stoc	kpile Method B (Stockpile Face)		
23.	Created horizontal surfaces with vertical faces?		
24.	24. At least one increment taken from each of the top, middle, and bottom thirds of the stockpile.		
Stoc	kpile Method C – Alternate Tube Method (Fine Aggregate)		
25.	Outer layer removed?		
26.	Increments taken from at least five locations with a sampling tube?		
Gen	eral		
27.	Increments mixed thoroughly to form sample?		
First	Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ture of Examiner WAQTC #:		

Comments:

FOP for WAQTC TM 2

Sampling Freshly Mixed Concrete

Scope

This method covers procedures for obtaining representative samples of fresh concrete delivered to the project site. The method includes sampling from stationary, paving and truck mixers, and from agitating and non-agitating equipment used to transport central mixed concrete.

This method also covers the removal of large aggregate particles by wet sieving.

Sampling concrete 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.

Warning—Fresh Hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.

Apparatus

- Wheelbarrow
- Cover for wheelbarrow (plastic, canvas, or burlap)
- Buckets
- Shovel
- Cleaning equipment, including scrub brush, rubber gloves, water
- Apparatus for wet sieving, including: a sieve(s), meeting the requirements of FOP for AASHTO T 27/T 11, minimum of 2 ft² (0.19 m²) of sieving area, conveniently arranged and supported so that the sieve can be shaken rapidly by hand.

Procedure

- 1. 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 from 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 0.03 m³ (1 ft³).
- 2. Dampen the surface of the receptacle just before sampling, empty any excess water.
 - **Note 1:** Sampling should normally be performed as the concrete is delivered from the mixer to the conveying vehicle used to transport the concrete to the forms; however, specifications may require other points of sampling, such as at the discharge of a concrete pump.
- 3. Use one of the following methods to obtain the sample:
 - Sampling from stationary mixers

Obtain the sample after a minimum of $\frac{1}{2}$ m³ ($\frac{1}{2}$ yd³) 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. 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

Obtain the 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, the concrete may be sampled by placing a shallow container on the subgrade and discharging the concrete across the container.

Sampling from revolving drum truck mixers or agitators

Obtain the sample after a minimum of $\frac{1}{2}$ m³ ($\frac{1}{2}$ yd³) of concrete has been discharged. Obtain samples 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. Perform sampling 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

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

Sampling from pump or conveyor placement systems

Obtain sample after a minimum of $\frac{1}{2}$ m³ ($\frac{1}{2}$ yd³) of concrete has been discharged. Obtain samples after all of the pump slurry has been eliminated. Perform sampling 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.

- 4. Transport samples to the place where fresh concrete tests are to be performed and specimens are to be molded. They shall then be combined and remixed with a shovel the minimum amount necessary to ensure uniformity. Protect the sample from direct sunlight, wind, rain, and sources of contamination.
- 5. Complete test for temperature and start tests for slump and air content within 5 minutes of obtaining the sample. Start molding specimens for strength tests within 15 minutes of obtaining the sample. Complete the test methods as expeditiously as possible.

Wet Sieving

When required due to oversize aggregate, the concrete sample shall be wet sieved, after transporting but prior to remixing, for slump testing, air content testing or molding test specimens, by the following:

- 1. Place the sieve designated by the test procedure over the 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. A horizontal back and forth motion is preferred.
- 4. Discard oversize material including all adherent mortar.
- 5. Repeat until sample of sufficient size is obtained. Mortar adhering to the wetsieving equipment shall be included with the sample.
- 6. Using a shovel, remix the sample the minimum amount necessary to ensure uniformity.

Note 2: Wet sieving is not allowed for samples being used for density determinations according to the FOP for AASHTO T 121.

Report

- On forms approved by the agency
- · Sample ID
- Date
- Time
- Location
- · Quantity represented

Performance Exam Checklist FOP for WAQTC TM 2 Sampling Freshly Mixed Concrete

Part	cipant Name Exam Date		
Reco	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Proc	edure Element	Trial 1	Trial 2
1.	Receptacle dampened and excess water removed?		
2.	Obtain a representative sample from drum mixer:		
	a. Concrete sampled after ½ m³ (½ yd³) discharged?		
	b. Receptacle passed through entire discharge stream or discharge stream completely diverted into sampling container?		
3.	Obtain a representative sample from a paving mixer:		
	a. Concrete sampled after all the concrete has been discharged?		
	b. Material obtained from at least 5 different locations in the pile?		
	c. Avoid contaminating the sample with sub-grade materials.		
4.	4. Obtain a representative sample from a pump:		
	a. Concrete sampled after $\frac{1}{2}$ m ³ ($\frac{1}{2}$ yd ³) has been discharged?		
	b. All the pump slurry is out of the lines?		
	c. Receptacle passed through entire discharge stream or discharge stream completely diverted into sampling container?		
	d. Do not lower the pump arm from the placement position.		
5.	Samples transported to place of testing?		
6.	Sample(s) combined, or remixed, or both?		
7.	Sample protected?		
8.	Minimum size of sample used for strength tests 0.03 m ³ (1 ft ³)?		
9.	Completed temperature test within 5 minutes of obtaining sample?		
10.	Start tests for slump and air within 5 minutes of obtaining sample?		
11.	Start molding cylinders within 15 minutes of obtaining sample?		
12.	Protect sample against rapid evaporation and contamination?		

Proc	edu	ure Element	Trial 1	Trial 2		
13.	We					
	a.	Required sieve size determined for test method to be performed?				
	b.	Concrete placed on sieve and doesn't overload the sieve.				
	c.	Sieve shaken until no more material passes the sieve.				
	d.	Sieving continued until required testing size obtained.				
	e.	Oversized aggregate discarded.				
	f.	Sample remixed.				
First	Att	tempt: Pass Fail Second Attempt: Pass Fail				
Signa	ature	e of Examiner WAQTC #:				
	his checklist is derived, in part, from copyrighted material printed in ACI CP-1, published by the merican Concrete Institute.					

Comments:

Performance Exam Checklist AASHTO T 19M/T 19 Bulk Density ("Unit Weight") and Voids in Aggregate (Rodding Procedure Only)

WSDOT has adopted AASHTO T 19.

Part	icipant Name Exam Date		
Pro	cedure 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.	Sample is approximately 125 to 200 percent of quantity required to fill measure?		
4.	Sample is handled correctly to avoid segregation?		
5.	Sample is dried in accordance with WSDOT FOP for AASHTO T 255?		
Rod	lding Procedure		
6.	Mass of empty unit weight measure is determined and recorded (nearest 0.1 lb)?		
7.	Measure is filled in three equal layers?		
8.	Each layer is rodded throughout it's depth 25 times with a hemispherical end of rod but rodding does not penetrating into the next layer?		
9.	Rodding is evenly distributed over the surface of the sample?		
10.	Mass of unit weight measure plus contents is determined to the nearest 0.1 lb and recorded?		
11.	All calculations performed correctly?		
12.	Bulk density reported to the nearest 1 lb/ft ³ ?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	nature of Examiner		

Comments:

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 references notes which provide explanatory material. These notes shall not be considered as requirements of the standard.

2. Referenced Documents

- 2.1 AASHTO Standards
 - R 39 Making and Curing Concrete Test Specimens in the Laboratory
 - T 23 Making and Curing Concrete Test Specimens in the Field
 - T 24 Obtaining and Testing Drilled Cores and Sawed Beams of Concrete
 - T 231 Capping Cylindrical Concrete Specimens
- 2.2 ASTM Standards
 - 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

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.

¹This FOP is based on AASHTO T 22-11 and has been modified per WSDOT standards. To view the redline modifications, contact the WSDOT Quality Systems Manager at 360-709-5412.

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 T 23, T 24, T 231, 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 machine should be capable of achieving 170 percent 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 Section 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.
 - 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.
 - 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 \pm 1.0 percent 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:

$$E = A - B$$
$$Ep = 100(A - B)/B$$

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 loadindicating mechanism of the testing machine or loads within that portion of the range below 10 percent 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 percent 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.

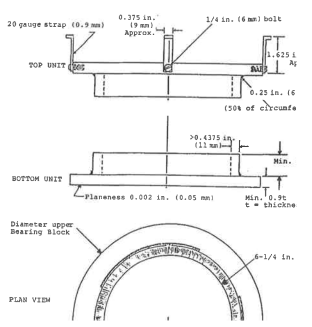
Note 3: It is desirable that the bearing faces of blocks used for compression testing of concrete have a Rockwell hardness of not less than 55 HRC.

- 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 percent 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:

Diameter of Test Specimens in (mm)	Maximum Diameter of Bearing Face in (mm)
2 (50)	4 (105)
3 (75)	5 (130)
4 (100)	6.5 (165)
6 (150)	10 (255)
8 (200)	11 (280)

- *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 \pm 5 percent of the radius of the sphere. The diameter of the sphere shall be at least 75 percent 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).
 - **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.
- 5.2.2.7 If the ball portion of the upper bearing block is a two-piece design composed of a spherical portion and a bearing plate, a mechanical means shall be provided to ensure that the spherical portion is fixed and centered on the bearing plate.



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

Schematic Sketch of a Typical Spherical Bearing Block Figure 1

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 percent of the full scale load (Note 8). The dial shall be readable within 1 percent 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 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 1.0 percent 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 percent (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, ground, or capped in accordance with T 231 to meet that tolerance or if the ends meet the requirements of A6, then neoprene caps with steel controllers may be used instead of capping. 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.
- 6.3 The height of the cylinder shall be determined to 0.01 in. The mass of the cylinder shall be determined to the nearest 0.1 lb or better.

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:

Test Age Permissible Tolerance		
12 h ± 0.25 h or 2.1%		
24 h ± 0.5 h or 2.1%		
3 days + 2 h or 2.8%		
7 days	+ 6 h or 3.6%	
28 days	+ 20 h or 3.0%	
56 days	+ 40 h or 3.0%	
90 days	+ 2 days 2.2%	

Note: The 28-day compressive break may be extended by up to 48 hours if the scheduled 28-day break falls on a Saturday, Sunday, or Holiday. The Regional Materials Engineer must authorize the time extension in writing.

- 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). 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 $(0.25 \pm 0.05 \text{ MPa/s})$ (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.
- Apply the compressive load until the load indicator shows that the load is decreasing steadily and the specimen displays a well-defined fracture pattern (Figure 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 percent 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 Figure 2. If the fracture pattern is not one of the typical patterns shown in Figure 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):

L/D: 1.75 1.50 1.25 1.00 Factor: 0.98 0.96 0.93 0.87 (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 2,000 to 6,000 psi(15 to 45 MPa). For strengths higher than 6,000 psi (45 MPa), correction factors may be larger than the values listed above x.

8.3 Calculate the average compressive strength of the set of specimens to the nearest 10 psi or 0.1MPa.

Average Compressive Strength = $\frac{(CS_1 + CS_2)}{2}$

Where:

 CS_1 = Compressive Strength of Specimen 1 CS_2 = Compressive Strength of Specimen 2

Calculate the density of the specimen to the nearest 1 lb/ft³ (10 kg/m³) as follows:

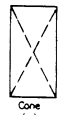
Density = $\frac{W}{V}$ where:

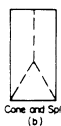
W = mass of specimen, lb (kg)

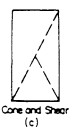
V = volume of specimen computed from the average diameter and average length or from weighing the cylinder in air and submerged, ft³ (m³)

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 Newton.
 - 9.1.5 Compressive strength calculated to the nearest 10 psi or 0.1MPa.
 - 9.1.6 Average compressive strength for the set of specimens calculated to the nearest 10 psi or 0.1 MPa.
 - 9.1.7 Type of fracture, if other than the usual cone (see Figure 2).











Sketches of Types of Fracture Figure 2

- 9.1.8 Defects in either specimen or caps.
- 9.1.9 Age of specimen.
- 9.1.10 Report 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.

Appendix A

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 Field²
 - C 39 Test Method for Compressive Strength of Cylindrical Concrete Specimens²
 - C 192/C 192M Practice for Making and Curing Concrete Test Specimens in the Laboratory²
 - C 617 Practice for Capping Cylindrical Concrete Specimens²
 - D 2000 Classification System for Rubber Products in Automotive Applications³
 - D 2240 Test Method for Rubber Property—Durometer Hardness⁴

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 $\frac{1}{2} \pm \frac{1}{16}$ in (13 ± 2 mm) thick and the diameter shall not be more than $\frac{1}{16}$ in (2 mm) smaller than the inside diameter of the retaining ring.

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

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

Shore A Durometer	Classification D 2000 Line Call-Out
50	M2BC514
60	M2BC614
70	M2BC714

The tolerance on Shore A durometer hardness is \pm 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.

²Annual Book of ASTM Standards, Volume 04.02.

³Annual Book of ASTM Standards, Volume 09.02.

⁴Annual Book of ASTM Standards, Volume 09.01.

- 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.
 - 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 percent or greater than 107 percent 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 in² (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 ½ in (13 mm) neoprene pads are shown in Figure 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 12 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 192/C 192M.

Neither end of a cylinder shall depart from perpendicularity to the axis by more than 0.5° (approximately equivalent to ½ 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 percent.

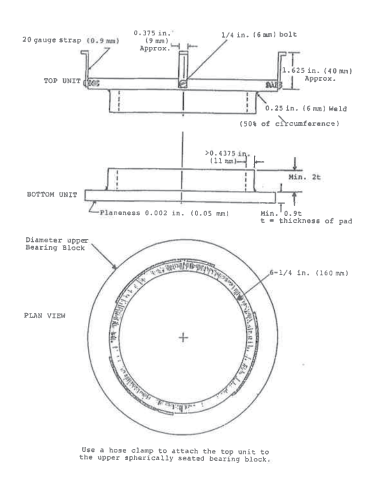
Note 2: One method of measuring the perpendicularly of ends of cylinders is to place a try square across any diameter and measure the departure of the longer blade from an element of the cylindrical surface. An alternative method is to place the end of the cylinder on a plane surface and support the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

Cylinder Compressive Strength, psi (MPa)	Shore A Durometer Hardness	Qualification Tests Required	Maximum Reuses ^A
1500 to 6000 (10 to 40)	50	None	100
2500 to 7000 (17 to 50)	60	None	100
4000 to 7000 (28 to 50)	70	None	100
7000 to 12000 (50 to 80)	70	Required	50
Greater than 12000 (80)		Not Permitted	

AMaximum number of reuses. Will be less if pads wear, crack or split (Note 6).

Requirements for Use of Polychloroprene (Neoprene) Pads *Table 1*



Example of Steel Retaining Rings for 6 by 12 in (150 by 300 mm) Cylinders (Nonmandatory)

Figure 1

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 $\frac{3}{8}$ 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 percent of the anticipated specimen strength, check to see that the axis of the cylinder is vertical within a tolerance of ½ in in 12 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 Section 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.

Evam Data

Participant Nama

Performance Exam Checklist

Compressive Strength of Cylindrical Concrete Specimens FOP for AASHTO T 22

i articipani ivanic		L'Aim 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. Specimens kept moist between removal from moist storage and testing?
- 4. Is the diameter of the cylinder reported to the nearest 0.01 inch by averaging two diameters taken at about mid-height?
- 5. Is the length of the cylinder reported to the nearest 0.01 inches?
- 6. Is the mass of the cylinder reported to the nearest 0.1 lbs or better?
- 7. Ends of cylinders checked for perpendicularity to axis?
- 8. Ends of cylinders checked for depressions greater than 0.2 inch?
- 9. Ends of cylinders checked for plane?
- 10. If ends did not meet plane, was correct method chosen to correct plane?
- 11. Are lower and upper bearing surface wiped clean?
- 12. Is the axis of the cylinder aligned with center of the spherical block?
- 13. Is the spherical block rotated prior to it contacts with the cylinder?
- 14. Is the load applied continuously and without shock?
- 15. Is the load applied at the specified rate and maintain for the latter half of the anticipated load.
- 16. Is no rate adjustment made while the cylinder is yielding?
- 17. Is the maximum load recorded?
- 18. Are cylinders tested to failure and the type of fracture recorded?
- 19. Specimens broken within the permissible time tolerance?
- 20. All calculations performed correctly?

Unbonded Caps – AASHTO 22 Appendix A

- 1. Pads examined for splits or cracks?
- 2. Cylinders centered in retaining rings?
- 3. Is cylinders checked for alignment with a small load applied?

First Attempt: Pass	Fail	Second Attempt:	Pass	Fail
Signature of Examiner				
Comments:				

WSDOT Errata to FOP for AASHTO T 23

Method of Making and Curing Concrete Test Specimens in the Field

WAQTC FOP for AASHTO T 23 has been adopted by WSDOT with the following changes:

Apparatus and Test Specimens

Initial curing facilities:

Include details below:

Cure Box – The cure box shall be a commercially manufactured cure box meeting AASHTO T 23 standards and the following requirements:

- 1. The interior shall be rustproof with a moisture-proof seal between the lid and the box.
- 2. The lid shall lock or have loops for padlocks that allow the box to be locked.
- 3. The box shall be equipped with a heating and cooling system. If the system uses a water circulating system, the box shall be equipped with a bottom drain and an overflow port. The cure box shall provide an environment that prevents loss of moisture from the specimens. The curing temperature and moist environment shall be controlled by the use of heating and cooling devices installed in the cure box.

Procedure - Initial Curing

Method 2 – Initial cure by burying in earth or by using a curing box over the cylinder – Method not recognized by WSDOT.

Include item below when required:

Field Curing

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:

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

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.

Beams – 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.

FOP for AASHTO T 23

Method of Making and Curing Concrete Test Specimens in The Field

Scope

This procedure covers the method for making, initially curing, and transporting concrete test specimens in the field in accordance with AASHTO T 23-17.

Warning—Fresh Hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.

Apparatus and Test Specimens

- Concrete cylinder molds: Conforming to AASHTO M 205 with a length equal to
 twice the diameter. Standard specimens shall be 150 mm (6 in) by 300 mm (12 in)
 cylinders. Mold diameter must be at least three times the maximum aggregate size
 unless wet sieving is conducted according to the FOP for WAQTC TM 2. Agency
 specifications may allow cylinder molds of 100 mm (4 in) by 200 mm (8 in) when the
 nominal maximum aggregate size does not exceed 25 mm (1 in).
- Beam molds: Rectangular in shape with ends and sides at right angles to each other. Must be sufficiently rigid to resist warpage. Surfaces must be smooth. Molds shall produce length no more than 1.6 mm (1/16 in) shorter than that required (greater length is allowed). Maximum variation from nominal cross section shall not exceed 3.2 mm (1/8 in). Ratio of width to depth may not exceed 1:5; the smaller dimension must be at least 3 times the maximum aggregate size. Standard beam molds shall result in specimens having width and depth of not less than 150 mm (6 in). Agency specifications may allow beam molds of 100 mm (4 in) by 100 mm (4 in) when the nominal maximum aggregate size does not exceed 38 mm (1.5 in). Specimens shall be cast and hardened with the long axes horizontal.
- Standard tamping rod: 16 mm (% in) in diameter and 400 mm (16 in) to 600 mm (24 in) long, having a hemispherical tip of the same diameter as the rod for preparing 150 mm (6 in) x 300 mm (12 in) cylinders.
- Small tamping rod: 10 mm (¾ in) diameter and 305 mm (12 in) to 600 mm (24 in) long, having a hemispherical tip of the same diameter as the rod for preparing 100 mm (4 in) x 200 mm (8 in) cylinders.
- Vibrator: At least 9000 vibrations per minute, with a diameter no more than ¼ the diameter or width of the mold and at least 75 mm (3 in) longer than the section being vibrated for use with low slump concrete.
- Scoop: a receptacle of appropriate size so that each representative increment of the concrete sample can be placed in the container without spillage.
- · Trowel or float
- Mallet: With a rubber or rawhide head having a mass of 0.57 \pm 0.23 kg (1.25 \pm 0.5 lb).
- Rigid base plates and cover plates: may be metal, glass, or plywood.

- Initial curing facilities: Temperature-controlled curing box or enclosure capable of
 maintaining the required range of 16 to 27°C (60 to 80°F) during the entire initial
 curing period (for concrete with compressive strength of 40 Mpa (6000 psi) or
 more, the temperature shall be 20 to 26°C (68 to 78°F). As an alternative, sand
 or earth for initial cylinder protection may be used provided that the required
 temperature range is maintained and the specimens are not damaged.
- Thermometer: Capable of registering both maximum and minimum temperatures during the initial cure.

Procedure – Making Specimens – General

- 1. Obtain the sample according to the FOP for WAQTC TM 2.
- 2. Wet Sieving per the FOP for WAQTC TM 2 is required for 150 mm (6 in) diameter specimens containing aggregate with a nominal maximum size greater than 50 mm (2 in); screen the sample over the 50 mm (2 in) sieve.
- 3. Remix the sample after transporting to testing location.
- 4. Begin making specimens within 15 minutes of obtaining the sample.
- 5. Set molds upright on a level, rigid base in a location free from vibration and relatively close to where they will be stored.
- 6. Fill molds in the required number of layers, attempting to slightly overfill the mold on the final layer. Add or remove concrete prior to completion of consolidation to avoid a deficiency or excess of concrete.
- 7. There are two methods of consolidating the concrete rodding and internal vibration. If the slump is greater than 25 mm (1 in), consolidation may be by rodding or vibration. When the slump is 25 mm (1 in) or less, consolidate the sample by internal vibration. Agency specifications may dictate when rodding or vibration will be used.

Procedure – Making Cylinders –Self Consolidating Concrete

- 1. Use the scoop to slightly overfill the mold. Evenly distribute the concrete in a circular motion around the inner perimeter of the mold.
- 2. Strike off the surface of the molds with tamping rod, straightedge, float, or trowel.
- 3. Immediately begin initial curing.

Procedure - Making Cylinders - Rodding

- For the standard 150 mm (6 in) by 300 mm (12 in) specimen, fill each mold in three approximately equal layers, moving the scoop or trowel around the perimeter of the mold to evenly distribute the concrete. For the 100 mm (4 in) by 200 mm (8 in) specimen, fill the mold in two layers. When filling the final layer, slightly overfill the mold.
- 2. Consolidate each layer with 25 strokes of the appropriate tamping rod, using the rounded end. Distribute strokes evenly over the cross section of the concrete. Rod the first layer throughout its depth without forcibly hitting the bottom. For subsequent layers, rod the layer throughout its depth penetrating approximately 25 mm (1 in) into the underlying layer.

- 3. After rodding each layer, tap the sides of each mold 10 to 15 times with the mallet (reusable steel molds) or lightly with the open hand (single-use light-gauge molds).
- 4. Strike off the surface of the molds with tamping rod, straightedge, float, or trowel.
- 5. Immediately begin initial curing.

Procedure – Making Cylinders – Internal Vibration

- 1. Fill the mold in two layers.
- 2. Insert the vibrator at the required number of different points for each layer (two points for 150 mm (6 in) diameter cylinders; one point for 100 mm (4 in) diameter cylinders). When vibrating the bottom layer, do not let the vibrator touch the bottom or sides of the mold. When vibrating the top layer, the vibrator shall penetrate into the underlying layer approximately 25 mm (1 in)
- 3. Remove the vibrator slowly, so that no large air pockets are left in the material.
 - **Note 1:** 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.
- 4. After vibrating each layer, tap the sides of each mold 10 to 15 times with the mallet (reusable steel molds) or lightly with the open hand (single-use light-gauge molds).
- 5. Strike off the surface of the molds with tamping rod, straightedge, float, or trowel.
- 6. Immediately begin initial curing.

Procedure - Making Flexural Beams - Rodding

- 1. Fill the mold in two approximately equal layers with the second layer slightly overfilling the mold.
- 2. Consolidate each layer with the tamping rod once for every 1300 mm² (2 in²) using the rounded end. Rod each layer throughout its depth, taking care to not forcibly strike the bottom of the mold when compacting the first layer. Rod the second layer throughout its depth, penetrating approximately 25 mm (1 in) into the lower layer.
- 3. After rodding each layer, strike the mold 10 to 15 times with the mallet and spade along the sides and end using a trowel.
- 4. Strike off the surface of the molds with tamping rod, straightedge, float, or trowel.
- 5. Immediately begin initial curing.

Procedure – Making Flexural Beams – Vibration

- 1. Fill the mold to overflowing in one layer.
- 2. Consolidate the concrete by inserting the vibrator vertically along the centerline at intervals not exceeding 150 mm (6 in). Take care to not over-vibrate, and withdraw the vibrator slowly to avoid large voids. Do not contact the bottom or sides of the mold with the vibrator.
- 3. After vibrating, strike the mold 10 to 15 times with the mallet.
- 4. Strike off the surface of the molds with tamping rod, straightedge, float, or trowel.
- 5. Immediately begin initial curing.

Procedure – Initial Curing

- When moving cylinder specimens made with single use molds support the bottom of the mold with trowel, hand, or other device.
- For initial curing of cylinders, there are two methods, use of which depends on the agency. In both methods, the curing place must be firm, within ¼ in of a level surface, and free from vibrations or other disturbances.
- Maintain initial curing temperature of 16 to 27°C (60 to 80°F) or 20 to 26°C (68 to 78°F) for concrete with strength of 40 Mpa (6000 psi) or more.
- · Prevent loss of moisture.

Method 1 - Initial cure in a temperature controlled chest-type curing box

- 1. Finish the cylinder using the tamping rod, straightedge, float, or trowel. The finished surface shall be flat with no projections or depressions greater than 3.2 mm (1/4 in).
- 2. Place the mold in the curing box. When lifting light-gauge molds be careful to avoid distortion (support the bottom, avoid squeezing the sides).
- 3. Place the lid on the mold to prevent moisture loss.
- 4. Mark the necessary identification data on the cylinder mold and lid.

Method 2 - Initial cure by burying in earth or by using a curing box over the cylinder

Note 2: This procedure may not be the preferred method of initial curing due to problems in maintaining the required range of temperature.

- 1. Move the cylinder with excess concrete to the initial curing location.
- 2. Mark the necessary identification data on the cylinder mold and lid.
- 3. Place the cylinder on level sand or earth, or on a board, and pile sand or earth around the cylinder to within 50 mm (2 in) of the top.
- 4. Finish the cylinder using the tamping rod, straightedge, float, or trowel. Use a sawing motion across the top of the mold. The finished surface shall be flat with no projections or depressions greater than 3.2 mm (1/4) in).
- 5. If required by the agency, place a cover plate on top of the cylinder and leave it in place for the duration of the curing period, or place the lid on the mold to prevent moisture loss.

Procedure – Transporting Specimens

- Initially cure the specimens for 24 to 48 hours. Transport specimens to the laboratory for final cure. Specimen identity will be noted along with the date and time the specimen was made and the maximum and minimum temperatures registered during the initial cure.
- Protect specimens from jarring, extreme changes in temperature, freezing, or moisture loss during transport.
- Secure cylinders so that the axis is vertical.
- Do not exceed 4 hours transportation time.

Final Curing

- Upon receiving cylinders at the laboratory, remove the cylinder from the mold and apply the appropriate identification.
- For all specimens (cylinders or beams), final curing must be started within 30 minutes of mold removal. Temperature shall be maintained at 23° ± 2°C (73 ± 3°F). Free moisture must be present on the surfaces of the specimens during the entire curing period. Curing may be accomplished in a moist room or water tank conforming to AASHTO M 201.
- For cylinders, during the final 3 hours prior to testing the temperature requirement may be waived, but free moisture must be maintained on specimen surfaces at all times until tested.
- Final curing of beams must include immersion in lime-saturated water for at least 20 hours before testing.

Report

- On forms approved by the agency
- Pertinent placement information for identification of project, element(s) represented, etc.
- Sample ID
- Date and time molded.
- Test ages.
- Slump, air content, and density.
- Temperature (concrete, initial cure max. and min., and ambient).
- · Method of initial curing.
- Other information as required by agency, such as: concrete supplier, truck number, invoice number, water added, etc.

Performance Exam Checklist

Making and Curing Concrete Test Specimens in the Field FOP for AASHTO T 23 (4 X 8)

Part	ticipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Molds placed on a level, rigid, horizontal surface free of vibration?		
2.	Representative sample selected?		
3.	Making of specimens begun within 15 minutes of sampling?		
Firs	t layer		
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 approximately half full?		
6.	Layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes?		
7.	Sides of the mold tapped 10-15 times after rodding?		
	a. With mallet for reusable steel molds		
	b. With the open hand for flexible light-gauge molds		
Sec	ond layer		
8.	Concrete placed in the mold, moving a scoop or trowel around the perimeter of the mold to evenly distribute the concrete as discharged?		
9.	Mold slightly overfilled on the last layer?		
10.	Layer rodded 25 times with hemispherical end of rod, uniformly distributing strokes and penetrating 25 mm (1 in) into the underlying layer?		
11	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		
12.	Concrete struck off with tamping rod, float or trowel?		
13.	Specimens covered with non-absorptive, non-reactive cap or plate?		
14.	Initial curing addressed?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
	checklist is derived, in part, from copyrighted material printed in ACI CP-1, publish rican Concrete Institute.	ned by th	ne

Comments:

Participant Name

Performance Exam Checklist FOP for AASHTO T 23 (6 X 12) Making and Curing Concrete Test Specimens in the Field

ı aı ı	Exam Dute		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Molds placed on a level, rigid, horizontal surface free of vibration?		
2.	Representative sample selected?		
3.	Making of specimens begun within 15 minutes of sampling?		
Firs	t layer		
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 approximately one third full?		
6.	Layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes?		
7.	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		
Sec	ond layer		
8.	Concrete placed in the mold, moving a scoop or trowel around the perimeter of the mold to evenly distribute the concrete as discharged?		
9.	Mold filled approximately two thirds full?		
10.	Layer rodded 25 times with hemispherical end of rod, uniformly distributing strokes and penetrating 25 mm (1 in) into the underlying layer?		
11	Sides of the mold tapped 10-15 times after rodding?		
	a. With mallet for reusable steel molds		
	b. With the open hand for flexible light-gauge molds		
Thir	d layer		
12.	Concrete placed in the mold, moving a scoop or trowel around the perimeter of the mold to evenly distribute the concrete as discharged?		
13.	Mold slightly overfilled on the last layer?		
14.	Layer rodded 25 times with hemispherical end of rod, uniformly distributing strokes and penetrating 25 mm (1 in) into the underlying layer?		

Evam Date

Procedure Element	Trial 1	Trial 2
15. Sides of the mold tapped 10-15 times after rodding?		
a. With mallet for reusable steel molds		
b. With the open hand for flexible light-gauge molds		
16. Concrete struck off with tamping rod, straightedge, float, or trowel?		
17. Specimens covered with non-absorptive, non-reactive cap or plate?		
18. Initial curing addressed?		
First Attempt: Pass Fail Second Attempt: Pass Fail		
Signature of Examiner WAQTC #:		
This checklist is derived, in part, from copyrighted material printed in ACI CP-1, publ American Concrete Institute.	ished by th	ne

Comments:

WSDOT Errata to WAQTC FOP for AASHTO T 27_T 11

Sieve Analysis of Fine and Coarse Aggregates

WAQTC FOP for AASHTO T 27_T 11 has been adopted by WSDOT with the following changes:

Procedure Method C - Method not recognized by WSDOT.

Sample Preparation

Table 1 Test Sample Sizes for Aggregate Gradation Test – *Shall conform to the following table.*

Nominal Maximum Size*in (mm)		Minimum Dry	y Mass Ib (kg)
US No. 4	(4.75)	1	(0.5)
1/4	(6.3)	2	(1)
3/8	(9.5)	2	(1)
1/2	(12.5)	5	(2)
5/8	(16.0)	5	(2)
3/4	(19.0)	7	(3)
1	(25.0)	13	(6)
11/4	(31.5)	17	(7.5)
1½	(37.5)	20	(9)
2	(50)	22	(10)
2½	(63)	27	(12)
3	(75)	33	(15)
3½	(90)	44	(20)

FOP for AASHTO T 27

Sieve Analysis of Fine and Coarse Aggregates

FOP for AASHTO T 11

Materials Finer Than 75 µm (No. 200) Sieve In Mineral Aggregate By Washing

Scope

A sieve analysis, or 'gradation,' measures distribution of aggregate particle sizes within a given sample.

Accurate determination of the amount of material smaller than 75 μ m (No. 200) cannot be made using just AASHTO T 27. If quantifying this material is required, use AASHTO T 11 in conjunction with AASHTO T 27.

This FOP covers sieve analysis in accordance with AASHTO T 27-14 and materials finer than 75 μ m (No. 200) in accordance with AASHTO T 11-05 performed in conjunction with AASHTO T 27. The procedure includes three methods: A, B, and C.

Apparatus

- Balance or scale: Capacity sufficient for the masses shown in Table 1, accurate to 0.1 percent of the sample mass or readable to 0.1 g, and meeting the requirements of AASHTO M 231
- Sieves: Meeting the requirements of ASTM E11
- Mechanical sieve shaker: Meeting the requirements of AASHTO T 27
- Suitable drying equipment (refer to FOP for AASHTO T 255)
- Containers and utensils: A pan or vessel of sufficient size to contain the test sample covered with water and permit vigorous agitation without loss of test material or water
- · Optional: mechanical washing device

Sample Sieving

- In all procedures, the test sample is shaken in nested sieves. Sieves are selected
 to furnish information required by specification. Intermediate sieves are added for
 additional information or to avoid overloading sieves, or both.
- The sieves are nested in order of increasing size from the bottom to the top, and the test sample, or a portion of the test sample, is placed on the top sieve.
- The loaded sieves are shaken in a mechanical shaker for approximately 10 minutes, refer to Annex A; Time Evaluation.
- Care must be taken so that sieves are not overloaded, refer to Annex B; Overload Determination. The test sample may be sieved in increments and the mass retained for each sieve added together from each test sample increment to avoid overloading sieves.

Sample Preparation

Obtain samples according to the FOP for AASHTO T 2 and reduce to test sample size, shown in Table 1, according to the FOP for AASHTO R 76.

Table 1	Test Sample Sizes for Aggregate Gradation Test
	1000 0000000000000000000000000000000000

Nominal Maximum Size* mm (in)		Minimum Dr	y Mass g (lb)
125	(5)	300,000	(660)
100	(4)	150,000	(330)
90	(3½)	100,000	(220)
75	(3)	60,000	(130)
63	(2½)	35,000	(77)
50	(2)	20,000	(44)
37.5	(1½)	15,000	(33)
25.0	(1)	10,000	(22)
19.0	(3/4)	5000	(11)
12.5	(½)	2000	(4)
9.5	(3/8)	1000	(2)
6.3	(1/4)	1000	(2)
4.75	(No. 4)	500	(1)

^{*}Nominal maximum size: One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps between specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size.

Test sample sizes in Table 1 are standard for aggregate sieve analysis, due to equipment restraints samples may need to be divided into several "subsamples." For example, a gradation that requires 100 kg (220 lbs) of material would not fit into a large tray shaker all at once.

Some agencies permit reduced test sample sizes if it is proven that doing so is not detrimental to the test results. Some agencies require larger test sample sizes. Check agency guidelines for required or permitted test sample sizes.

Selection of Procedure

Agencies may specify which method to perform. If a method is not specified, perform Method A.

Overview

Method A

- Determine dry mass of original test sample
- Wash over a 75mm (No. 200) sieve
- Determine dry mass of washed test sample
- Sieve washed test sample
- · Calculate and report percent retained and passing each sieve

Method B

- Determine dry mass of original test sample
- Wash over a 75mm (No. 200) sieve
- Determine dry mass of washed test sample
- Sieve test sample through coarse sieves, 4.75 mm (No. 4) sieves and larger
- Determine dry mass of fine material, minus 4.75 mm (No. 4)
- Reduce fine material
- Determine mass of reduced portion
- Sieve reduced portion
- Calculate and report percent retained and passing each sieve

Method C

- Determine dry mass of original test sample
- Sieve test sample through coarse sieves, 4.75 mm (No. 4) sieves and larger
- Determine mass of fine material, minus 4.75 mm (No. 4)
- · Reduce fine material
- Determine mass of reduced portion
- Wash reduced portion over a 75mm (No. 200) sieve
- Determine dry mass of washed reduced portion
- Sieve washed reduced portion
- Calculate and report percent retained and passing each sieve

Procedure Method A

- 1. Dry the test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the total dry mass of the sample to the nearest 0.1 percent or 0.1 g. Designate this mass as *M*.
 - When the specification does not require the amount of material finer than 75 μ m (No. 200) be determined by washing, skip to Step 11.
- 2. Nest a sieve, such as a 2.0 mm (No. 10), above the 75 μm (No. 200) sieve.
- 3. Place the test sample in a container and cover with water.
 - **Note 1:** 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 75 μ m (No. 200) 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.
- 4. Agitate vigorously to ensure complete separation of the material finer than 75 μ m (No. 200) from coarser particles and bring the fine material into suspension above the coarser material. Avoid degradation of the sample when using a mechanical washing device.
- 5. Immediately pour the wash water containing the suspended material over the nested sieves; be careful not to pour out the coarser particles or over fill the 75 μ m (No. 200) sieve.

- 6. Add water to cover material remaining in the container, agitate, and repeat Step 5. Continue until the wash water is reasonably clear.
- 7. Remove the upper sieve and return material retained to the washed test sample.
- 8. Rinse the material retained on the 75 μ m (No. 200) sieve until water passing through the sieve is reasonably clear and detergent or dispersing agent is removed, if used.
- 9. Return all material retained on the 75 μ m (No. 200) sieve to the container by rinsing into the washed sample.
 - **Note 2:** Excess water may be carefully removed with a bulb syringe; the removed water must be discharged back over the 75 μ m (No. 200) sieve to prevent loss of fines.
- 10. Dry the washed test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the dry mass.
- 11. Select sieves required by the specification and those necessary to avoid overloading. With a pan on bottom, nest the sieves increasing in size starting with the 75 μ m (No. 200).
- 12. Place the test sample, or a portion of the test sample, on the top sieve. Sieves may already be in the mechanical shaker, if not place sieves in mechanical shaker and shake for the minimum time determined to provide complete separation for the sieve shaker being used (approximately 10 minutes, the time determined by Annex A).
 - **Note 3:** Excessive shaking (more than 10 minutes) may result in degradation of the sample.
- 13. Determine and record the individual or cumulative mass retained for each sieve and in the pan. Ensure that all material trapped in full openings of the sieve are removed and included in the mass retained.
 - **Note 4:** For sieves 4.75 mm (No. 4) and larger, check material trapped in less than a full opening by sieving over a full opening. Use coarse wire brushes to clean the $600 \mu m$ (No. 30) and larger sieves, and soft bristle brushes for smaller sieves.
 - **Note 5:** In the case of coarse/fine aggregate mixtures, distribute the minus 4.75 mm (No. 4) among two or more sets of sieves to prevent overloading of individual sieves.
- 14. Perform the Check Sum calculation Verify the total mass after sieving agrees with the dry mass before sieving to within 0.3 percent. The dry mass before sieving is the dry mass after wash or the original dry mass (M) if performing the sieve analysis without washing. Do not use test results for acceptance if the Check Sum result is greater than 0.3 percent.
- 15. Calculate the total percentages passing, and the individual or cumulative percentages retained to the nearest 0.1 percent by dividing the individual sieve masses or cumulative sieve masses by the total mass of the initial dry sample (M).
- 16. Report total percent passing to 1 percent except report the 75 μ m (No. 200) sieve to 0.1 percent.

Method A Calculations

Check Sum

$$Check \ Sum = \frac{dry \ mass \ before \ seiving - total \ mass \ after \ sieving}{dry \ mass \ before \ sieving} \times 100$$

Percent Retained

$$IPR = \frac{IMR}{M} \times 100$$
 or $CPR = \frac{CMR}{M} \times 100$

Where:

IPR = Individual Percent Retained CPR = Cumulative Percent Retained M = Total Dry Sample mass before washing IMR = Individual Mass Retained

CMR = Cumulative Mass Retained

Percent Passing (PP)

$$PP = PPP - IPR$$
 or $PP = 100 - CPR$

Where:

PP = Percent Passing PPP = Previous Percent Passing

Method A Example Individual Mass Retained

Dry mass of total sample before washing (M): 5168.7 g

Dry mass of sample after washing: 4911.3 g

Total mass after sieving is equals

Sum of Individual Masses Retained (IPR), including pan: 4905.9 g

Amount of 75μm (No. 200) minus washed out (5168.7 g - 4911.3 g): 257.4 g

Check Sum

Check Sum =
$$\frac{4911.3 \ g - 4905.9 \ g}{4911.3 \ g} \times 100 = 0.1\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Individual Percent Retained (IPR) for 9.5 mm (% in) sieve:

$$CPR = \frac{1343.9 \ g}{5168.7 \ g} \times 100 = 26.0\%$$

Percent Passing (PP) 9.5 mm (% in) sieve:

$$PP = 86.0\% - 12.0\% = 74.0\%$$

Reported Percent Passing = 74 percent

As shown in the following table:

Gradation on All Sieves Individual Mass

Sieve Si:	ze mm (in)	Individual Mass Retained g (IMR)	Individual Percent Retained (IPR)	Percent Passing (PP)	Reported Percent Passing*
19.0	(3/4)	0	0	100.0	100
12.5	(1/2)	724.7	14.0	86.0	86
9.5	(3/8)	619.2	12.0	74.0	74
4.75	(No. 4)	1189.8	23.0	51.0	51
2.36	(No. 8)	877.6	17.0	34.0	34
1.18	(No. 16)	574.8	11.1	22.9	23
0.600	(No. 30)	329.8	6.4	16.5	17
0.300	(No. 50)	228.5	4.4	12.1	12
0.150	(No. 100)	205.7	4.0	8.1	8
0.075	(No. 200)	135.4	2.6	5.5	5.5
F	an	20.4			

^{*}Report total percent passing to 1 percent except report the 75 µm (No. 200) sieve to 0.1 percent.

Method A Example Cumulative Mass Retained

Dry mass of total sample before washing (M): 5168.7 g

Dry mass of sample after washing: 4911.3 g

Total mass after sieving is equals

Final Cumulative Mass Retained (CMR) in pan: 4905.9 g

Amount of 75μm (No. 200) minus washed out (5168.7 g - 4911.3 g): 257.4 g

Check Sum

Check Sum =
$$\frac{4911.3 \ g - 4905.9 \ g}{4911.3 \ g} \times 100 = 0.1\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Cumulative Percent Retained (CPR) for 9.5 mm (% in) sieve:

$$CPR = \frac{1343.9 \ g}{5168.7 \ g} \times 100 = 26.0\%$$

Percent Passing (PP) 9.5 mm (% in) sieve:

$$PP = 86.0\% - 12.0\% = 74.0\%$$

Reported Percent Passing = 74 percent

As shown in the following table:	As	shown	in	the	follo	owing	table:
----------------------------------	----	-------	----	-----	-------	-------	--------

Gradation on All Sieves									
Sieve Size mm (in)		Cumulative Mass Retained g (CMR)	Cumulative Percent Retained (CPR)	Percent Passing (PP)	Reported Percent Passing*				
19.0	(3/4)	0	0.0	100.0	100				
12.5	(1/2)	724.7	14.0	86.0	86				
9.5	(3/8)	1343.9	26.0	74.0	74				
4.75	(No. 4)	2533.7	49.0	51.0	51				
2.36	(No. 8)	3411.3	66.0	34.0	34				
1.18	(No. 16)	3986.1	77.1	22.9	23				
0.600	(No. 30)	4315.9	83.5	16.5	17				
0.300	(No. 50)	4544.4	87.9	12.1	12				
0.150	(No. 100)	4750.1	91.9	8.1	8				
0.075	(No. 200)	4885.5	94.5	5.5	5.5				
Pan		4905.9							
* Report total percent passing to 1 percent except report the 75 µm (No. 200) sieve to 0.1 percent.									

Procedure Method B

- 1. Dry the test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the total dry mass of the sample to the nearest 0.1 percent or 0.1 g. Designate this mass as M.
 - When the specification does not require the amount of material finer than 75 μ m (No. 200) be determined by washing, skip to Step 11.
- 2. Nest a protective sieve, such as a 2.0 mm (No. 10), above the 75 μ m (No. 200) sieve.
- 3. Place the test sample in a container and cover with water.
 - Note 1: 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 75 μm (No. 200) 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.
- 4. Agitate vigorously to ensure complete separation of the material finer than 75 μ m (No. 200) from coarser particles and bring the fine material into suspension above the coarser material. Avoid degradation of the sample when using a mechanical washing device.
- 5. Immediately pour the wash water containing the suspended material over the nested sieves; be careful not to pour out the coarser particles or over fill the 75 μ m (No. 200) sieve.
- 6. Add water to cover material remaining in the container, agitate, and repeat Step 5. Continue until the wash water is reasonably clear.
- 7. Remove the upper sieve and return material retained to the washed test sample.

- 8. Rinse the material retained on the 75 μ m (No. 200) sieve until water passing through the sieve is reasonably clear and detergent or dispersing agent is removed, if used.
- 9. Return all material retained on the 75 μ m (No. 200) sieve to the container by rinsing into the washed sample.
 - **Note 2:** Excess water may be carefully removed with a bulb syringe; the removed water must be discharged back over the 75 μ m (No. 200) sieve to prevent loss of fines.
- 10. Dry the washed test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the dry mass.
- 11. Select sieves required by the specification and those necessary to avoid overloading. With a pan on bottom, nest the sieves increasing in size starting with the 4.75 mm (No. 4).
- 12. Place the test sample, or a portion of the test sample, on the top sieve. Sieves may already be in the mechanical shaker, if not place the sieves in the mechanical shaker and shake for the minimum time determined to provide complete separation for the sieve shaker being used (approximately 10 minutes, the time determined by Annex A).
 - **Note 3:** Excessive shaking (more than 10 minutes) may result in degradation of the sample.
- 13. Determine and record the individual or cumulative mass retained for each sieve. Ensure that all particles trapped in full openings of the sieve are removed and included in the mass retained.
 - **Note 4:** For sieves 4.75 mm (No. 4) and larger, check material trapped in less than a full opening by sieving over a full opening. Use coarse wire brushes to clean the $600 \mu m$ (No. 30) and larger sieves, and soft hair bristle for smaller sieves.
- 14. Determine and record the mass of the minus 4.75 mm (No. 4) material in the pan. Designate this mass as M_1 .
- 15. Perform the Coarse Check Sum calculation Verify the total mass after coarse sieving agrees with the dry mass before sieving to within 0.3 percent. The dry mass before sieving is the dry mass after wash or the original dry mass (M) if performing the sieve analysis without washing. Do not use test results for acceptance if the Check Sum result is greater than 0.3 percent.
- 16. Reduce the minus 4.75 mm (No. 4) according to the FOP for AASHTO R 76 to produce a sample with a minimum mass of 500 g. Determine and record the mass of the minus 4.75 mm (No. 4) split, designate this mass as M_2 .
- 17. Select sieves required by the specification and those necessary to avoid overloading. With a pan on bottom, nest the sieves increasing in size starting with the 75 μ m (No. 200) up to, but not including, the 4.75 mm (No. 4) sieve.
- 18. Place the test sample portion on the top sieve and place the sieves in the mechanical shaker. Shake for the minimum time determined to provide complete separation for the sieve shaker being used (approximately 10 minutes, the time determined by Annex A).

- 19. Determine and record the individual or cumulative mass retained for each sieve and in the pan. Ensure that all particles trapped in full openings of the sieve are removed and included in the mass retained.
 - **Note 4:** For sieves 4.75 mm (No. 4) and larger, check material trapped in less than a full opening by sieving over a full opening. Use coarse wire brushes to clean the 600 μ m (No. 30) and larger sieves, and soft hair bristle for smaller sieves.
- 20. Perform the Fine Check Sum calculation Verify the total mass after sieving agrees with the dry mass before sieving (M_2) to within 0.3 percent. Do not use test results for acceptance if the Check Sum result is greater than 0.3 percent.
- 21. Calculate to the nearest 0.1 percent, the Individual Mass Retained (IMR) or Cumulative Mass Retained (CMR) of the size increment of the reduced sample and the original sample.
- 22. Calculate the total percent passing.
- 23. Report total percent passing to 1 percent except report the 75 μm (No. 200) sieve to 0.1 percent.

Method B Calculations

Check Sum

$$\textit{Coarse Check Sum} = \frac{\textit{dry mass before sieveing} - \textit{total mass after coarse sieving}}{\textit{dry mass before sieving}} \times 100$$

Fine Check Sum =
$$\frac{M_2 - total \ mass \ after \ fine \ sieving}{M_2} \times 100$$

Percent Retained for 4.75 mm (No. 4) and larger

$$IPR = \frac{IMR}{M} \times 100$$
 or $CPR = \frac{CMR}{M} \times 100$

Where:

IPR = Individual Percent Retained

CPR = Cumulative Percent Retained

M = Total dry test sample mass before washing

IMR = Individual Mass Retained

CMR = Cumulative Mass Retained

Percent Passing (PP) for 4.75 mm (No. 4) and larger

$$PP = PPP - IPR$$
 or $PP = 100 - CPR$

Where:

PP = Percent Passing

PPP = Previous Percent Passing

Minus 4.75mm (No. 4) adjustment factor (R)

The mass of material retained for each sieve is multiplied by the adjustment factor, the total mass of the minus 4.75 mm (No. 4) from the pan, M_1 , divided by the mass of the reduced split of minus 4.75 mm (No. 4), M_2 . For consistency, this adjustment factor is carried to three decimal places.

$$R = \frac{M_1}{M_2}$$

Where:

R = minus 4.75 mm (No. 4) adjustment factor

 M_1 = total mass of minus 4.75 mm (No. 4) before reducing

 M_2 = mass of the reduced split of minus 4.75 mm (No. 4)

Adjusted Individual Mass Retained (AIMR):

$$AIMR = R \times B$$

Where:

AIMR = Adjusted Individual Mass Retained

R = minus 4.75 mm (No. 4) adjustment factor

B = individual mass of the size increment in the reduced portion sieved

Adjusted Cumulative Mass Retained (ACMR)

$$ACMR = (R \times B) + D$$

Where:

ACMR = Adjusted Cumulative Mass Retained

R = minus 4.75 mm (No. 4) adjustment factor

B = cumulative mass of the size increment in the reduced portion sieved

D = cumulative mass of plus 4.75mm (No. 4) portion of sample

Method B Example Individual Mass Retained

Dry mass of total sample, before washing: 3214.0 g

Dry mass of sample after washing: 3085.1 g

Total mass after sieving

Sum of Individual Masses Retained (IPR) and pan 3085.0 g

Amount of 75 μm (No. 200) minus washed out (3214.0 g – 3085.1 g): 128.9 g

Coarse Check Sum

Coarse Check Sum =
$$\frac{3085.1 \ g - 3085.0 \ g}{3085.1 \ g} \times 100 = 0.0\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Individual Percent Retained (IPR) for 9.5 mm (% in) sieve

$$IPR = \frac{481.4 \ g}{3214.0 \ g} \times 100 = 15.0\%$$

Percent Passing (PP) for 9.5 mm (% in) sieve:

$$PP = 95.0\% - 15.0\% = 80.0\%$$

As shown in the following table:

Gradation on Coarse Sieves Individual Mass

Sieve Size mm (in)		Individual Mass Retained g (IMR)	Individual Percent Retained (IPR)	Percent Passing (PP)			
16.0	(5/8)	0	0	100			
12.5	(1/2)	161.1	5.0	95.0			
9.50	(3/8)	481.4	15.0	80.0			
4.75	(No. 4)	475.8	14.8	65.2			
Pa	an	1966.7 (M ₁)					
7	Total mass after sieving = sum of sieves + pan = 3085.0						
	Dry mass of total sample, before washing (M): 3214.0 g						

Fine Test Sample

The pan, M_1 (1966.7 g), was reduced according to the FOP for AASHTO R 76, to at least 500 g. In this case, the reduced mass was determined to be **512.8 g**. This is M_2 .

The reduced mass was sieved.

Total mass after sieving equals

Sum of Individual Masses Retained (IPR) and pan

511.8 g

Fine Check Sum

Fine Check Sum =
$$\frac{512.8 \ g - 511.8 \ g}{512.8 \ g} \times 100 = 0.2\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Adjustment Factor (R) for Adjusted Individual Mass Retained (AIMR) on minus 4.75 (No. 4) sieves

The mass of material retained for each sieve is multiplied by the adjustment factor (R) carried to three decimal places.

$$R = \frac{M_1}{M_2} = \frac{1,966.7 \ g}{512.8 \ g} = 3.835$$

Where:

R = minus 4.75 mm (No. 4) adjustment factor

 M_1 = total mass of minus 4.75 mm (No. 4) from the pan

 M_2 = mass of the reduced split of minus 4.75 mm (No. 4)

Each "individual mass retained" on the fine sieves must be multiplied by R to obtain the Adjusted Individual Mass Retained.

Adjusted Individual Mass Retained (AIMR) for 2.00 mm (No. 10) sieve

$$AIMR = 3.835 \times 207.1 g = 794.2 g$$

Individual Percent Retained (IPR) for 2.00 mm (No. 10) sieve:

$$IPR = \frac{794.2 \ g}{3214.0 \ g} \times 100 = 24.7\%$$

Percent Passing (PP) 2 mm (No. 10) sieve:

$$PP = 65.2\% - 24.7\% = 40.5\%$$

As shown in the following table:

Final Gradation on All Sieves Individual Mass

		Individual Mass		Individual Percent	Percent	Reported Percent		
Sieve Siz	e mm (in)	Retained, g (IMR)	Retained (AIMR)	Retained (IPR)	Passing (PP)	Passing*		
16.0	(5/8)	0		0.0	100.0	100		
12.5	(1/2)	161.1		5.0	95.0	95		
9.5	(3/8)	481.4		15.0	80.0	80		
4.75	(No. 4)	475.8		14.8	65.2	65		
2.00	(No. 10)	207.1 × 3.835	794.2	24.7	40.5	41		
0.425	(No. 40)	187.9 × 3.835	720.6	22.4	18.1	18		
0.210	(No. 80)	59.9 × 3.835	229.7	7.1	11.0	11		
0.075	(No. 200)	49.1 × 3.835	188.3	5.9	5.1	5.1		
Pan 7.8 × 3.835			29.9					
	Total Sum of masses on fine sieves + pan = 511.8							
		Dry mass of to	tal sample, before	washing: 3214	1.0 g			

^{*} Report total percent passing to 1 percent except report the 75 μm (No. 200) sieve to 0.1 percent.

Method B Example Cumulative Mass Retained

Dry mass of total sample, before washing: 3214.0 g

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

Total mass after sieving equals

Cumulative Mass Retained (CMR) on the 4.75 (No. 4) plus the pan: 3085.0 g

Amount of 75 μm (No. 200) minus washed out (3214.0 g – 3085.1 g): 128.9 g

Coarse Check Sum

Coarse Check Sum =
$$\frac{3085.1 g - 3085.0 g}{3085.1 g} \times 100 = 0.0\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Cumulative Percent Retained (CPR) for 9.5 mm (% in) sieve

$$CPR = \frac{642.5 \, g}{3214.0 \, g} \times 100 = 20.0\%$$

Percent Passing (PP) for 9.5 mm (% in) sieve

$$PP = 100.0\% - 20.0\% = 80.0\%$$

Reported Percent Passing = 80 percent

As shown in the following table:

Gradation on Coarse Sieves Cumulative Mass

Sieve Size mm (in)		Cumulative Mass Retained g (CMR)	Cumulative Percent Retained (CPR)	Percent Passing (PP)			
16.0	(5/8)	0	0	100			
12.5	(1/2)	161.1	5.0	95.0			
9.50	(3/8)	642.5	20.0	80.0			
4.75	(No. 4)	1118.3 (D)	34.8	65.2			
Mass	Mass in Pan 1966.7 (M ₁)						
	Cumulative sieved mass: 1118.3 + 1966.7 = 3085.0						
	Dry mass of total sample, before washing (M): 3214.0 g						

The mass of minus 4.75 mm (No. 4) material in the pan, M_1 (1966.7 g), was reduced according to the FOP for AASHTO R 76, to at least 500 g. In this case, the reduced mass was determined to be **512.8** g. This is M_2 .

The reduced mass was sieved.

Total mass after sieving equals

Final Cumulative Mass Retained (CMR) in pan:

511.8 g

Fine Check Sum

Fine Check Sum =
$$\frac{512.8 \ g - 511.8 \ g}{512.8 \ g} \times 100 = 0.2\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

The cumulative mass of material retained for each sieve is multiplied by the adjustment factor (R) carried to three decimal places and added to the cumulative mass retained on the 4.75 mm (No. 4) sieve, D, to obtain the Adjusted Cumulative Mass Retained (ACMR).

Adjustment factor (R) for Cumulative Mass Retained (CMR) in minus 4.75 (No. 4) sieves

$$R = \frac{M_1}{M_2} = \frac{1,966.7 \ g}{512.8 \ g} = 3.835$$

Where:

R = minus 4.75 mm (No. 4) adjustment factor M_1 = total mass of minus 4.75 mm (No. 4) from the pan M_2 = mass of the reduced split of minus 4.75 mm (No. 4)

Adjusted Cumulative Mass Retained (ACMR) for the 2.00 mm (No. 10) sieve

$$ACMR = 3.835 \times 207.1 g = 794.2 g$$

Total Cumulative Mass Retained (TCMR) for the 2.00 mm (No. 10) sieve

$$TCMR = 794.2 g + 1118.3 g = 1912.5 g$$

Cumulative Percent Retained (CPR) for 2.00 mm (No. 10) sieve:

$$CPR = \frac{1912.5 \ g}{3214.0 \ g} \times 100 = 59.5\%$$

Percent Passing (PP) 2.00 mm (No. 10) sieve:

$$PP = 100.0\% - 59.5\% = 40.5\%$$

Reported Percent Passing = 41 percent

As shown in the following table:

Final Gradation on All Sieves Cumulative Mass

Sieve Size mm (in)		Cumulative Mass Retained g (CMR)	Adjusted Cumulative Mass Retained, g (ACMR)	Total Cumulative Mass Retained g (TCMR)	Cumulative Percent Retained (CPR)	Percent Passing (PP)	Reported Percent Passing*	
16.0	(5/8)	0		0	0.0	100.0	100	
12.5	(1/2)	161.1		161.1	5.0	95.0	95	
9.5	(3/8)	642.5		642.5	20.0	80.0	80	
4.75	(No. 4)	1118.3		1118.3	34.8	65.2	65	
2.00	(No. 10)	207.1 × 3.835	794.2 + 1118.3	1912.5	59.5	40.5	41	
0.425	(No. 40)	395.0 × 3.835	1514.8 + 1118.3	2633.1	81.9	18.1	18	
0.210	(No. 80)	454.9 × 3.835	1744.5 + 1118.3	2862.8	89.1	10.9	11	
0.075	(No. 200)	504.0 × 3.835	1932.8 + 1118.3	3051.1	94.9	5.1	5.1	
Pa	Pan 511.8 × 3.835		1962.8 + 1118.3	3081.1				
	Total sum of masses on fine sieves + pan = 511.8							
	Dry mass of total sample, before washing: 3214.0 g							

 $^{^{\}ast}$ Report total percent passing to 1 percent except report the 75 μm (No. 200) sieve to 0.1 percent.

Procedure Method C

- 1 Dry the test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the total dry mass of the sample to the nearest 0.1 percent or 0.1 g. Designate this mass as *M*.
- 2. Break up any aggregations or lumps of clay, silt or adhering fines to pass the 4.75 mm (No. 4) sieve.
- 3. Select sieves required by the specification and those necessary to avoid overloading. With a pan on bottom, nest the sieves increasing in size starting with the 4.75 mm (No. 4) sieve.
- 4. Place the sample, or a portion of the sample, on the top sieve. Sieves may already be in the mechanical shaker, if not place the sieves in the mechanical shaker and shake for the minimum time determined to provide complete separation for the sieve shaker being used (approximately 10 minutes, the time determined by Annex A).
 - **Note 3:** Excessive shaking (more than 10 minutes) may result in degradation of the sample.
- 5. Determine and record the cumulative mass retained for each sieve. Ensure that all material trapped in full openings of the sieve are removed and included in the mass retained.
 - **Note 4:** For sieves 4.75 mm (No. 4) and larger, check material trapped in less than a full opening sieving over a full opening. Use coarse wire brushes to clean the 600 μ m (No. 30) and larger sieves, and soft bristle brush for smaller sieves.
- 6. Determine and record the mass of the minus 4.75 mm (No. 4) material in the pan. Designate this mass as M_1 .
- 7. Perform the Coarse Check Sum calculation Verify the total mass after coarse sieving agrees with the dry mass before sieving (M) within 0.3 percent.
- 8. Reduce the minus 4.75 mm (No. 4) according to the FOP for AASHTO R 76, to produce a sample with a minimum mass of 500 g.
- 9. Determine and record the mass of the minus 4.75 mm (No. 4) split, designate this mass as M_3 .
- 10. Nest a protective sieve, such as a 2.0 mm (No. 10), above the 75 μ m (No. 200) sieve.
- 11. Place the test sample in a container and cover with water.
 - **Note 1:** 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 75 μ m (No. 200) 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.
- 12. Agitate vigorously to ensure complete separation of the material finer than 75 μ m (No. 200) from coarser particles and bring the fine material into suspension above the coarser material. Avoid degradation of the sample when using a mechanical washing device.

- 13. Immediately pour the wash water containing the suspended material over the nested sieves; be careful not to pour out the coarser particles or over fill the 75 μ m (No. 200) sieve.
- 14. Add water to cover material remaining in the container, agitate, and repeat Step 12. Repeat until the wash water is reasonably clear.
- 15. Remove the upper sieve and return material retained to the washed test sample.
- 16. Rinse the material retained on the 75 μ m (No. 200) sieve until water passing through the sieve is reasonably clear and detergent or dispersing agent is removed, if used.
- 17. Return all material retained on the 75 μ m (No. 200) sieve to the container by flushing into the washed sample.
 - **Note 2:** Excess water may be carefully removed with a bulb syringe; the removed water must be discharged back over the 75 μ m (No. 200) sieve to prevent loss of fines.
- 18. Dry the washed test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the dry mass, designate this mass as *dry mass before sieving*.
- 19. Select sieves required by the specification and those necessary to avoid overloading. With a pan on bottom, nest the sieves increasing in size starting with the 75 μ m (No. 200) sieve up to, but not including, the 4.75 mm (No. 4) sieve.
- 20. Place the sample on the top sieve. Place the sieves in the mechanical shaker and shake for the minimum time determined to provide complete separation for the sieve shaker being used (approximately 10 minutes, the time determined by Annex A).
 - **Note 3:** Excessive shaking (more than 10 minutes) may result in degradation of the sample.
- 21. Determine and record the cumulative mass retained for each sieve. Ensure that all material trapped in full openings of the sieve are removed and included in the mass retained.
 - **Note 4:** For sieves 4.75 mm (No. 4) and larger, check material trapped in less than a full opening by sieving over a full opening. Use coarse wire brushes to clean the 600 μ m (No. 30) and larger sieves, and soft bristle brushes for smaller sieves.
- 22. Perform the Fine Check Sum calculation Verify the total mass after fine sieving agrees with the dry mass before sieving within 0.3 percent. Do not use test results for acceptance if the Check Sum is greater than 0.3 percent.
- 23. Calculate the Cumulative Percent Retained (CPR) and Percent Passing (PP) for the 4.75 mm (No. 4) and larger.
- 24. Calculate the Cumulative Percent Retained (CPR_{-#4}) and the Percent Passing (PP_{-#4}) for minus 4.75 mm (No. 4) split and Percent Passing (PP) for the minus 4.75 mm (No. 4).
- 25. Report total percent passing to 1 percent except report the 75 μ m (No. 200) sieve to 0.1 percent.

Method C Calculations

Check Sum

$$Coarse\ check\ sum = \frac{M-total\ mass\ after\ coarse\ sieving}{M} \times 100$$

$$Fine\ check\ sum = \frac{dry\ mass\ before\ sieving-total\ mass\ after\ fine\ sieving}{dry\ mass\ before\ sieving} \times 100$$

where:

M = Total dry sample mass before washing

Cumulative Percent Retained (CPR) for 4.75 mm (No. 4) sieve and larger

$$CPR = \frac{CMR}{M} \times 100$$

where:

CPR = Cumulative Percent Retained of the size increment for the total sample CMR = Cumulative Mass Retained of the size increment for the total sample M = Total dry sample mass before washing

Percent Passing (PP) 4.75 mm (No. 4) sieve and larger

$$PP = 100 - CPR$$

where:

PP = Percent Passing of the size increment for the total sample CPR = Cumulative Percent Retained of the size increment for the total sample

Or, calculate PP for sieves larger than 4.75 mm (No. 4) sieve without calculating CPR

$$\frac{M - CMR}{M} \times 100$$

Cumulative Percent Retained (CPR_{#4}) for minus 4.75 mm (No. 4) split

$$CPR_{-\#4} = \frac{CMR_{-\#4}}{M_3} \times 100$$

Where:

CPR_{.#4} = Cumulative Percent Retained for the sieve sizes of M_3 CMR_{.#4} = Cumulative Mass Retained for the sieve sizes of M_3 M_3 = Total mass of the minus 4.75 mm (No. 4) split before washing

Percent Passing (PP_{-#4}) for minus 4.75 mm (No. 4) split

$$PP_{-#4} = 100 - CPR_{-#4}$$

where:

 $PP_{-#4}$ = Percent Passing for the sieve sizes of M_3 $CPR_{-#4}$ = Cumulative Percent Retained for the sieve sizes of M_3

Percent Passing (PP) for sieves smaller than 4.75 mm (No. 4) sieve

$$PP = \frac{(PP_{-\#4} \times \#4 \ PP)}{100}$$

where:

PP = Total Percent Passing

 $PP_{-#4}$ = Percent Passing for the sieve sizes of M_3 #4 PP = Total Percent Passing the 4.75 mm (No. 4) sieve

Or, calculate PP for sieves smaller than 4.75 mm (No. 4) sieve without calculating CPR_{-#4} and PP_{-#4}

$$PP = \frac{\#4 \ PP}{M_3} \times (M_3 - CMR_{-\#4})$$

Where:

PP = Total Percent Passing

#4 PP = Total Percent Passing the 4.75 mm (No. 4) sieve

 M_3 = Total mass of the minus 4.75 mm (No. 4) split before washing

 $CMR_{-\#4}$ = Cumulative Mass Retained for the sieve sizes of M_3

Method C Example

Dry Mass of total sample (M):

3304.5 g

Total mass after sieving equals

Cumulative Mass Retained (CMR) on the 4.75 (No. 4) plus the pan:

3085.0 g

Coarse Check Sum

Coarse Check Sum =
$$\frac{3304.5 \ g - 3304.5 \ g}{3304.5 \ g} \times 100 = 0.0\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Cumulative Percent Retained (CPR) for the 9.5 mm (% in) sieve:

$$CPR = \frac{604.1 \, g}{3304.5 \, g} \times 100 = 18.3\%$$

Percent Passing (PP) for the 9.5 mm (% in) sieve:

$$PP = 100.0\% - 18.3\% = 81.7\%$$

Reported Percent Passing = 82 percent

Example for Alternate Percent Passing (PP) formula for the 9.5 mm (3/8 in) sieve:

$$PP = \frac{3304.5 - 604.1}{3304.5} \times 100 = 81.7\%$$

Reported Percent Passing = 82 percent

Gradation on Coarse Sieves Cumulative Mass

Sieve Siz	e mm (in)	Cumulative Mass Retained, g (CMR)		Percent Passing (PP)	Reported Percent Passing*		
Sieve Siz		Retained, g (Civin)	(CFR)	(FF)	rassilig		
16.0	(5/8)	0	0.0	100.0	100		
12.5	1/2)	125.9	3.8	96.2	96		
9.5	(3/8)	604.1	18.3	81.7	82		
4.75	(No. 4)	1295.6	39.2	60.8 (#4 PP)	61		
Mass in Pan		2008.9					
	Cumulative sieved mass: 1295.6 + 2008.9 = 3304.5						
	Total Dry Sample (M) = 3304.5						

Fine Test Sample

The pan (2008.9 g) was reduced according to the FOP for AASHTO R 76, to at least 500 g. In this case, the reduced mass was determined to be **527.6** g. This is M_3 .

Dry Mass of minus 4.75mm (No. 4) reduced portion before wash (M_3): 527.6 g

Dry Mass of minus 4.75mm (No. 4) reduced portion after wash: 495.3 g

Total mass after sieving equals

Final Cumulative Mass Retained (CMR) in pan: 495.1 g

Fine Check Sum

Fine Check Sum =
$$\frac{495.3 \ g - 495.1 \ g}{495.3 \ g} \times 100 = 0.04\%$$

The result is less than 0.3 percent therefore the results can be used for acceptance purposes.

Cumulative Percent Retained ($CPR_{-#4}$) for minus 4.75 mm (No. 4) for the 2.0 mm (No. 10) sieve:

$$CPR_{-#4} = \frac{194.3 \ g}{527.6 \ g} \times 100 = 36.8\%$$

Percent Passing (PP_{-#4}) for minus 4.75 mm (No. 4) for the 2.0 mm (No. 10) sieve:

$$PP_{-#4} = 100.0\% - 36.8\% = 63.2\%$$

Gradation on Minus No. 4 Sieves Cumulative Mass

Sieve Size mm (in)		Cumulative Mass Retained g (CMR _{-#4})	Cumulative Percent Retained _{-#4} (CPR _{-#4})	Percent Passing _{-#4} (PP _{-#4})
2.0	(No. 10)	194.3	36.8	63.2
0.425	(No. 40)	365.6	69.3	30.7
0.210	(No. 80)	430.8	81.7	18.3
0.075	(No. 200)	484.4	91.8	8.2
Pan		495.1		

Dry mass before washing (M_3): 527.6 g

Dry mass after washing: 495.3 g

Percent Passing (PP) for the 2.0 mm (No. 10) sieve for the entire test sample:

#4 PP (Total Percent Passing the 4.75 mm (No. 4) sieve) = 60.8 percent

$$PP = \frac{63.2\% \times 60.8\%}{100} = 38.4\%$$

Reported Percent Passing = 38 percent

As shown in the following table:

Final Gradation on All Sieves Cumulative Mass

Sieve Size mm (in)		Cumulative Mass Retained, g (CMR)	Cumulative Percent Retained (CPR)	Percent Passing (PP _{-#4})	Percent Passing (PP)	Reported Percent Passing*
16.0	(5/8)	0	0.0		100.0	100
12.5	(1/2)	125.9	3.8		96.2	96
9.5	(3/8)	604.1	18.3		81.7	82
4.75	(No. 4)	1295.6	39.2		60.8 (#4 PP)	61
2.0	(No. 10)	194.3	36.8	63.2	38.4	38
0.425	(No. 40)	365.6	69.3	30.7	18.7	19
0.210	(No. 80)	430.8	81.7	18.3	11.1	11
0.075	(No. 200)	484.4	91.8	8.2	5.0	5.0
Р	an	495.1				

 $^{^{\}ast}$ Report total percent passing to 1 percent except report the 75 μm (No. 200) sieve to 0.1 percent.

Example for Alternate Percent Passing (PP) for the 2.0 mm (No. 10) sieve for the entire test sample:

#4 PP (Total Percent Passing the 4.75 mm (No. 4) sieve) = 60.8 percent

$$PP = \frac{60.8\%}{527.6} \times (527.6 - 194.3) = 38.4\%$$

Reported Percent Passing = 38 percent

As shown in the following table:

Final Gradation on All Sieves Cumulative Mass

Sieve Size mm (in)		Cumulative Mass Retained, g (CMR)	Cumulative Percent Retained (CPR)	Determined Percent Passing (PP)	Reported Percent Passing*			
16.0	(5/8)	0	0.0	100.0	100			
12.5	(1/2)	125.9	3.8	96.2	96			
9.5	(3/8)	604.1	18.3	81.7	82			
4.75	(No. 4)	1295.6	39.2	60.8 (#4 PP)	61			
2.0	(No. 10)	194.3		38.4	38			
0.425	(No. 40)	365.6		18.7	19			
0.210	(No. 80)	430.8		11.2	11			
0.075	(No. 200)	484.4		5.0	5.0			
Pan 495.1		495.1						
Dry mas	Dry mass before washing (M ₃): 527.6 g							

^{*} Report total percent passing to 1 percent except report the 75 µm (No. 200) sieve to 0.1 percent.

Fineness Modulus

Fineness Modulus (FM) is used in determining the degree of uniformity of the 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 fine aggregate in PCC.

The sum of the cumulative percentages retained on specified sieves in the following table divided by 100 gives the FM.

Sample Calculation

		Example A				
		Percent				
			R	etained		
				On Spec'd		
Sieve Siz	ze mm (in)	Passing		Sieves*		
75*	(3)	100	0	0		
37.5*	(1½)	100	0	0		
19*	(3/4)	15	85	85		
9.5*	(3/8)	0	100	100		
4.75*	(No.4)	0	100	100		
2.36*	(No.8)	0	100	100		
1.18*	(No.16)	0	100	100		
0.60*	(No. 30	0	100	100		
0.30*	(No. 50)	0	100	100		
0.15*	(100)	0	100	100		
				∑ = 785		
				FM = 7.85		

Example B						
	Percent					
	R	etained				
Passing	On Spec'o					
100	0	0				
100	0	0				
100	0	0				
100	0	0				
100	0	0				
87	13	13				
69	31	31				
44	56	56				
18	82	82				
4	96	96				
		∑ = 278				
		FM = 2.78				

In decreasing size order, each * sieve is one-half the size of the preceding * sieve.

Report

- Results on forms approved by the agency
- Sample ID
- · Percent passing for each sieve
- · Individual mass retained for each sieve
- Individual percent retained for each sieve or
- Cumulative mass retained for each sieve
- Cumulative percent retained for each sieve
- FM to the nearest 0.01

Report percentages to the nearest 1 percent except for the percent passing the 75 μ m (No. 200) sieve, which shall be reported to the nearest 0.1 percent.

ANNEX A Time Evaluation

The sieving time for each mechanical sieve shaker shall be checked at least annually to determine the time required for complete separation of the test sample by the following method:

- 1. Shake the sample over nested sieves for approximately 10 minutes.
- 2. Provide a snug-fitting pan and cover for each sieve, and hold in a slightly inclined position in one hand.
- 3. Hand-shake each sieve by striking the side of the sieve sharply and with an upward motion against the heel of the other hand at the rate of about 150 times per minute, turning the sieve about one sixth of a revolution at intervals of about 25 strokes.

If more than 0.5 percent by mass of the total sample before sieving passes any sieve after one minute of continuous hand shaking adjust shaker time and re-check.

In determining sieving time for sieve sizes larger than 4.75 mm (No. 4), limit the material on the sieve to a single layer of particles.

ANNEX B Overload Determination

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

- For sieves with openings smaller than 4.75 mm (No. 4), the mass retained on any sieve shall not exceed 7 kg/m^2 (4 g/in²) of sieving surface.
- For sieves with openings 4.75 mm (No. 4) and larger, the mass, in grams shall not exceed the product of 2.5 × (sieve opening in mm) × (effective sieving area). See Table B1.

Table B1 Maximum Allowable Mass of Material Retained on a Sieve, g Nominal Sieve Size, mm (in) Exact size is smaller (see AASHTO T 27)

				·		
		203 dia (8)	305 dia (12)	305 by 305 (12 × 12)	350 by 350 (14 × 14)	372 by 580 (16 × 24)
			S	ieving Area m	12	
Sieve Size mm (in)		0.0285	0.0670	0.0929	0.1225	0.2158
90	(3½)	*	15,100	20,900	27,600	48,500
75	(3)	*	12,600	17,400	23,000	40,500
63	(2½)	*	10,600	14,600	19,300	34,000
50	(2)	3600	8400	11,600	15,300	27,000
37.5	(1½)	2700	6300	8700	11,500	20.200
25.0	(1)	1800	4200	5800	7700	13,500
19.0	(3/4)	1400	3200	4400	5800	10,200
16.0	(5/8)	1100	2700	3700	4900	8600
12.5	(1/2)	890	2100	2900	3800	6700
9.5	(3/8)	670	1600	2200	2900	5100
6.3	(1/4)	440	1100	1500	1900	3400
4.75	(No. 4)	330	800	1100	1500	2600
-4.75	(-No. 4)	200	470	650	860	1510

PERFORMANCE EXAM CHECKLIST

METHOD A

FOP for AASHTO T 27 Sieve Analysis of Fine and Coarse Aggregates

FOP for AASHTO T 11 Materials Finer Than 75 μm (No. 200) Sieve in Mineral Aggregate by Washing

Part	icipant Name Exam Date		
Reco	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Prod	cedure Element	Trial 1	Trial 2
1.	Minimum sample mass meets requirement of Table 1?		
2.	Test sample dried to a constant mass by FOP for AASHTO T 255?		
3.	Test sample cooled and mass determined to nearest 0.1 percent or 0.1 g?		
4.	Test sample placed in container and covered with water?		
5.	Contents of the container vigorously agitated?		
6.	Complete separation of coarse and fine particles achieved?		
7.	Wash water poured through nested sieves such as 2 mm (No. 10) and 75 μ m (No. 200)?		
8.	Operation continued until wash water is clear?		
9.	Material retained on sieves returned to washed sample?		
10.	Washed test sample dried to a constant mass by FOP for AASHTO T 255?		
11.	Washed test sample cooled and mass determined to nearest 0.1 percent or 0.1 g?		
12.	Test sample placed in nest of sieves specified? (Additional sieves may be used to prevent overloading as allowed in FOP.)		
13.	Material sieved in verified mechanical shaker for proper time?		
14.	Mass of material on each sieve and pan determined to 0.1 g?		
15.	Total mass of material after sieving agrees with mass before sieving to within 0.3 percent?		
16.	Percentages calculated to the nearest 0.1 percent and reported to the nearest whole number, except 75 μm (No. 200) which is reported to the nearest 0.1 percent?		
17.	Percentage calculations based on original dry sample mass?		
18.	Calculations performed properly?		
	t Attempt: Pass Fail Second Attempt: Pass Fail ature of Examiner WAQTC #:		

Comments:

Performance Exam Checklist

METHOD B

FOP for AASHTO T 27 Sieve Analysis of Fine and Coarse Aggregates

FOP for AASHTO T 11 Materials Finer Than 75 μm (No. 200) Sieve in Mineral Aggregate by Washing

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Minimum sample mass meets requirement of Table 1?		
2.	Test sample dried to a constant mass by FOP for AASHTO T 255?		
3.	Test sample cooled and mass determined to nearest 0.1 percent or 0.1 g?		
4.	Test sample placed in container and covered with water?		
5.	Contents of the container vigorously agitated?		
6.	Complete separation of coarse and fine particles achieved?		
7.	Wash water poured through nested sieves such as 2 mm (No. 10) and 75 μ m (No. 200)?		
8.	Operation continued until wash water is clear?		
9.	Material retained on sieves returned to washed sample?		
10.	Washed test sample dried to a constant mass by FOP for AASHTO T 255?		
11.	Washed test sample cooled and mass determined to nearest 0.1 percent or 0.1 g ?		
12.	Test sample placed in nest of sieves specified? (Additional sieves may be used to prevent overloading as allowed in FOP.)		
13.	Material sieved in verified mechanical shaker for proper time?		
14.	Mass of material on each sieve and pan determined to the nearest 0.1 percent or 0.1 g?		
15.	Total mass of material after sieving agrees with mass before sieving to within 0.3 percent?		
16.	Material in pan reduced in accordance with FOP for AASHTO R 76 to at least 500 g and weighed to the nearest 0.1 g?		
17.	Test sample placed in nest of sieves specified? (Additional sieves may be used to prevent overloading as allowed in FOP.)		
18.	Material sieved in verified mechanical shaker for proper time?		
19.	Mass of material on each sieve and pan determined to the nearest percent		

Proc	edure Element	iriai 1	Iriai 2
20.	Total mass of material after sieving agrees with mass before sieving to within 0.3 percent?		
21.	Percentages calculated to the nearest 0.1 percent and reported to the nearest whole number, except 75 μm (No. 200) which is reported to the nearest 0.1 percent?		
22.	Percentage calculations based on original dry sample mass?		
23.	Calculations performed properly?		

First Attempt: Pass Fail Second Attempt: Pass Fail

Signature of Examiner _____ WAQTC #:______

Comments:

WSDOT Errata to FOP for AASHTO T 30

Mechanical Analysis of Extracted Aggregate

WAQTC FOP for AASHTO T 30 has been adopted by WSDOT with the following changes:

Procedure

15. Step not recognized by WSDOT.

FOP for AASHTO T 30

Mechanical Analysis of Extracted Aggregate

Scope

This procedure covers mechanical analysis of aggregate recovered from asphalt mix samples in accordance with AASHTO T 30-15. This FOP utilizes the aggregate recovered from the ignition oven used in AASHTO T 308. AASHTO T 30 was developed for analysis of extracted aggregate and thus includes references to extracted bitumen and filter element, which do not apply in this FOP.

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

Apparatus

- Balance or scale: Capacity sufficient for the sample mass, accurate to 0.1 percent of the sample mass or readable to 0.1 g
- Sieves, meeting the requirements of FOP for AASHTO T 27/T 11.
- Mechanical sieve shaker, meeting the requirements of FOP for AASHTO T 27/T 11.
- Mechanical Washing Apparatus (optional)
- Suitable drying equipment, meeting the requirements of the 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

Sample Sieving

- In this procedure, it is required to shake the sample over nested sieves. Sieves are selected to furnish information required by specification. Intermediate sieves are added for additional information or to avoid overloading sieves, or both.
- The sieves are nested in order of increasing size from the bottom to the top, and the test sample, or a portion of the test sample, is placed on the top sieve.
- The loaded sieves are shaken in a mechanical shaker for approximately 10 minutes, refer to Annex A: Time Evaluation.

Mass Verification

Using the aggregate sample obtained from the FOP for AASHTO T 308, determine and record the mass of the sample, $M_{(T30)}$, to 0.1 g. This mass shall agree with the mass of the aggregate remaining after ignition, $M_{\rm f}$ from T 308, within 0.10 percent. If the variation exceeds 0.10 percent the results cannot be used for acceptance.

Calculation

$$\textit{Mass verification} = \frac{M_{f\,(T308)}\text{-}M_{(T30)}}{M_{f\,(T308)}} \times 100$$

Where:

 $M_{f(T308)}$ = Mass of aggregate remaining after ignition from the FOP for AASHTO T 308 $M_{(T30)}$ = Mass of aggregate sample obtained from the FOP for AASHTO T 308

Example:

Mass verification =
$$\frac{2422.5 g - 2422.3 g}{2422.5 g} \times 100 = 0.01\%$$

Where:

$$M_{f(T308)} = 2422.5 g$$

 $M_{(T30)} = 2422.3 g$

Procedure

- 1. Nest a sieve, such as a 2.0 mm (No. 10) or 1.18 mm (No. 16), above the $75\mu m$ (No. 200) sieve.
- 2. Place the test sample in a container and cover with water. Add a detergent, dispersing agent, or other wetting solution to the water to assure a thorough separation of the material finer than the $75\mu m$ (No. 200) 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.
- 3. Agitate vigorously to ensure complete separation of the material finer than $75\mu m$ (No. 200) from coarser particles and bring the fine material into suspension above the coarser material. Avoid degradation of the sample when using a mechanical washing device. Maximum agitation is 10 min.
 - **Note 1:** When mechanical washing equipment is used, the introduction of water, agitating, and decanting may be a continuous operation. Use care not to overflow or overload the $75\mu m$ (No. 200) sieve.
- 4. Immediately pour the wash water containing the suspended material over the nested sieves; be careful not to pour out the coarser particles or over fill the 75 μ m (No. 200) sieve.
- 5. Add water to cover material remaining in the container, agitate, and repeat Step 4. Continue until the wash water is reasonably clear.
- 6. Remove the upper sieve, return material retained to the washed sample.
- 7. Rinse the material retained on the 75 μ m (No. 200) sieve until water passing through the sieve is reasonably clear and detergent or dispersing agent is removed.

- 8. Return all material retained on the 75 μ m (No. 200) sieve to the washed sample by rinsing into the washed sample.
- 9. Dry the washed test sample to constant mass according to the FOP for AASHTO T 255. Cool to room temperature. Determine and record the "dry mass after washing."
- 10. Select sieves required by the specification and those necessary to avoid overloading. With a pan on bottom, nest the sieves increasing in size starting with the 75 μ m (No. 200).
- 11. Place the test sample, or a portion of the test sample, on the top sieve. Place sieves in mechanical shaker and shake for the minimum time determined to provide complete separation for the sieve shaker being used (approximately 10 minutes, the time determined by Annex A).
 - **Note 2:** Excessive shaking (more than 10 minutes) may result in degradation of the sample.
- 12. Determine and record the individual or cumulative mass retained for each sieve including the pan. Ensure that all material trapped in full openings of the sieves are removed and included in the mass retained.
 - **Note 3:** For sieves 4.75 mm (No. 4) and larger, check material trapped in less than a full opening by sieving over a full opening. Use coarse wire brushes to clean the $600 \mu m$ (No. 30) and larger sieves, and soft bristle brushes for smaller sieves.
- 13. Perform the Check Sum calculation Verify the total mass after sieving of material agrees with the dry mass after washing within 0.2 percent. Do not use test results for acceptance if the Check Sum result is greater than 0.2 percent.
- 14. Calculate the total percentages passing, and the individual or cumulative percentages retained, to the nearest 0.1 percent by dividing the individual sieve masses or cumulative sieve masses by the total mass of the initial dry sample.
- 15. Apply the Aggregate Correction Factor (ACF) to the calculated percent passing, as required in the FOP for AASHTO T 308 "Correction Factor," to obtain the reported percent passing.
- 16. Report total percent passing to 1 percent except report the 75 μ m (No. 200) sieve to 0.1 percent.

Calculations

Check Sum

$$check \; sum = \frac{dry \; mass \; after \; washing - total \; mass \; after \; sieving}{dry \; mass \; after \; washing} \; \times 100$$

Percent Retained

Individual

$$IPR = \frac{IMR}{M_{T30}} \times 100$$

Cumulative

$$CPR = \frac{CMR}{M_{T30}} \times 100$$

Where:

IPR = Individual Percent Retained

CPR = Cumulative Percent Retained

 M_{T30} = Total dry sample mass before washing

IMR = Individual Mass Retained CMR = Cumulative Mass Retained

Percent Passing

Individual

$$PP = PCP - IPR$$

Cumulative

$$PP = 100 - CPR$$

Where:

PP = Calculated Percent Passing

PCP = Previous Calculated Percent Passing

Reported Percent Passing

$$RPP = PP + ACF$$

Where:

RPP = Reported Percent Passing

ACF = Aggregate Correction Factor (if applicable)

Example

Dry mass of total sample, before washing (M_{T30}): 2422.3 g Dry mass of sample, after washing out the 75 μ m (No. 200) minus: 2296.2 g

Amount of 75 μm (No. 200) minus washed out (2422.3 g – 2296.2g): 126.1 g

Check sum

check sum =
$$\frac{2296.2 g - 2295.3 g}{2296.2 g} \times 100 = 0.04\%$$

This is less than 0.2 percent therefore the results can be used for acceptance purposes.

Percent Retained for the 75 µm (No. 200) sieve

$$IPR = \frac{63.5 \ g}{2422.3 \ g} \times 100 = 2.6\%$$
or

$$CPR = \frac{2289.6 \ g}{2422.3 \ g} \times 100 = 94.5\%$$

Percent Passing using IPR and PCP for the 75 μm (No. 200) sieve

$$PP = 8.1\% - 2.6\% = 5.5\%$$

Percent Passing using CPR for the 75 µm (No. 200) sieve

$$PP = 100.0\% - 94.5\% = 5.5\%$$

Reported Percent Passing

$$RPP = 5.5\% = (-0.6\%) = 4.9\%$$

Gradation on All Screens

Sieve Siz	ze mm (in)	Mass Retained (g) (MR)	Percent	Mass	Cumulative Percent Retained (CPR)	Percent Passing (PP)	Agg. Corr. Factor from T 308 (ACF)	Reported Percent Passing (RPP)
19.0	(3/4)	0.0		0.0	0	100.0		100
12.5	(1/2)	346.9	14.3	346.9	14.3	85.7		86
9.5	(3/8)	207.8	8.6	554.7	22.9	77.1		77
4.75	(No. 4)	625.4	25.8	1180.1	48.7	51.3		51
2.36	(No. 8)	416.2	17.2	1596.3	65.9	34.1		34
01.18	(No. 16)	274.2	11.3	1870.5	77.2	22.8		23
0.600	(No. 30)	152.1	6.3	2022.6	83.5	16.5		16
0.300	(No. 50)	107.1	4.4	2129.7	87.9	12.1		12
0.150	(No. 100)	96.4	4.0	2226.1	91.9	8.1		8
75 μm	(No. 200)	63.5	2.6	2289.6	94.5	5.5	-0.6	4.9
Р	Pan	5.7		2295.3				

^{*} Report total percent passing to 1 percent except report the 75 μ m (No. 200) sieve to 0.1 percent.

Report

- Results on forms approved by the agency
- Sample ID
- Depending on the agency, this may include:
 - Individual mass retained on each sieve
 - Individual percent retained on each sieve
 - Cumulative mass retained on each sieve
 - Cumulative percent retained on each sieve
 - Aggregate Correction Factor for each sieve from AASHTO T 308
 - Calculated percent passing each sieve to 0.1 percent
- Percent passing to the nearest 1 percent, except 75 μm (No. 200) sieve to the nearest 0.1 percent.

ANNEX A Time Evaluation

The minimum time requirement should be evaluated for each shaker at least annually by the following method:

- 1. Shake the sample over nested sieves for approximately 10 minutes.
- 2. Provide a snug-fitting pan and cover for each sieve, and hold in a slightly inclined position in one hand.
- 3. Hand-shake each sieve by striking the side of the sieve sharply and with an upward motion against the heel of the other hand at the rate of about 150 times per minute, turning the sieve about one sixth of a revolution at intervals of about 25 strokes.

If more than 0.5 percent by mass of the total sample before sieving passes any sieve after one minute of continuous hand sieving adjust shaker time and re-check.

In determining sieving time for sieve sizes larger than 4.75 mm (No. 4), limit the material on the sieve to a single layer of particles.

ANNEX B Overload Determination

- For sieves with openings smaller than 4.75 mm (No. 4), the mass retained on any sieve shall not exceed 7 kg/m^2 (4 g/in²) of sieving surface.
- For sieves with openings 4.75 mm (No. 4) and larger, the mass (in kg) shall not exceed the product of 2.5 × (sieve opening in mm) × (effective sieving area). See Table B1.

Additional sieves may be necessary to keep from overloading the specified sieves. The sample may also be sieved in increments or sieves with a larger surface area.

Table B1 Maximum Allowable Mass of Material Retained on a Sieve, g Nominal Sieve Size, mm (in) Exact size is smaller (see AASHTO T 27)

		203 dia (8)	305 dia (12)	305 by 305 (12 × 12)	350 by 350 (14 × 14)	372 by 580 (16 × 24)	
		Sieving Area m ²					
Sieve Siz	ze mm (in)	0.0285	0.0670	0.0929	0.1225	0.2158	
90	(3½)	*	15,100	20,900	27,600	48,500	
75	(3)	*	12,600	17,400	23,000	40,500	
63	(2½)	*	10,600	14,600	19,300	34,000	
50	(2)	3600	8400	11,600	15,300	27,000	
37.5	(1½)	2700	6300	8700	11,500	20,200	
25.0	(1)	1800	4200	5800	7700	13,500	
19.0	(3/4)	1400	3200	4400	5800	10,200	
16.0	(5/8)	1100	2700	3700	4900	8600	
12.5	(1/2)	890	2100	2900	3800	6700	
9.5	(3/8)	670	1600	2200	2900	5100	
6.3	(1/4)	440	1100	1500	1900	3400	
4.75	(No. 4)	330	800	1100	1500	2600	
-4.75	(-No. 4)	200	470	650	860	1510	

Performance Exam Checklist FOP for AASHTO T 30 Mechanical Analysis of Extracted Aggregate

Part	ticipant Name Exam Date	Exam Date					
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.						
Pro	Procedure Element		Trial 2				
1.	Total dry mass determined to 0.1 g						
2.	Dry mass agrees with sample mass after ignition ($M_{\rm f}$) from AASHTO T 308 within 0.1 percent?						
3.	Sample placed in container and covered with water?						
4.	Wetting agent added?						
5.	Contents of container agitated vigorously?						
6.	Wash water poured through proper nest of two sieves?						
7.	Washing continued until wash water is clear and no wetting agent remaining?						
8.	Retained material returned to washed sample?						
9.	Washed material coarser than 75 μ m (No. 200) dried to constant mass at 110 ± 5°C (230 ± 9°F)?						
10.	Sample cooled to room temperature?						
11.	Dry mass after washing determined to 0.1 g?						
12.	2. Material sieved on specified sieves?						
13.	Mass of each fraction of aggregate, including minus 75 μ m (No. 200), determined and recorded to 0.1 g?						
14.	Total mass of material after sieving agrees with mass before sieving to within 0.2 percent?						
15.	Percent passing each sieve determined correctly to the nearest 0.1 percent?						
16.	Aggregate correction factor applied?						
17.	Percent passing on each sieve reported correctly to the nearest 1 percent and nearest 0.1 percent on the 75 μ m (No. 200)?						
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail						
Signa	ature of Examiner WAQTC #:						

Comments:

WSDOT Errata to FOP for AASHTO R 47

Reducing Samples of Hot Mix Asphalt (HMA) to Testing Size

WAQTC FOP for AASHTO R 47 has been adopted by WSDOT with the following changes:

Procedure

Include items below:

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.

FOP for AASHTO R 47

Reducing Samples of Hot Mix Asphalt (HMA) to Testing Size

Scope

This procedure covers sample reduction of Hot Mix Asphalt (HMA) to testing size in accordance with AASHTO R 47-14. The reduced portion is to be representative of the original sample.

Apparatus

- Thermostatically controlled oven capable of maintaining a temperature of at least 110°C (230°F) or high enough to heat the material to a pliable condition for splitting.
- Non-contact temperature measuring device.
- Metal spatulas, trowels, metal straightedges, or drywall taping knives, or a combination thereof; for removing HMA samples from the quartering device, cleaning surfaces used for splitting, etc.
- Square-tipped, flat-bottom scoop, shovel or trowel for mixing HMA prior to quartering.
- Miscellaneous equipment including hot plate, non-asbestos heat-resistant gloves or mittens, pans, buckets, and cans.
- Sheeting: Non-stick heavy paper, heat-resistant plastic, or other material as approved by the agency.
- Agency-approved release agent, free of solvent or petroleum-based material that could affect asphalt binder.
- Mechanical Splitter Type A (Quartermaster): having four equal-width chutes
 discharging into four appropriately sized sample receptacles. Splitter is to be
 equipped with a receiving hopper that will hold the sample until the release lever
 is activated with four sample receptacles of sufficient capacity to accommodate
 the reduced portion of the HMA sample from the mechanical splitter. Refer to
 AASHTO R 47, Figures 1 through 3, for configuration and required dimensions of
 the mechanical splitter.
- Mechanical Splitter Type B (Riffle): having a minimum of eight equal-width chutes
 discharging alternately to each side with a minimum chute width of at least
 50 percent larger than the largest particle size. A hopper or straight-edged pan with
 a width equal to or slightly smaller than the assembly of chutes in the riffle splitter
 to permit uniform discharge of the HMA through the chutes without segregation or
 loss of material. Sample receptacles of sufficient width and capacity to receive the
 reduced portions of HMA from the riffle splitter without loss of material.
- Quartering Template: formed in the shape of a cross with equal length sides at right
 angles to each other. Template shall be manufactured of metal that will withstand
 heat and use without deforming. The sides of the quartering template should be
 sized so that the length exceeds the diameter of the flattened cone of HMA by an
 amount allowing complete separation of the quartered sample. Height of the sides
 must exceed the thickness of the flattened cone of HMA.
- Non-stick mixing surface that is hard, heat-resistant, clean, level, and large enough to permit HMA samples to be mixed without contamination or loss of material.

Sampling

Obtain samples according to the FOP for AASHTO T 168.

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. Do not exceed either the temperature or time limits specified in the test method(s) to be performed.

Selection of Procedure (Method)

Refer to agency requirements when determining the appropriate method(s) of sample reduction. In general, the selection of a particular method to reduce a sample depends on the initial size of the sample vs. the size of the sample needed for the specific test to be performed. It is recommended that, for large amounts of material, the initial reduction be performed using a mechanical splitter. This decreases the time needed for reduction and minimizes temperature loss. Further reduction of the remaining HMA may be performed by a combination of the following methods, as approved by the agency. The methods for reduction are:

- · Mechanical Splitter Method
 - Type A (Quartermaster)
 - Type B (Riffle Splitter)
- · Quartering Method
 - Full Quartering
 - By Apex
- Incremental Method

Procedure

Mechanical Splitter Type A (Quartermaster)

- 1. Clean the splitter and apply a light coating of approved release agent to the surfaces that will contact HMA.
- 2. Close and secure hopper gates.
- 3. Place the four sample receptacles in the splitter so that there is no loss of material.
- 4. Remove the sample from the agency-approved container(s) and place in the mechanical splitter hopper. Avoid segregation, loss of HMA or the accidental addition of foreign material.
- 5. Release the handle, allowing the HMA to drop through the divider chutes and discharge into the four receptacles.
- 6. Any HMA that is retained on the surface of the splitter shall be removed and placed into the appropriate receptacle.
- 7. Close and secure the hopper gates.
- 8. Reduce the remaining HMA as needed by this method or a combination of the following methods as approved by the agency.

- 9. Combine the material contained in the receptacles from opposite corners and repeat the splitting process until an appropriate sample size is obtained.
- 10. Retain and properly identify the remaining unused portion of the HMA sample for further testing if required by the agency.

Mechanical Splitter Type B (Riffle)

- 1. When heating of the testing equipment is desired, it shall be heated to a temperature not to exceed 110°C (230°F).
- 2. Clean the splitter and apply a light coating of approved release agent to the surfaces that will come in contact with HMA (hopper or straight-edged pan, chutes, receptacles).
- 3. Place two empty receptacles under the splitter.
- 4. Carefully empty the HMA from the agency-approved container(s) into the hopper or straight-edged pan without loss of material. Uniformly distribute from side to side of the hopper or pan.
- 5. Discharge the HMA at a uniform rate, allowing it to flow freely through the chutes.
- 6. Any HMA that is retained on the surface of the splitter shall be removed and placed into the appropriate receptacle.
- 7. Reduce the remaining HMA as needed by this method or a combination of the following methods as approved by the agency.
- 8. Using one of the two receptacles containing HMA, repeat the reduction process until the HMA contained in one of the two receptacles is the appropriate size for the required test.
- 9. After each split, remember to clean the splitter hopper and chute surfaces if needed.
- 10. Retain and properly identify the remaining unused HMA sample for further testing if required by the agency.

Quartering Method

- 1. When heating of the testing equipment is desired, it shall be heated to a temperature not to exceed the maximum mixing temperature.
- 2. If needed, apply a light coating of release agent to quartering template.
- 3. Dump the sample from the agency approved container(s) into a conical pile on a hard, "non-stick," clean, level surface where there will be neither a loss of material nor the accidental addition of foreign material. The surface can be made non-stick by the application of an approved asphalt release agent, or sheeting.
- 4. Mix the material thoroughly by turning the entire sample over a minimum of four times with a flat-bottom scoop; or by alternately lifting each corner of the sheeting and pulling it over the sample diagonally toward the opposite corner, causing the material to be rolled. Create a conical pile by either depositing each scoop or shovelful of the last turning on top of the preceding one, or lifting both opposite corners.
- 5. Flatten the conical pile to a uniform diameter and thickness where the diameter is four to eight times the thickness. Make a visual observation to ensure that the material is homogeneous.

- 6. Divide the flattened cone into four equal quarters using the quartering template. Press the template down until it is in complete contact with the surface on which the sample has been placed, assuring complete separation.
 - **Note 1:** Straightedges may be used in lieu of the quartering device to completely separate the material in approximately equal quarters.
- 7. Reduce the sample by quartering the sample completely or by removing the sample from the apex.

8. Full Quartering

- 8a. Remove two diagonally opposite quarters, including all of the fine material.
- 8b. Remove the quartering template and combine the remaining quarters, again forming a conical pile.
- 8c. Repeat steps 4, 5, 6, 8a, and 8b until a sample of the required size has been obtained. The final sample must consist of the two remaining diagonally opposite quarters.
- 8d. Retain and properly identify the remaining unused portion of the HMA sample for further testing if required by the agency.

9. By Apex

- 9a. Using a straightedge, slice through a quarter of the HMA from the center point to the outer edge of the quarter.
- 9b. Pull or drag the material from the quarter with two straight edges or hold one edge of the straightedge in contact with quartering device.
- 9c. Remove an equal portion from the opposite quarter and combine these increments to create the required sample size.
- 9d. Continue using the apex method with the unused portion of the HMA until samples have been obtained for all required tests.
- 9e. Retain and properly identify the remaining unused portion of the HMA sample for further testing if required by the agency.

Incremental Method

- Cover a hard, clean, level surface with sheeting. This surface shall be large enough that there will be neither a loss of material nor the accidental addition of foreign material.
- 2. Place the sample from the agency approved container(s) into a conical pile on that surface
- 3. Mix the material thoroughly by turning the entire sample over a minimum of four times with a flat-bottom scoop; or by alternately lifting each corner of the sheeting and pulling it over the sample diagonally toward the opposite corner, causing the material to be rolled. Create a conical pile by either depositing each scoop or shovelful of the last turning on top of the preceding one, or lifting both opposite corners.
- 4. Grasp the sheeting and roll the conical pile into a cylinder (loaf), then flatten the top. Make a visual observation to determine that the material is homogenous.

- 5. Pull the sheeting so at least ¼ 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 straightedge, slice off approximately ¼ of the length of the loaf and place in a container to be saved.
- 6. Pull material off the edge of the counter and drop into an appropriate size sample pan or container for the test to be performed. Continue removing material from the loaf until the proper size sample has been acquired. As an alternate, using a straightedge, slice off an appropriate size sample from the length of the loaf and place in a sample pan or container.
- 7. Repeat step 6 until all the samples for testing have been obtained.
 - **Note 2:** 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 grossly in excess of the minimum or exceeds the maximum test size, use the sample as reduced for the test.
- 8. Retain and properly identify the remaining unused portion of the HMA sample for further testing if required by the agency.

Performance Exam Checklist

FOP for AASHTO R 47

Reducing Samples of Hot Mix Asphalt (HMA) To Testing Size

Part	Participant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Prod	edure Element	Trial 1	Trial 2
1.	Sample made soft enough to separate easily without exceeding temperature limits?		
Med	hanical Splitter Method Type A (Quartermaster)		
2.	Splitter cleaned and surfaces coated with release agent?		
3.	Hopper closed and receptacles in place?		
4.	Sample placed into hopper without segregation or loss of material?		
5.	Hopper handle released allowing the HMA to uniformly flow into receptacles?		
6.	Splitter surfaces cleaned of all retained HMA, allowing it to fall into appropriate receptacles?		
7.	Further reduction with the quartermaster:		
	a. Material in receptacles from opposite corners combined?		
	b. Splitting process repeated until appropriate sample size is obtained?		
8.	Remaining HMA stored in suitable container and properly labeled?		
Med	hanical Splitter Method Type B (Riffle)		
9.	Splitting apparatus and tools, if preheated, not exceeding 110°C (230°F)?		
10.	Splitter cleaned and surfaces coated with release agent?		
11.	Two empty receptacles placed under splitter?		
12.	Sample placed in hopper or straight edged pan without loss of material and uniformly distributed from side to side?		
13.	Material discharged across chute assembly at controlled rate allowing free flow of HMA through chutes?		
14.	Splitter surfaces cleaned of all retained HMA allowing it to fall into appropriate receptacles?		
15.	Further reduction with the riffle splitter:		
	a. Material from one receptacle discharged across chute assembly at controlled rate, allowing free flow of HMA through chutes?		
	b. Splitting process continued until appropriate sample size obtained, with splitter surfaces cleaned of all retained HMA after every split?		
16.	Remaining unused HMA stored in suitable container, properly labeled?		

Proc	edure Element	Trial 1	Trial 2
Qua	rtering Method		
17.	Testing equipment preheated to a temperature not to exceed mix temperature?		
18.	Sample placed in a conical pile on a hard, non-stick, heat-resistant splitting surface such as metal or sheeting?		
19.	Sample mixed by turning the entire sample over a minimum of 4 times?		
20.	Conical pile formed and then flattened uniformly to diameter equal to about 4 to 8 times thickness?		
21	Sample divided into 4 equal portions either with a metal quartering template or straightedges such as drywall taping knives?		
22.	Reduction by Full Quartering:		
	a. Two diagonally opposite quarters removed and returned to sample container?		
	b. Two other diagonally opposite quarters combined and process continued until appropriate sample size has been achieved?		
23.	Reduction by Apex:		
	a. Using two straightedges or a splitting device and one straightedge, was one		
	of the quarters split from apex to outer edge of material? b. Similar amount of material taken from opposite quarter?		
	c. Increments combined to produce appropriate sample size?		
24			
	Remaining unused HMA stored in suitable container, properly labeled?		
	emental Method		
25.	Sample placed on hard, non-stick, heat-resistant splitting surface covered with sheeting?		
26.	Sample mixed by turning the entire sample over a minimum of 4 times?		
27.	Conical pile formed?		
28.	HMA rolled into loaf and then flattened?		
29.	The first quarter of the loaf removed by slicing off or dropping off edge of counter and set aside?		
30.	Proper sample size sliced off or dropped off edge of counter into sample container?		
31.	Process continued until all samples are obtained?		
32.	All remaining unused HMA stored in suitable container, properly labeled?		
First	: Attempt: Pass Fail Second Attempt: Pass Fail		
	•		
Signa	ture of Examiner WAQTC #:		

Comments

FOP for AASHTO R 66

Sampling Asphalt Materials

Scope

This procedure covers obtaining samples of liquid asphalt materials in accordance with AASHTO R 66-16. Sampling of solid and semi-solid asphalt materials – included in AASHTO R 66 – is not covered here.

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

Warning: Always use appropriate safety equipment and precautions for hot liquids.

Terminology

- Asphalt binder: Asphalt cement or modified asphalt cement that binds the aggregate particles into a dense mass.
- Asphalt emulsion: A mixture of asphalt binder and water.
- Cutback asphalt: Asphalt binder that has been modified by blending with a chemical solvent.

Procedure

- 1. Coordinate sampling with contractor or supplier.
- 2. Allow a minimum of 4 L (1 gal) to flow before obtaining a sample(s).
- 3. Obtain samples of:
 - Asphalt binder from the line between the storage tank and the mixing plant 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 L (1 qt) 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.

- Emulsified asphalt: Use wide-mouth plastic jars with screw caps. Protect the samples from freezing since water is a part of the emulsion. The sample container should be completely filled to minimize a skin formation on the sample.
- Asphalt binder and cutbacks: Use metal cans.

Note: The 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.

Report

- On forms approved by the agency
- Sample ID
- Date
- Time
- Location
- Quantity represented

FOP for AASHTO R 75

Developing a Family of Curves

Scope

This procedure provides a method to develop a family of curves in accordance with AASHTO R 75-16 using multiple moisture density relationships developed using the same method, A, B, C, or D, from the FOP for AASHTO T 99/T 180.

All curves used in a family must be developed using a single Method: A, B, C, or D of a procedure for AASHTO T 99 or T 180. See the FOP for AASHTO T 99/T 180.

Terminology

family of curves — a group of soil moisture-density relationships (curves) determined using AASHTO T 99 or T 180, which reveal certain similarities and trends characteristic of the soil type and source.

spine — smooth line extending through the point of maximum density/optimum moisture content of a family of moisture-density curves.

Procedure

- 1. Sort the curves by Method (A, B, C, or D of the FOP for T 99/T 180). At least three curves are required to develop a family.
- 2. Select the highest and lowest maximum dry densities from those selected to assist in determining the desired scale of the subsequent graph.
- 3. Plot the maximum density and optimum moisture points of the selected curves on the graph.
- 4. Draw a smooth, "best fit," curved line through the points creating the spine of the family of curves.
- 5. Remove maximum density and optimum moisture points that were not used to establish the spine.
- 6. Add the moisture/density curves associated with the points that were used to establish the spine. It is not necessary to include the portion of the curves over optimum moisture.
 - **Note 1:** Intermediate template curves using slopes similar to those of the original moisture-density curves may be included when maximum density points are more than 2.0 lb/ft³ apart. Template curves are indicated by a dashed line.
- 7. Plot the 80 percent of optimum moisture range when desired:
 - a. Using the optimum moisture of an existing curve, calculate 80 percent of optimum moisture and plot this value on the curve. Repeat for each curve in the family.
 - b. Draw a smooth, "best fit," curved line connecting the 80 percent of optimum moisture points plotted on the curves that parallel the spine.

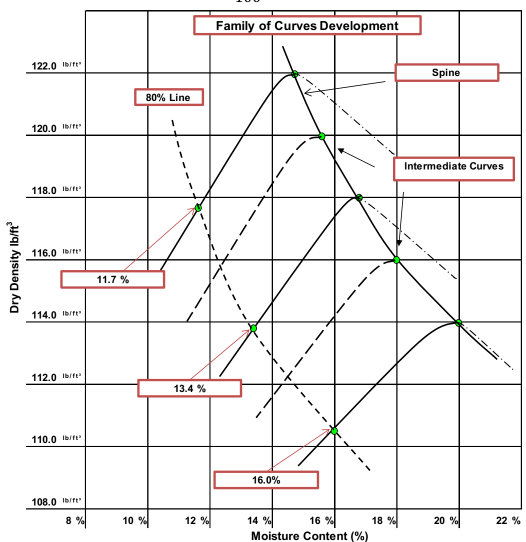
Calculations

Calculate 80 percent of optimum moisture of each curve:

Example:

Optimum moisture of the highest density curve = 14.6 percent

$$80\% \ point = \frac{80}{100} \times 14.6\% = 11.7\%$$



Performance Exam Checklist FOP for AASHTO R 75 Developing a Family of Curves

Par	Participant Name Exam Date					
Red	cord the symbols "P" for passing or "F" for failing on each step of the checklist.					
Pro	cedure Element	Trial 1	Trial 2			
1.	Curves sorted by method and procedure (A, B, C, or D of the FOP for T 99/T 180)?					
	a. At least three curves per family?					
2.	Maximum density and optimum moisture points plotted on the graph?					
3.	Spine drawn correctly?					
4.	Maximum density and optimum moisture points removed that were not used for the spine?					
5.	Moisture/density curves added?					
6.	Optimum moisture range?					
	a. 80 percent of optimum moisture calculated for each curve?					
	b. Curved line through 80 percent of optimum moisture drawn correctly?					
Firs	st Attempt: Pass Fail Second Attempt: Pass Fail					
Sign	ature of Examiner WAQTC #:					
Con	nments:					

FOP FOR AASHTO R 76

Reducing Samples of Aggregates to Testing Size

Scope

This procedure covers the reduction of samples to the appropriate size for testing in accordance with AASHTO R 76-16. Techniques are used that minimize variations in characteristics between test samples and field samples. Method A (Mechanical Splitter) and Method B (Quartering) are covered.

This FOP applies to fine aggregate (FA), coarse aggregate (CA), and mixes of the two (FA/CA), and may also be used on soils.

Apparatus

Method A - Mechanical Splitter

Splitter chutes:

- Even number of equal width chutes
- Discharge alternately to each side
- Minimum of 8 chutes total for CA and FA/CA, 12 chutes total for FA
- · Width:
 - Minimum 50 percent larger than largest particle
 - Maximum chute width of 19 mm (¾ in) for fine aggregate passing the 9.5 mm
 (¾ in) sieve

Feed control:

- Hopper or straightedge pan with a width equal to or slightly less than the overall width of the assembly of chutes
- Capable of feeding the splitter at a controlled rate

Splitter receptacles/pans:

Capable of holding two halves of the sample following splitting

The splitter and accessory equipment shall be so designed that the sample will flow smoothly without restriction or loss of material.

Method B - Quartering

- Straightedge scoop, shovel, or trowel
- · Broom or brush
- Canvas or plastic sheet, approximately 2 by 3 m (6 by 9 ft)

Method Selection

Samples of CA may be reduced by either Method A or Method B.

Samples of FA which are drier than the saturated surface dry (SSD) condition, as described in AASHTO T 84, shall be reduced by a mechanical splitter according to Method A. As a quick approximation, if the fine aggregate will retain its shape when molded with the hand, it is wetter than SSD.

Samples of FA/CA which are drier than SSD may be reduced by Method A or Method B.

Samples of FA and FA/CA that are at SSD or wetter than SSD shall be reduced by Method B, or the entire sample may be dried to the SSD condition – using temperatures that do not exceed those specified for any of the tests contemplated – and then reduced to test sample size using Method A.

Table 1

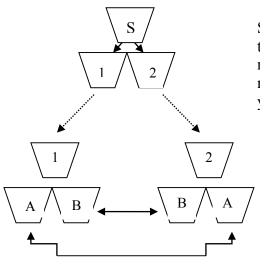
	Drier than SSD	Wetter than SSD
Fine Aggregate (FA)	Method A (Mechanical)	Method B (Quartering)
Mixture of FA/CA	Either Method	Method B (Quartering)
Coarse Aggregate (CA)	Either Method	Either Method

Procedure

Method A - Mechanical Splitter

- 1. Place the sample in the hopper or pan and uniformly distribute it from edge to edge so that approximately equal amounts 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 pans below.
- 2. Reduce the sample from one of the two pans as many times as necessary to reduce the sample to meet the minimum size specified for the intended test. The portion of the material collected in the other pan may be reserved for reduction in size for other tests.
- 3. As a check for effective reduction, determine the mass of each reduced portion. If the percent difference of the two masses is greater than 5 percent, corrective action must be taken. In lieu of the check for effective reduction, use the method illustrated in Figure 1.

Figure 1



Sample (S) is an amount greater than or equal to twice the mass needed for testing. Sample (S) is reduced in a mechanical splitter to yield parts (1) and (2)

Part (1) is further reduced yielding (A) and (B) while part (2) is reduced to yield (B) and (A).

Final testing sample is produced by combining alternate pans, i.e. A/A or B/B only.

Calculation

$$\frac{Smaller\ Mass}{Larger\ Mass} = Ratio \quad (1 - ratio) \times 100 = \%\ Difference$$

Splitter check: 5127 g total sample mass

Splitter pan #1: 2583 g Splitter pan #2: 2544 g

$$\frac{2544 \text{ g}}{2583 \text{ g}} = 0.985 \qquad (1 - 0.985) \times 100 = 1.5\%$$

Procedure

Method B - Quartering

Use either of the following two procedures or a combination of both.

Procedure # 1: Quartering on a clean, hard, level surface:

- 1. Place the sample on a hard, clean, level surface where there will be neither loss of material nor the accidental addition of foreign material.
- 2. Mix the material thoroughly by turning the entire sample over a minimum of four times. With the last turning, shovel the entire sample into a conical pile by depositing each shovelful on top of the preceding one.
- 3. Flatten the conical pile to a uniform thickness and diameter by pressing down with a shovel. The diameter should be four to eight times the thickness.
- 4. Divide the flattened pile into four approximately equal quarters with a shovel or trowel.
- 5. Remove two diagonally opposite quarters, including all fine material, and brush the cleared spaces clean.
- 6. Successively mix and quarter the remaining material until the sample is reduced to the desired size.
- 7. The final test sample consists of two diagonally opposite quarters.

Procedure # 2: Quartering on a canvas or plastic sheet:

- 1. Place the sample on the sheet.
- 2. Mix the material thoroughly a minimum of four times by pulling each corner of the sheet horizontally over the sample toward the opposite corner. After the last turn, form a conical pile.
- 3. Flatten the conical pile to a uniform thickness and diameter by pressing down with a shovel. The diameter should be four to eight times the thickness.
- 4. Divide the flattened pile into four approximately equal quarters with a shovel or trowel, or, insert a stick or pipe beneath the sheet and under the center of the pile, then lift both ends of the stick, dividing the sample into two roughly equal parts. Remove the stick leaving a fold of the sheet 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 roughly equal quarters.
- 5. Remove two diagonally opposite quarters, being careful to clean the fines from the sheet.
- 6. Successively mix and quarter the remaining material until the sample size is reduced to the desired size.
- 7. The final test sample consists of two diagonally opposite quarters.

Performance Exam Checklist FOP for AASHTO R 76 Reducing Field Samples of Aggregates to Testing Size

Par	Participant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
Me	thod 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?		
Me	thod B - Quartering		
1.	Sample placed on clean, hard, and level surface?		
2.	Mixed by turning over 4 times with shovel or by pulling sheet horizontally 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?		
8.	Cleared space between quarters brushed clean?		
9.	Process continued until desired sample size is obtained when two opposite quarters combined?		
	The sample may be placed upon a sheet and a stick or pipe may be placed under the sheet to divide the pile into quarters.		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	ature of Examiner WAQTC #:		
Com	nments:		

FOP for AASHTO T 85

Specific Gravity and Absorption of Coarse Aggregate

Scope

This procedure covers the determination of specific gravity and absorption of coarse aggregate in accordance with AASHTO T 85-14. Specific gravity may be expressed as bulk specific gravity ($G_{\rm sb}$), bulk specific gravity, saturated surface dry ($G_{\rm sb}$ SSD), or apparent specific gravity ($G_{\rm sa}$). $G_{\rm sb}$ and absorption are based on aggregate after soaking in water. This procedure is not intended to be used with lightweight aggregates.

Terminology

Absorption – the increase in the mass of aggregate due to water being absorbed into the pores of the material, but not including water adhering to the outside surface of the particles, expressed as a percentage of the dry mass. The aggregate is considered "dry" when it has been maintained at a temperature of 110 ± 5 °C (230 ± 9 °F) for sufficient time to remove all uncombined water.

Saturated Surface Dry (SSD) – condition of an aggregate particle when the permeable voids are filled with water, but no water is present on exposed surfaces.

Specific Gravity – the ratio of the mass, in air, of a volume of a material to the mass of the same volume of gas-free distilled water at a stated temperature.

Apparent Specific Gravity (G_{sa}) – the ratio of the mass, in air, of a volume of the impermeable portion of aggregate to the mass of an equal volume of gas-free distilled water at a stated temperature.

Bulk Specific Gravity (G_{sb})– the ratio of the mass, in air, of a volume of aggregate (including the permeable and impermeable voids in the particles, but not including the voids between particles) to the mass of an equal volume of gas-free distilled water at a stated temperature.

Bulk Specific Gravity (SSD) (G_{sb} SSD) – the ratio of the mass, in air, of a volume of aggregate, including the mass of water within the voids filled to the extent achieved by submerging in water for 15 to 19 hours (but not including the voids between particles), to the mass of an equal volume of gas-free distilled water at a stated temperature.

Apparatus

- Balance or scale: with a capacity of 5 kg, sensitive to 1 g. Meeting the requirements of AASHTO M 231.
- Sample container: a wire basket of 3.35 mm (No. 6) or smaller mesh, with a capacity of 4 to 7 L (1 to 2 gal) to contain aggregate with a nominal maximum size of 37.5 mm (1½ in) or smaller; or a larger basket for larger aggregates, or both.
- Water tank: watertight and large enough to completely immerse aggregate and basket, equipped with an overflow valve to keep water level constant.
- Suspension apparatus: wire used to suspend apparatus shall be of the smallest practical diameter.
- Sieves 4.75 mm (No. 4) or other sizes as needed, meeting the requirements of FOP for AASHTO T 27/T 11.
- · Large absorbent towel

Sample Preparation

- 1. Obtain the sample in accordance with the FOP for AASHTO T 2 (see Note 1).
- 2. Mix the sample thoroughly and reduce it to the approximate sample size required by Table 1 in accordance with the FOP for AASHTO R 76.
- 3. Reject all material passing the appropriate sieve by dry sieving.
- 4. Thoroughly wash sample to remove dust or other coatings from the surface.
- 5. Dry the test sample to constant mass at a temperature of $110 \pm 5^{\circ}$ C ($230 \pm 9^{\circ}$ F) and cool in air at room temperature for 1 to 3 hours.
 - **Note 1:** Where the absorption and specific gravity values are to be used in proportioning concrete mixtures in which the aggregates will be in their naturally moist condition, the requirement for initial drying to constant mass may be eliminated, and, if the surfaces of the particles in the sample have been kept continuously wet until test, the 15-to-19 hour soaking may also be eliminated.
- 6. Re-screen the sample over the appropriate sieve. Reject all material passing that sieve.
- 7. The sample shall meet or exceed the minimum mass given in Table 1.

Note 2: If this procedure is used only to determine the G_{sb} of oversized material for the FOP for AASHTO T 99/T 180, the material can be rejected over the appropriate sieve. For T 99/T 180 Methods A and B, use the 4.75 mm (No. 4) sieve; T 99/T 180 Methods C and D use the 19 mm ($\frac{3}{4}$ in).

Table 1

Nominal Maxir	num Size* mm (in)	Minimum Mass of	Test Sample, g (lb)
12.5	(½) or less	2000	(4.4)
19.0	(3/4)	3000	(6.6)
25.0	(1)	4000	(8.8)
37.5	(1½)	5000	(11)
50	(2)	8000	(18)
63	(2½)	12,000	(26)
75	(3)	18,000	(40)

^{*} One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size.

Procedure

- 1. Immerse the aggregate in water at room temperature for a period of 15 to 19 hours.
 - **Note 3:** When testing coarse aggregate of large nominal maximum size requiring large test samples, it may be more convenient to perform the test on two or more subsamples, and then combine the values obtained.
- 2. Place the empty basket into the water bath and attach to the balance. Inspect the immersion tank to ensure the water level is at the overflow outlet height. Tare the balance with the empty basket attached in the water bath.
- 3. Remove the test sample from the water and roll it in a large absorbent cloth until all visible films of water are removed. Wipe the larger particles individually. If the test sample dries past the SSD condition, immerse in water for 30 min, and then resume the process of surface-drying.
 - **Note 4:** A moving stream of air may be used to assist in the drying operation, but take care to avoid evaporation of water from aggregate pores.
- 4. Determine the SSD mass of the sample, and record this and all subsequent masses to the nearest 0.1 g or 0.1 percent of the sample mass, whichever is greater. Designate this mass as "B."
- 5. Immediately place the SSD test sample in the sample container and weigh it in water maintained at $23.0 \pm 1.7^{\circ}$ C ($73.4 \pm 3^{\circ}$ F). Shake the container to release entrapped air before recording the weight. Re-inspect the immersion tank to insure the water level is at the overflow outlet height. Designate this submerged weight as "C."
 - **Note 5:** The container should be immersed to a depth sufficient to cover it and the test sample during mass determination. Wire suspending the container should be of the smallest practical size to minimize any possible effects of a variable immersed length.
- 6. Remove the sample from the basket. Ensure all material has been removed. Place in a container of known mass.
- 7. Dry the test sample to constant mass in accordance with the FOP for AASHTO T 255/T 265 (Aggregate section) and cool in air at room temperature for 1 to 3 hours. Designate this mass as "A."

Calculations

Perform calculations and determine values using the appropriate formula below.

Bulk specific gravity (G_{sh})

$$G_{sb} = \frac{A}{B - C}$$

Bulk specific gravity, SSD (G_{sb} SSD)

$$G_{sb}SSD = \frac{B}{B - C}$$

Apparent specific gravity (G_{sa})

$$G_{sa} = \frac{A}{A - C}$$

Absorption

Absorption =
$$\frac{B-A}{A} \times 100$$

Where:

A = oven dry mass, g B = SSD mass, g C = weight in water, g

Sample Calculations

Sample	Α	В	С	B - C	A - C	B - A
1	2030.9	2044.9	1304.3	740.6	726.6	14.0
2	1820.0	1832.5	1168.1	664.4	651.9	12.5
3	2035.2	2049.4	1303.9	745.5	731.3	14.2

Sample	G _{sb}	G _{sb} SSD	G _{sa}	Absorption
1	2.742	2.761	2.795	0.7
2	2.739	2.758	2.792	0.7
3	2.730	2.749	2.783	0.7

These calculations demonstrate the relationship between G_{sb} , G_{sb} SSD, and G_{sa} . G_{sb} is always lowest, since the volume includes voids permeable to water. G_{sb} SSD is always intermediate. G_{sa} is always highest, since the volume does not include voids permeable to water. When running this test, check to make sure the values calculated make sense in relation to one another.

Report

- · Results on forms approved by the agency
- Sample ID
- Specific gravity values to 3 decimal places
- Absorption to 0.1 percent

Performance Exam Checklist FOP for AASHTO T 85 Specific Gravity and Absorption of Coarse Aggregate

Part	Participant Name Exam Date		
Reco	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Prod	cedure Element	Trial 1	Trial 2
1.	Sample obtained by FOP for AASHTO T 2 and reduced by FOP for AASHTO R 76 or from FOP for AASHTO T 99/T 180?		
2.	Screened on the appropriate size sieve?		
3.	Sample mass appropriate?		
4.	Particle surfaces clean?		
5.	Dried to constant mass 110 ± 5 °C (230 ± 9 °F) and cooled to room temperature?		
6.	Re-screen over appropriate sieve?		
7.	Covered with water for 15 to 19 hours?		
8.	Basket placed into immersion tank and attached to balance?		
9.	Immersion tank inspected for proper water height?		
10.	Balance tared with basket in tank and temperature checked 23.0 ± 1.7 °C (73.4 ± 3°F)?		
11.	Sample removed from water and rolled in cloth to remove visible films of water?		
12.	Larger particles wiped individually?		
13.	Evaporation avoided?		
14.	Sample mass determined to 0.1 g?		
15.	Sample immediately placed in basket, in immersion tank?		
16.	Entrapped air removed before weighing by shaking basket while immersed?		
17.	Immersion tank inspected for proper water height?		
18.	Immersed sample weight determined to 0.1 g?		
19.	All the sample removed from basket?		
20.	Sample dried to constant mass and cooled to room temperature?		
21.	Sample mass determined to 0.1 g?		
22.	Proper formulas used in calculations?		
First	Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		

Comments:

WSDOT FOP for AASHTO T 891 **Determining the Liquid Limit of Soils** WSDOT has adopted AASHTO T 89.

¹This FOP is based on AASHTO T 23-08.

Performance Exam Checklist

Determining the Liquid Limit of Soils AASHTO T 89 (Method B Only)

Part	ticipant Name Exam Date				
Pre	paration	Yes	Νo		
1.	The tester has a copy of the current procedure on hand?	103	110		
2.	All equipment is functioning according to the test procedure, and if required, has the current calibration/verification tags present?				
3.	Sample obtained using AASHTO R 58?				
4.	Minimum sample mass meets requirement of AASHTO T 89 Method B?				
5.	Sample mixed with 15 to 20 mL of distilled or demineralized water?				
6.	Additional water added at 1 to 3 mL as necessary until mass is uniform and of a stiff consistency?				
7.	No dry soil added after test has begun?				
8.	If soil was too wet, was sample discarded or allowed to dry?				
Pro	ocedure	Yes	No		
1.	Sample placed in cup and spread to 10 mm maximum thickness?				
2.	Care taken to avoid entrapment of air bubbles?				
3.	Soil in cup divided through centerline of follower to the bottom of the cup in no more than six strokes?				
4.	Liquid Limit Device counter zeroed and base checked for level?				
5.	Was cup lifted and dropped at two revolutions per second until gap at bottom of groove closed about 0.5 in (13mm) in 22 to 28 blows?				
6.	Blows to closure recorded?				
7.	Was closure in acceptable blow count material?				
8.	Was material removed from cup and placed in a covered container?				
9.	Was procedure repeated a second time from step 1-6 without adding water?				
10	Was second closure within two blows of first closure? If not was test rerun?				
11.	. Was sample removed from device and moisture content determined per T 265?				
12.	Were all calculations performed correctly?				
Firs	st Attempt: Pass Fail Second Attempt: Pass Fail				
Sig	nature of Examiner				

Comments:

WSDOT FOP for AASHTO T 90

Determining the Plastic Limit and Plasticity Index of Soils

WSDOT has adopted AASHTO T 90.

Performance Exam Checklist

AASHTO T 90

Determining the Plastic Limit and Plasticity Index of Soils

Part	icipant Name _			Exam Date		
Pre	paration				Yes	No
1.	The tester has	а сору	of the curr	rent procedure on hand?		
2.			_	cording to the test procedure, and if required, cation tags present?		
3.	Sample obtain	ed usin	g AASHT(O R 58?		
4.	Minimum sam	ple ma	ss meets re	equirement of AASHTO T 90?		
5.	Sample mixed ball allowed to			demineralized water until plastic enough to form undisturbed?		
6.	8 g ball formed	d from	the moist s	sample material?		
7.	Ball broken in	to 1.5-2	g portions	s and formed into ellipsoidal masses?		
Pro	cedure				Yes	No
1.				rokes per minute (using one of the techniques than 2 minutes to form a 3 mm diameter thread?		
2.				pieces and pieces squeezed together into ellipsoidal crumbles under and soil can no longer be rolled into		
3.	Tested materia until all 8 g of	-		d covered container and procedure steps 1-6 repeated?		
4.	Sample dried i	in accor	dance with	h T 265 to determine moisture content?		
5.	Were all calcul	lations	performed	correctly?		
Firs	t Attempt: Pas	SS	Fail	Second Attempt: Pass Fail		
Sign	nature of Exami	iner				

Comments:

MOISTURE-DENSITY RELATIONS OF SOILS:

FOP for AASHTO T 99
USING A 2.5 kg (5.5 lb) RAMMER AND A 305 mm (12 in) DROP

FOP for AASHTO T 180
USING A 4.54 kg (10 lb) RAMMER AND A 457 mm (18 in) DROP

Scope

This procedure covers the determination of the moisture-density relations of soils and soil-aggregate mixtures in accordance with two similar test methods:

- AASHTO T 99-17: Methods A, B, C, and D
- AASHTO T 180-17: Methods A, B, C, and D

This test method applies to soil mixtures having 40 percent or less retained on the 4.75 mm (No. 4) sieve for methods A or B, or, 30 percent or less retained on the 19 mm (¾ in) with methods C or D. The retained material is defined as oversize (coarse) material. If no minimum percentage is specified, 5 percent will be used. Samples that contain oversize (coarse) material that meet percent retained criteria should be corrected by using Annex A, Correction of Maximum Dry Density and Optimum Moisture for Oversized Particles. Samples of soil or soil-aggregate mixture are prepared at several moisture contents and compacted into molds of specified size, using manual or mechanical rammers that deliver a specified quantity of compactive energy. The moist masses of the compacted samples are multiplied by the appropriate factor to determine wet density values. Moisture contents of the compacted samples are determined and used to obtain the dry density values of the same samples. Maximum dry density and optimum moisture content for the soil or soil-aggregate mixture is determined by plotting the relationship between dry density and moisture content.

Apparatus

- Mold Cylindrical mold made of metal with the dimensions shown in Table 1 or Table 2. If permitted by the agency, the mold may be of the "split" type, consisting of two half-round sections, which can be securely locked in place to form a cylinder. Determine the mold volume according to *Annex B*, *Standardization of the Mold*.
- Mold assembly Mold, base plate, and a detachable collar.
- Rammer –Manually or mechanically-operated rammers as detailed in Table 1 or Table 2. A manually-operated rammer shall be equipped with a guide sleeve to control the path and height of drop. The guide sleeve shall have at least four vent holes no smaller than 9.5 mm (¾ in) in diameter, spaced approximately 90 degrees apart and approximately 19 mm (¾ in) from each end. A mechanically-operated rammer will uniformly distribute blows over the sample and will be calibrated with several soil types, and be adjusted, if necessary, to give the same moisture-density results as with the manually operated rammer. For additional information concerning calibration, see the FOP for AASHTO T 99 and T 180.
- Sample extruder A jack, lever frame, or other device for extruding compacted specimens from the mold quickly and with little disturbance.

• Balance(s) or scale(s) of the capacity and sensitivity required for the procedure used by the agency.

A balance or scale with a capacity of 11.5 kg (25 lb) and a sensitivity of 1 g for obtaining the sample, meeting the requirements of AASHTO M 231, Class G 5.

A balance or scale with a capacity of 2 kg and a sensitivity of 0.1 g is used for moisture content determinations done under both procedures, meeting the requirements of AASHTO M 231, Class G 2.

- Drying apparatus A thermostatically controlled drying oven, capable of maintaining a temperature of 110 ± 5°C (230 ± 9°F) for drying moisture content samples in accordance with the FOP for AASHTO T 255/T 265.
- Straightedge A steel straightedge at least 250 mm (10 in) long, with one beveled edge and at least one surface plane within 0.1 percent of its length, used for final trimming.
- Sieve(s) 4.75 mm (No. 4) and/or 19.0 mm (¾ in), meeting the requirements of FOP for AASHTO T 27/T 11.
- Mixing tools Miscellaneous tools such as a mixing pan, spoon, trowel, spatula, etc., or a suitable mechanical device, for mixing the sample with water.
- Containers with close-fitting lids to prevent gain or loss of moisture in the sample.

Table 1 Comparison of Apparatus, Sample, and Procedure – Metric

	T 99	T 180	
Mold Volume, m ³	Methods A, C: 0.000943 ± 0.000014	Methods A, C: 0.000943 ± 0.000014	
	Methods B, D: 0.002124 ± 0.000025	Methods B, D: 0.002124 ± 0.000025	
Mold Diameter, mm	Methods A, C: 101.60 ± 0.40	Methods A, C: 101.60 ± 0.4	
	Methods B, D: 152.40 ± 0.70	Methods B, D: 152.40 ± 0.70	
Mold Height, mm	116.40 ± 0.50	116.40 ± 0.50	
Detachable Collar Height, mm	50.80 ± 0.64	50.80 ± 0.64	
Rammer Diameter, mm	50.80 ± 0.25	50.80 ± 0.25	
Rammer Mass, kg	2.495 ± 0.009	4.536 ± 0.009	
Rammer Drop, mm	305	457	
Layers	3	5	
Blows per Layer	Methods A, C: 25	Methods A, C: 25	
	Methods B, D: 56	Methods B, D: 56	
Material Size, mm	Methods A, B: 4.75 minus	Methods A, B: 4.75 minus	
	Methods C, D: 19.0 minus	Methods C, D: 19.0 minus	
Test Sample Size, kg	Method A: 3 Method B: 7		
	Method C: 5 (1) Method D: 11	(1)	
Energy, kN-m/m ³	592	2,693	

⁽¹⁾ This may not be a large enough sample depending on your nominal maximum size for moisture content samples.

	Т 99	T 180
Mold Volume, ft ³	Methods A, C: 0.0333 ± 0.0005	Methods A, C: 0.0333 ± 0.0005
	Methods B, D: 0.07500 ± 0.0009	Methods B, D: 0.07500 ± 0.0009
Mold Diameter, in	Methods A, C: 4.000 ± 0.016	Methods A, C: 4.000 ± 0.016
	Methods B, D: 6.000 ± 0.026	Methods B, D: 6.000 ± 0.026
Mold Height, in	4.584 ± 0.018	4.584 ± 0.018
Detachable Collar Height, in	2.000 ± 0.025	2.000 ± 0.025
Rammer Diameter, in	2.000 ± 0.025	2.000 ± 0.025
Rammer Mass, Ib	5.5 ± 0.02	10 ± 0.02
Rammer Drop, in	12	18
Layers	3	5
Blows per Layer	Methods A, C: 25	Methods A, C: 25
	Methods B, D: 56	Methods B, D: 56
Material Size, in	Methods A, B: No. 4 minus	Methods A, B: No.4 minus
	Methods C, D: ¾ minus	Methods C, D: ¾ minus
Test Sample Size, lb	Method A: 7 Method B: 16 Method C: 12 ₍₁₎ Method D: 25 ₍₁₎	
Energy, lb-ft/ft ³	12,375	56,250

Table 2Comparison of Apparatus, Sample, and Procedure – English

Sample

If the sample is damp, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus maintained at a temperature not exceeding 60°C (140°F). Thoroughly break up aggregations in a manner that avoids reducing the natural size of individual particles.

Obtain a representative test sample of the mass required by the agency by passing the material through the sieve required by the agency. See Table 1 or Table 2 for test sample mass and material size requirements.

In instances where the material is prone to degradation, i.e., granular material, a compaction sample with differing moisture contents should be prepared for each point.

Note 1: Both T 99 and T 180 have four methods (A, B, C, D) that require different masses and employ different sieves.

Note 2: If the sample is plastic (clay types), it should stand for a minimum of 12 hours after the addition of water to allow the moisture to be absorbed. In this case, several samples at different moisture contents should be prepared, put in sealed containers and tested the next day.

⁽¹⁾ This may not be a large enough sample depending on your nominal maximum size for moisture content samples.

Procedure

During compaction, rest the mold firmly on a dense, uniform, rigid, and stable foundation or base. This base shall remain stationary during the compaction process.

- 1. Determine the mass of the clean, dry mold. Include the base plate, but exclude the extension collar. Record the mass to the nearest 1 g (0.005 lb).
- 2. Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately 4 to 8 percentage points below optimum moisture content. See Note 2. For many materials, this condition can be identified by forming a cast by hand.
- 3. Form a specimen by compacting the prepared soil in the mold assembly in approximately equal layers. For each layer:
 - a. Spread the loose material uniformly in the mold.
 - **Note 3:** It is recommended to cover the remaining material with a non-absorbent sheet or damp cloth to minimize loss of moisture.
 - b. Lightly tamp the loose material with the manual rammer or other similar device, this establishes a firm surface.
 - c. Compact each layer with uniformly distributed blows from the rammer. See Table 1 for mold size, number of layers, number of blows, and rammer specification for the various test methods. Use the method specified by the agency.
 - d. Trim down material that has not been compacted and remains adjacent to the walls of the mold and extends above the compacted surface.
- 4. Remove the extension collar. Avoid shearing off the sample below the top of the mold. The material compacted in the mold should not be over 6 mm (¼ in) above the top of the mold once the collar has been removed.
- 5. Trim the compacted soil even with the top of the mold with the beveled side of the straightedge.
- 6. Determine and record the mass of the mold, base plate, and wet soil to the nearest 1 g (0.005 lb) or better.
- 7. Determine and record the wet mass (M_w) of the sample by subtracting the mass in Step 1 from the mass in Step 6.
- 8. Calculate the wet density, in kg/m³ (lb/ft³), by dividing the wet mass by the measured volume (V_m).
- 9. Extrude the material from the mold. For soils and soil-aggregate mixtures, slice vertically through the center and take a representative moisture content sample from one of the cut faces, ensuring that all layers are represented. For granular materials, a vertical face will not exist. Take a representative sample. This sample must meet the sample size requirements of the test method used to determine moisture content.

Note 4: When developing a 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 from the mixing bowl may be preferred in order to determine the amount of moisture available for compaction.

- 10. Determine and record the moisture content of the sample in accordance with the FOP for AASHTO T 255/T 265.
- 11. Thoroughly break up the remaining portion of the molded specimen until it will again pass through the sieve, as judged by eye, and add to the remaining portion of the sample being tested. See Note 2.
- 12. Add sufficient water to increase the moisture content of the remaining soil by 1 to 2 percentage points and repeat steps 3 through 11.
- 13. Continue determinations until there is either a decrease or no change in the wet mass. There will be a minimum of three points on the dry side of the curve and two points on the wet side. For non-cohesive, drainable soils, one point on the wet side is sufficient.

Calculations

Wet Density

$$D_w = \frac{M_w}{V_m}$$

Where:

 $D_w = \text{wet density, kg/m}^3 \text{ (lb/ft}^3)$

 M_w = wet mass

 $V_{\rm m}$ = volume of the mold, Annex B

Dry Density

$$D_d = \left(\frac{D_w}{w + 100}\right) \times 100 \quad or \quad D_d = \frac{D_w}{\left(\frac{W}{100}\right) + 1}$$

Where:

D_d = dry density, kg/m³ (lb/ft³) w = moisture content, as a percentage

Example for 4-inch mold, Methods A or C

Wet mass, $M_{w} = 1.944 \text{ kg} (4.25 \text{ lb})$

Moisture content, w = 11.3 percent

Measured volume of the mold, $V_m = 0.000946 \text{ m}^3 (0.0334 \text{ ft}^3)$

Wet Density

$$D_w = \frac{1.944 \, kg}{0.000946 \, m^3} = 2055 \, kg/m^3 \quad D_w = \frac{4.25 \, lb}{0.0334 \, ft^3} = 127.2 \, lb/ft^3$$

Dry Density

$$D_d = \left(\frac{2055 \, kg/m^3}{11.3 + 100}\right) \times 100 = 1846 \, kg/m^3 \ D_d = \left(\frac{127.2 \, lb/ft^3}{11.3 + 100}\right) \times 100 = 114.3 \, lb/ft^3$$

Or

$$D_d = \left(\frac{2055 \, kg/m^3}{\frac{11.3}{100} + 1}\right) = 1846 \, kg/m^3 \quad D_d = \left(\frac{127.2 \, lb/ft^3}{\frac{11.3}{100} + 1}\right) = 114.3 \, lb/ft^3$$

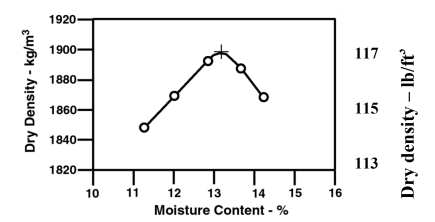
Moisture-Density Curve Development

When dry density is plotted on the vertical axis versus moisture content on the horizontal axis and the points are connected with a smooth line, a moisture-density curve is developed. The coordinates of the peak of the curve are the maximum dry density, or just "maximum density," and the "optimum moisture content" of the soil.

Example

Given the following dry density and corresponding moisture content values develop a moisture-density relations curve and determine maximum dry density and optimum moisture content.

Dry D	Moisture	
kg/m ³	lb/ft ³	Content, %
1846	114.3	11.3
1868	115.7	12.1
1887	116.9	12.8
1884	116.7	13.6
1871	115.9	14.2



In this case, the curve has its peak at:

Maximum dry density = $1890 \text{ kg/m}^3 (117.0 \text{ lb/ft}^3)$

Optimum moisture content = 13.2 percent

Note that both values are approximate, since they are based on sketching the curve to fit the points.

Report

- Results on forms approved by the agency
- Sample ID
- Maximum dry density to the closest 1 kg/m³ (0.1 lb/ft³)
- Optimum moisture content to the closest 0.1 percent

ANNEX A

Correction of Maximum DRY Density and Optimum Moisture for Oversized Particles

This section corrects the maximum dry density and moisture content of the material retained on the 4.75 mm (No. 4) sieve, Methods A and B; or the material retained on the 19 mm (¾ in) sieve, Methods C and D. The maximum dry density, corrected for oversized particles and total moisture content, are compared with the field-dry density and field moisture content.

This correction can be applied to the sample on which the maximum dry density is performed. A correction may not be practical for soils with only a small percentage of oversize material. The agency shall specify a minimum percentage below which the method is not needed. If not specified, this method applies when more than 5 percent by weight of oversize particles is present.

Bulk specific gravity (G_{sb}) of the oversized particles is required to determine the corrected maximum dry density. Use the bulk specific gravity as determined using the FOP for AASHTO T 85 in the calculations. For construction activities, an agency established value or specific gravity of 2.600 may be used.

This correction can also be applied to the sample obtained from the field while performing in-place density.

- Use the sample from this procedure or a sample obtained according to the FOP for AASHTO T 310.
- 2. Sieve the sample on the 4.75 mm (No. 4) sieve for Methods A and B or the 19 mm (¾ in) sieve, Methods C and D.
- 3. Determine the dry mass of the oversized and fine fractions (M_{DC} and M_{DF}) by one of the following:
 - a. Dry the fractions, fine and oversized, in air or by use of a drying apparatus that is maintained at a temperature not exceeding 60°C (140°F).
 - b. Calculate the dry masses using the moisture samples.

To determine the dry mass of the fractions using moisture samples.

- 1. Determine the moist mass of both fractions, fine (M_{Mf}) and oversized (M_{Mc}) :
- 2. Obtain moisture samples from the fine and oversized material.
- 3. Determine the moisture content of the fine particles (MC_f) and oversized particles (MC_C) of the material by FOP for AASHTO T 255/T 265 or agency approved method.
- 4. Calculate the dry mass of the oversize and fine particles.

$$M_D = \frac{M_m}{1 + MC}$$

Where:

 M_D = mass of dry material (fine or oversize particles)

 M_m = mass of moist material (fine or oversize particles)

MC = moisture content of respective fine or oversized, expressed as a decimal

5. Calculate the percentage of the fine (P_f) and oversized (P_c) particles by dry weight of the total sample as follows: See Note 2.

$$P_f = \frac{100 \times M_{DF}}{M_{DF} + M_{DC}} \qquad \frac{100 \times 15.4 \ lb}{15.4 \ lbs + 5.7 \ lb} = 73\% \qquad \frac{100 \times 6.985 \ kg}{6.985 \ kg + 2.602 \ kg} = 73\%$$

And

$$P_c = \frac{100 \times M_{DC}}{M_{DF} + M_{DC}} \qquad \frac{100 \times 5.7 \ lb}{15.4 \ lbs + 5.7 \ lb} = 27\% \qquad \frac{100 \times 2.585 kg}{6.985 \ kg + 2.585 \ kg} = 27\%$$

Or for P_c :

$$P_c = 100 - P_f$$

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 fine particles M_{DC} = mass of oversize particles

Optimum Moisture Correction Equation

1. Calculate the corrected moisture content as follows:

$$MC_T = \frac{\left(MC_F \times P_f\right) + \left(MC_c \times P_c\right)}{100} \qquad \frac{\left(13.2\% \times 73.0\%\right) + \left(2.1\% \times 27.0\%\right)}{100} = 10.2\%$$

Where:

MC_T = corrected moisture content of combined fines and oversized particles, expressed as a percent moisture

 MC_f = moisture content of fine particles, as a percent moisture MC_C = moisture content of oversized particles, as a percent moisture

Note 1: Moisture content of oversize material can be assumed to be two (2) percent for most construction applications.

Note 2: In some field applications agencies will allow the percentages of oversize and fine materials to be determined with the materials in the wet state.

Density Correction Equation

1. Calculate the corrected dry density of the total sample (combined fine and oversized particles) as follows:

$$D_d = \frac{100\%}{\left[\left(\frac{P_f}{D_f} \right) + \left(\frac{P_c}{k} \right) \right]}$$

Where:

 D_d = corrected total dry density (combined fine and oversized particles) kg/m³ (lb/ ft^3)

 D_f = dry density of the fine particles kg/m³ (lb/ft³), determined in the lab

P = percent of oversize particles, of sieve used, by weight.

 P_f = percent of fine particles, of sieve used, by weight.

 $k = Metric: 1,000 * Bulk Specific Gravity (<math>G_{sb}$) (oven dry basis) of coarse particles (kg/m^3).

k = English: 62.4 * Bulk Specific Gravity (G_{sb}) (oven dry basis) of coarse particles (Ib/ft^3)

Note 3: If the specific gravity is known, then this value will be used in the calculation. For most construction activities the specific gravity for aggregate may be assumed to be 2.600.

Calculation

Example

Metric:

Maximum laboratory dry density (D_f): 1890 kg/m³ Percent coarse particles (P_c): 27 percent Percent fine particles (P_f): 73 percent

Mass per volume coarse particles (k): $(2.697)(1000) = 2697 \text{ kg/m}^3$

$$\begin{split} D_d &= \frac{100\%}{\left[\left(\frac{P_f}{D_f}\right) + \left(\frac{P_c}{k}\right)\right]} \\ D_d &= \frac{100\%}{\left[\left(\frac{73\%}{1890\,kg/m^3}\right) + \left(\frac{27\%}{2697\,kg/m^3}\right)\right]} \\ D_d &= \frac{100\%}{\left[0.03862\,kg/m^3 + 0.01001\,kg/m^3\right]} \\ D_d &= 2056.3\,kg/m^3\,report\,\,2056\,kg/m^3 \end{split}$$

English:

Maximum laboratory dry density (D_f): 117.0 lb/ft³ Percent coarse particles (P_c): 27 percent

Percent fine particles (P_f) : 73 percent

Mass per volume of coarse particles (k): $(2.697) (62.4) = 168.3 \text{ lb/ft}^3$

$$D_{d} = \frac{100\%}{\left[\left(\frac{P_{f}}{D_{f}}\right) + \left(\frac{P_{c}}{k}\right)\right]}$$

$$D_{d} = \frac{100\%}{\left[\left(\frac{73\%}{117.0 \, lb/ft^{3}}\right) + \left(\frac{27\%}{168.3 \, lb/ft^{3}}\right)\right]}$$

$$D_{d} = \frac{100\%}{\left[0.6239 \, lb/ft^{3} + 0.1604 \, lb/ft^{3}\right]}$$

$$D_{d} = \frac{100\%}{0.7843 \, lb/ft^{3}}$$

$$D_{d} = 127.50 \, lb/ft^{3} \quad Report \ 127.5 \, lb/ft^{3}$$

Report

- · Results on forms approved by the agency
- Sample ID
- Corrected maximum dry density to the closest 1 kg/m³ (0.1 lb/ft³)
- · Corrected optimum moisture to the 0.1 percent

ANNEX B

Standardization of The Mold

Standardization is a critical step to ensure accurate test results when using this apparatus. Failure to perform the standardization procedure as described herein will produce inaccurate or unreliable test results.

Apparatus

- Mold and base plate
- Balance or scale Accurate to within 45 g (0.1 lb) or 0.3 percent of the test load, whichever is greater, at any point within the range of use.
- Cover plate A piece of plate glass, at least 6 mm (1/4 in) thick and at least 25 mm (1 in) larger than the diameter of the mold.
- Thermometers Standardized liquid-in-glass, or electronic digital total immersion type, accurate to 0.5°C (1°F)

Procedure

- 1. Create a watertight seal between the mold and base plate.
- 2. Determine and record the mass of the dry sealed mold, base plate, and cover plate.
- 3. Fill the mold with water at a temperature between 16°C and 29°C (60°F and 85°F) and cover with the cover plate in such a way as to eliminate bubbles and excess water.
- 4. Wipe the outside of the mold, base plate, and cover plate dry, being careful not to lose any water from the mold.
- 5. Determine and record the mass of the filled mold, base plate, cover plate, and water.
- 6. Determine and record the mass of the water in the mold by subtracting the mass in Step 2 from the mass in Step 5.
- 7. Measure the temperature of the water and determine its density from Table B1, interpolating as necessary.
- 8. Calculate the volume of the mold, V_m , by dividing the mass of the water in the mold by the density of the water at the measured temperature.

Calculations

$$V_m = \frac{M}{D}$$

Where:

V_m = volume of the mold M = mass of water in the mold

D = density of water at the measured temperature

Example

Mass of water in mold = 0.94061 kg (2.0737 lb)

Density of water at 23°C (73.4°F) = $997.54 \text{ kg/m}^3 (62.274 \text{ lb/ft}^3)$

$$V_m = \frac{0.94061 \, kg}{997.54 \, kg/m^3} = 0.000943 \, m^3$$
 $V_m = \frac{2.0737 \, lb}{62.274 \, lb/ft^3} = 0.0333 \, ft^3$

Table B1 Unit Mass of Water 15°C to 30°C

°C	(°F)	kg/m ³	(lb/ft ³)	°C	(°F)	kg/m ³	(lb/ft ³)
15	(59.0)	999.10	(62.372)	23	(73.4)	997.54	(62.274)
15.6	(60.0)	999.01	(62.366)	23.9	(75.0)	997.32	(62.261)
16	(60.8)	998.94	(62.361)	24	(75.2)	997.29	(62.259)
17	(62.6)	998.77	(62.350)	25	(77.0)	997.03	(62.243)
18	(64.4)	998.60	(62.340)	26	(78.8)	996.77	(62.227)
18.3	(65.0)	998.54	(62.336)	26.7	(80.0)	996.59	(62.216)
19	(66.2)	998.40	(62.328)	27	(80.6)	996.50	(62.209)
20	(68.0)	998.20	(62.315)	28	(82.4)	996.23	(62.192)
21	(69.8)	997.99	(62.302)	29	(84.2)	995.95	(62.175)
21.1	(70.0)	997.97	(62.301)	29.4	(85.0)	995.83	(62.166)
22	(71.6)	997.77	(62.288)	30	(86.0)	995.65	(62.156)

Report

- Mold ID
- · Date Standardized
- Temperature of the water
- Volume, V_m, of the mold

Performance Exam Checklist FOP for AASHTO T 99 MOISTURE-DENSITY RELATION OF SOILS

Part	icipant Name Exam Date		
Reco	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Prod	cedure Element	Trial 1	Trial 2
1.	If damp, sample dried in air or drying apparatus, not exceeding 60°C (140°F)?		
2.	Sample broken up and an adequate amount sieved over the appropriate sieve $(4.75 \text{ mm/No}. 4 \text{ or } 19.0 \text{ mm}/ \frac{3}{4} \text{ in})$ to determine oversize (coarse particle) percentage?		
3.	Sample passing the sieve has appropriate mass?		
4.	If soil is plastic (clay types):		
	a. Multiple samples mixed with water varying moisture content by 1 to 2 percent, bracketing the optimum moisture content?		
	b. Samples placed in covered containers and allowed to stand for at least 12 hours?		
5.	Sample determined to be 4 to 8 percent below expected optimum moisture content?		
6.	Mold placed on rigid and stable foundation?		
7.	Layer of soil (approximately one third compacted depth) placed in mold with collar attached, loose material lightly tamped?		
8.	Soil compacted with appropriate number of blows (25 or 56)?		
9.	Material adhering to the inside of the mold trimmed?		
10.	Layer of soil (approximately two thirds compacted depth) placed in mold with collar attached, loose material lightly tamped?		
11.	Soil compacted with appropriate number of blows (25 or 56)?		
12.	Material adhering to the inside of the mold trimmed?		
13.	Mold filled with soil such that compacted soil will be above the mold, loose material lightly tamped?		
14.	Soil compacted with appropriate number of blows (25 or 56)?		
15.	Collar removed without shearing off sample?		
16.	Approximately 6 mm ($\frac{1}{4}$ in) of compacted material above the top of the mold (without the collar)?		
17.	Soil trimmed to top of mold with the beveled side of the straightedge?		
18.	Mass of mold and contents determined to appropriate precision?		
19.	Wet density calculated from the wet mass?		

Pro	cedure Element	Trial 1	Trial 2
20.	Soil removed from mold using a sample extruder if needed?		
21.	Soil sliced vertically through center (non-granular material)?		
22.	Moisture sample removed ensuring all layers are represented?		
23.	Moist mass determined immediately to 0.1 g?		
24.	Moisture sample mass of correct size?		
25.	Sample dried and water content determined according to the FOP for T 255/T 265?		
26.	Remainder of material from mold broken up until it will pass through the sieve, as judged by eye, and added to remainder of original test sample?		
27.	Water added to increase moisture content of the remaining sample in 1 to 2 percent increments?		
28.	Steps 2 through 26 repeated for each increment of water added?		
29.	If material is degradable: Multiple samples mixed with water varying moisture content by 1 to 2 percent, bracketing the optimum moisture content?		
30.	Process continued until wet density either decreases or stabilizes?		
31.	Moisture content and dry density calculated for each sample?		
32.	Dry density plotted on vertical axis, moisture content plotted on horizontal axis, and points connected with a smooth curve?		
33.	Moisture content at peak of curve recorded as optimum water content and recorded to nearest 0.1 percent?		
34.	Dry density at optimum moisture content reported as maximum density to nearest 1 kg/m^3 (0.1 lb/ft ³)?		
35.	Corrected for coarse particles if applicable?		
First	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
_			

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 85 Portland Cement
 - M 152 Mixing Rooms, 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
 - M 240 Blended Hydraulic Cements
 - M 295 Coal Fly Ash and Raw or Calcined Natural Pozzolan for Use in Concrete
 - M 302 Ground Granulated Blast-Furnace Slag for Use in Concrete and Mortars
 - R11 Recommended Practice for Indicating Which Places of Figures Are to be Considered Significant in Specified Limiting Values
 - T 105 Chemical Analysis of Hydraulic Cement
 - T 162 Mechanical Mixing of Hydraulic Cement Pastes and Mortars of Plastic Consistency

¹ This Test Method is based on AASHTO T 106-09

2.2 ASTM Standards

- C 91 Masonry Cement
- 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
- C 1157 Hydraulic Cement
- C 1328 Plastic (Stucco) Cement
- C 1329 Mortar Cement
- **IEEE/ASTM SI 10**

Standard for Use of the International System of Units (SI): The Modern Metric System

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.

	2 in Cub	e 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 Compartment	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 Faces ^A	90 ± 0.5°	90 ± 0.5°	90 ± 0.5°	90 ± 0.5°	

Notes:

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.

Permissible Variations of Specimen Molds Table 1

- 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 ½ 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 ½50 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 ½5 in or (1 mm) and ½6 in or (1.6 mm). When the spacing is between ½6 in or (1.6 mm) and ½6 in or (3.2 mm), one third of the scale interval can be read with reasonable certainty. When the spacing is ½6 in or (3.2 mm) or more, one fourth of the scale interval can be read with reasonable certainty.
- 592 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. A hardened metal bearing block shall be used beneath the specimen to minimize wear of the lower platen of the machine. To facilitate accurate centering of the test specimen in the compression machine, one of the two surfaces of the bearing blocks shall have a diameter or diagonal of between 70.7 mm (2.83 in) (See Note 3) and 2.9 in (73.7 mm). When the upper block surface meets this requirement, the lower bearing surface shall be greater than 70.7 (2.83 in). When the lower block bearing surface meets this requirement, the diameter or diagonal of upper block bearing surface shall be between 70.7 and 79.4 (2.83 and 3\\ in). When the lower block is the only block with a diameter or diagonal between 70.7 and 73.7 mm [2.83 in and 2.9 in], the lower block shall be used to center the test specimen. In that case, the lower block shall be centered with respect to the upper bearing block and held in position by suitable means. 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 0.013 mm [0.0005 in] when the blocks are new and shall be maintained within a permissible variation of 0.001 in (0.025 mm).
 - **Note 3:** The diagonal of 2 in or (50 mm) cube is 2.83 in (70.7 mm).

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 \pm 3°F or (\pm 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:

	No. of Sp	pecimens
	6	9
Cement, g	500	740
Sand, g	1375	2035
Water, mL: Portland (0.485)	242	359
Air-entraining portland (0.460)	230	340
Other (to flow of 110 ± 5)	_	-

Mixing Proportions for 2 in Cubes *Table 2*

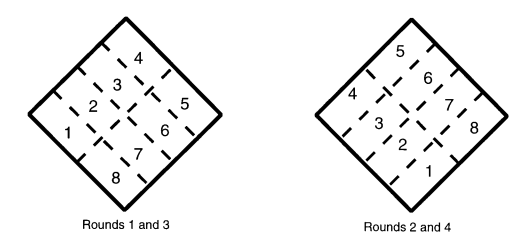
- 10.2 Preparation of Mortar
 - 10.2.1 Mechanically mix in accordance with the procedure given in T 162.
- 10.3 Determination of Flow
 - 10.3.1 Carefully wipe the flow-table top clean and dry, and place the flow mold at the center. Place a layer of mortar about 1 in or (25 mm) in thickness in the mold 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.
- 10.3.4 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.
- 10.3.5 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 Molding Test Specimens

10.4.1 Complete the consolidation of the mortar in the molds either by hand tamping or by a qualified alternative method. Alternative methods include but are not limited to the use of vibrating table or mechanical devices.



Order of Tamping in Molding of Test specimens
Figure 1

- Hand Tamping-Start molding the specimens within a total elapsed time of not more 10.4.2 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.
- 10.4.3 Alternative Methods Any consolidation method may be used that meets the qualification requirements of this section. The consolidation method consists of a specific procedure, equipment and consolidation device, as selected and used in a consistent manner by a specific laboratory. The mortar batch size of the method may be modified to accommodate the apparatus, provided the proportions maintain the same ratios as given in Section 10.1.2.
 - 10.4.3.1 Separate qualifications are required for the following classifications:
 - 10.4.3.2 *Class A, Non-air Entrained Cements* For use in concrete. Refer to M 85, M 240 and ASTM C 1157.
 - 10.4.3.3 *Class B, Air-entrained Cements* For use in concrete. Refer to M 85, M 240 and ASTM C 1157.
 - 10.4.3.4 Class C, Masonry, Mortar and Stucco Cements Refer to ASTM C 91, C 1328, and C 1329.
 - 10.4.3.5 An alternative method may only be used to test the cement types as given in Section 10.4.3.1 above, for which it has been qualified.
 - 10.4.3.6 It can also be used for Strength Activity Index determinations for fly ash and slag, refer to M 295 and M 302, provided the alternative method has qualified for both Class A and Class C cements.

- 10.4.4 *Qualification Procedure* Contact CCRL to purchase cement samples that have been used in the Proficiency Sample Program (PSP). Four samples (5 kg each) of the class to be qualified will be required to complete a single qualification (See Note 7).
 - 10.4.4.1 In one day, prepare replicate six-cube or nine-cube batches using one of the cements and cast a minimum of 36 cubes. Complete one round of tests on each cement on different days. Store and test all specimens as prescribed in the sections below. Test all cubes at the age of seven-days.
 - 10.4.4.2 Tabulate the compressive strength data and complete the mathematical analyses as instructed in Annex A1.
- 10.4.5 Re-qualification of the Alternate Compaction Method:
 - 10.4.5.1 Re-qualification of the method shall be required if any of the following occur:
 - Evidence that the method may not be providing data in accordance with the requirements of Table 2.
 - Results that differ from the reported final average of a CCRL-PSP sample with a rating of three or less.
 - Results that differ from the accepted value of a known reference sample with established strength values by more than twice the multi-laboratory 1s percent values of Table 2. Before starting the re-qualification procedure, evaluate all aspects of cube fabrication and testing process to determine if the offending result is due to some systematic error or just an occasional random event.
 - 10.4.5.2 If the compaction equipment is replaced, significantly modified, repaired, or has been recalibrated, re-qualify the equipment in accordance with Section 10.4.4.
 - **Note 7:** It is recommended that a large homogenous sample of cement be prepared at the time of qualification for use as a secondary standard and for method evaluation. Frequent testing of this sample will give early warning of any changes in the performance of the apparatus.
- 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:

Test Age	Permissible Tolerance
24 hrs	± ½ hr
3 days	± 1 hr
7 days	± 3 hr
28 days	± 12 hr

Testing Time tolerances *Table 3*

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 $73.4 \pm 3^{\circ}$ F or $(23 \pm 2^{\circ}\text{C})$ 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 8). 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 8:** *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 9: 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 = P/A (1)
where:
fm = compressive strength in psi or (MPa),
P = total maximum load in lbf or (N), and
A = area of loaded surface 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 10).
 - **Note 10:** 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 11:** 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 106-09 for Precision and Bias.

Performance Exam Checklist

Participant Name

Compressive Strength of Hydraulic Cement Mortar for AASHTO T 106

Pro	cedure 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.	Cubes are broken within permissible tolerance for time?		
4.	Cubes tested immediately after removal from moist closet?		
5.	Specimens covered with damp cloth while out of moist room closet?		
6.	Cubes wiped clean of sand, and wiped to surface dry condition prior to testing?		
7.	Faces to contact the bearing blocks are those that were in contact with the mold?		
8.	Faces that will contact the bearing blocks checked with a straightedge?		
9.	Cross-sectional area determined in respect to those faces contacting the bearing blocks?		
10.	Prior to testing each cube, the spherically seated block was checked for freedom to tilt?		
11.	Load rate of 200 to 400 lbf/s (900-1800 N/s) obtained during the first half of the anticipated load?		
12.	No adjustment in rate was made during the second half of the loading?		
13.	Compressive strength of cubes averaged and reported to the nearest 10 psi (0.1 MPa)?		
First	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	nature of Examiner		

Exam Date

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 $230 \pm 9^{\circ}F$ ($110 \pm 5^{\circ}C$) 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 $230 \pm 9^{\circ}F$ ($110 \pm 5^{\circ}C$) 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 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 $1\frac{3}{4}$ 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 $7\frac{1}{2}$ in $\pm \frac{1}{4}$ in (190.5 mm \pm 6.3 mm) diameter x 6 $\pm \frac{1}{2}$ in (152.4 mm \pm 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 D 3744)
- d. Sand equivalent graduated cylinder and rubber stopper
- e. Sand equivalent stock solution
- f. Sieves $-\frac{1}{2}$ in (12.5 mm), $\frac{3}{8}$ in (9.5 mm), $\frac{1}{4}$ 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 ASTM E11
- 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 $230 \pm 9^{\circ}F$ ($110 \pm 5^{\circ}C$)
- 1. 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 $230 \pm 9^{\circ}F$ ($110 \pm 5^{\circ}C$) to allow for clean separation from the fine material. Separate the material over the $\frac{1}{2}$ in (12.5 mm) sieve and discard that finer than the $\frac{1}{2}$ in (12.5 mm) and proceed to step 3d.
- b. If testing crushed and stockpiled material: dry at 230 ± 9 °F (110 ± 5 °C) 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 ½ in (12.5 mm) sieve and discard that finer than the ½ 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 ½ in (12.5 mm), ¾ in (9.5 mm), ¼ 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 $\frac{1}{2}$ - $\frac{3}{8}$ (12.5-9.5 mm), 550 g of $\frac{3}{8}$ - $\frac{1}{4}$ (9.5-6.3 mm), and 1100 g of $\frac{1}{4}$ -#10 (6.3-2.00 mm).
- h. Combine the $\frac{1}{2}$ - $\frac{3}{8}$ (12.5-9.5 mm) with the $\frac{3}{8}$ - $\frac{1}{4}$ (9.5-6.3 mm).
- i. Wash the ½-¼ (12.5-6.3 mm) and ¼-#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 $230 \pm 9^{\circ}$ F ($110 \pm 5^{\circ}$ C).
- k. Allow to cool to room temperature.
- 1. 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.
- 1. 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. $(381 \pm 2.5 \text{ mm})$ 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:

$$D_1 = \frac{(15-H_1)}{(15+1.75H_1)\times 100} \qquad D_2 = \frac{(15-H_2)}{(15+1.75H_2)\times 100}$$

Note: Table 1 may be used to determine the values of D1 and D2 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

H₂ = 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, D₁ & D₂ must agree within 6 points.
- b. Repeat the entire test if variation between the test samples exceeds 6 points, see following calculation:

Absolute Value
$$(D_1-D_2) > 6$$

$$D = \frac{(15-H)}{(15+1.75H)} \times 100$$

н	D	н	D	Н	D	н	D	Н	D
0.0	100	3.1	58	6.1	35	9.1	19	12.1	8
0.1	98	3.2	57	6.2	34	9.2	19	12.2	8
0.2	96	3.3	56	6.3	33	9.3	18	12.3	7
0.3	95	3.4	55	6.4	33	9.4	18	12.4	7
0.4	93	3.5	54	6.5	32	9.5	17	12.5	7
0.5	91	3.6	54	6.6	32	9.6	17	12.6	6
0.6	90	3.7	53	6.7	31	9.7	17	12.7	6
0.7	88	3.8	52	6.8	30	9.8	16	12.8	6
8.0	87	3.9	51	6.9	30	9.9	16	12.9	6
0.9	85	4.0	50	7.0	29	10.0	15	13.0	5
1.0	84								
1.1	82	4.1	49	7.1	29	10.1	15	13.1	5
1.2	81	4.2	48	7.2	28	10.2	15	13.2	5
1.3	79	4.3	48	7.3	28	10.3	14	13.3	4
1.4	78	4.4	47	7.4	27	10.4	14	13.4	4
1.5	77	4.5	46	7.5	27	10.5	13	13.5	4
1.6	75	4.6	45	7.6	26	10.6	13	13.6	4
1.7	74	4.7	44	7.7	26	10.7	13	13.7	3
1.8	73	4.8	44	7.8	25	10.8	12	13.8	3
1.9	71	4.9	43	7.9	25	10.9	12	13.9	3
2.0	70	5.0	42	8.0	24	11.0	12	14.0	3
2.1	69	5.1	41	8.1	24	11.1	11	14.1	2
2.2	68	5.2	41	8.2	23	11.2	11	14.2	2
2.3	67	5.3	40	8.3	23	11.3	11	14.3	2
2.4	66	5.4	39	8.4	22	11.4	10	14.4	1
2.5	65	5.5	39	8.5	22	11.5	10	14.5	1
2.6	63	5.6	38	8.6	21	11.6	10	14.6	1
2.7	62	5.7	37	8.7	21	11.7	9	14.7	1
2.8	61	5.8	37	8.8	20	11.8	9	14.8	0
2.9	60	5.9	36	8.9	20	11.9	9	14.9	0
3.0	59	6.0	35	9.0	20	12.0	8	15.0	0

Degradation Value "D" *Table 1*

Performance Exam Checklist

WSDOT TM 113

Method of Test for Determination of Degradation Value

Participant Name	Exam Date	
Procedure Element		Yes No

Equipment

- 1. Balance 5000g capacity, sensitive to 0.1g- Calibrated?
- 2. Degradation Shaker $-1\frac{3}{4}$ " throw, 300 ± 5 oscillations per minute Verified?
- 3. Canister plastic, 7½ in diameter x 6 in high, walls meet floor at 90 deg with min fillet, or steel meeting AASHTO T210, or ASTM D 3744?
- 4. Sand Equivalent Cylinder & Rubber Stopper?
- 5. Sand Equivalent Stock Solution?
- 6. Sieves $-\frac{1}{2}$, $\frac{3}{8}$, $\frac{1}{4}$, No. 10, No. 200 Verified?
- 7. Graduates 500 ml tall form & 100 ml?
- 8. Interval Timer Verified?
- 9. Funnel Large enough to hold the sieves with a mouth that fits in the 500 ml graduate?
- 10. Sieve Shaker(s) Verified?
- 11. Oven verified at $230 \pm 9^{\circ}$ F.- Calibrated?
- 12. Sprayer produces a low volume stream of water?
- 13. Containers suitable for drying and washing?

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 $\frac{1}{2}$?
- 3. Split out approx. 5000g?
- 4. Separate the material over the $\frac{1}{2}$, $\frac{3}{8}$, $\frac{1}{4}$, and No. 10?
- 5. Split or quarter approx. $550g^{1/2}-3/8$, $550g^{3/8}-1/4$, & $1100g^{1/4}-No. 10$?
- 6. Combine the $\frac{1}{2}$ - $\frac{3}{8}$ with the $\frac{3}{8}$ - $\frac{1}{4}$?
- 7. Hand wash the $\frac{1}{2}$ - $\frac{1}{4}$ and $\frac{1}{4}$ -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 ± 1 g portions, hand selection ok to 50g?
- 10. Combine to create two $1000 \pm 2g$, $\frac{1}{2}$ No. 10 test samples?
- 11. Place one sample into a canister, cover with 200 ± 5 ml water, cover & shake for 20 min?

Procedure Element Yes No

- 12. Empty canister into the nested No. 10 & No. 200 fitted in the funnel over the 500 ml grad.?
- 13. Rinse the lid into the canister and then the canister into the nested sieves?
- 14. Shake the sieves to spread the sample?
- 15. Wash using only 20-50 ml.?
- 16. Shake the sieves to release trapped water and then lift observing liquid for clarity?
- 17. Repeat 14-16 until water reaches the 500 ml mark water not to exceed 500 ml?
- 18. No loss of fines or liquid during the washing process?
- 19. Place 7 ± 1 ml of SE Stock Solution in a SE Graduated Cylinder?
- 20. Turn capped 500 ml upside down & back allowing bubble to traverse 10 cycles?
- 21. Immediately pour into a SE Cylinder to the 15 ± 0.1 mark no settling allowed?
- 22. Rubber stopper inserted and SE Cylinder turned upside down & back 20 cycles?
- 23. Place gently, remove stopper, start timer, allow to sit undisturbed for 20 min?
- 24. Record height of column to nearest 0.1 in?
- 25. Repeat for second sample?
- 26. Calculations performed correctly?
- 27. Second sample must be within 6 points?

First Attempt:	Pass	Fail	Second Attempt:	Pass	Fail
Signature of Exa	aminer				

FOP for AASHTO T 119

Slump of Hydraulic Cement Concrete

Scope

This procedure provides instructions for determining the slump of hydraulic cement concrete in accordance with AASHTO T 119-13. It is not applicable to non-plastic and non-cohesive concrete.

Warning—Fresh Hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.

Apparatus

- Mold: A metal frustum of a cone provided with foot pieces and handles. The mold
 must be constructed without a seam. The interior of the mold shall be relatively
 smooth and free from projections such as protruding rivets. The mold shall be free
 from dents. A mold that clamps to a rigid nonabsorbent base plate is acceptable
 provided the clamping arrangement is such that it can be fully released without
 movement of the mold.
- Mold: If other than metal, it must conform to AASHTO T 119, Sections 5.1.2.1 and 5.1.2.2.
- Tamping rod: 16 mm (% in) diameter and 400 mm (16 in) to 600 mm (24 in) long, having a hemispherical tip the same diameter as the rod. (Hemispherical means "half a sphere"; the tip is rounded like half of a ball.)
- Scoop: a receptacle of appropriate size so that each representative increment of the concrete sample can be placed in the container without spillage.
- Tape measure or ruler with at least 5 mm or \% in graduations
- Base: Flat, rigid, non-absorbent moistened surface on which to set the slump mold

Procedure

- Obtain the sample in accordance with the FOP for WAQTC TM 2. If the concrete mixture contains aggregate retained on the 37.5 mm (1½ in) sieve, the aggregate must be removed in accordance with the Wet Sieving portion of the FOP for WAQTC TM 2.
 - **Note 1:** Testing shall begin within five minutes of obtaining the sample.
- 2. Dampen the inside of the mold and place it on a dampened, rigid, nonabsorbent surface that is level and firm.
- 3. Stand on both foot pieces in order to hold the mold firmly in place.
- 4. Use the scoop to fill the mold ½ full by volume, to a depth of approximately 67 mm (25% in) by depth.
- 5. 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.
- 6. For this bottom layer, incline the rod slightly and make approximately half the strokes near the perimeter, and then progress with vertical strokes, spiraling toward the center.

- 7. Use the scoop to fill the mold \(^2\) full by volume, to a depth of approximately 155 mm (6\(^1\) in) by depth.
- 8. Consolidate this layer with 25 strokes of the tamping rod, penetrate approximately 25 mm (1 in) into the bottom layer. Distribute the strokes evenly.
- 9. Use the scoop to fill the mold to overflowing.
- 10. Consolidate this layer with 25 strokes of the tamping rod, penetrate approximately 25 mm (1 in) into the second layer. Distribute the strokes evenly. If the concrete falls below the top of the mold, stop, add more concrete, and continue rodding for a total of 25 strokes. Keep an excess of concrete above the top of the mold at all times. Distribute strokes evenly as before.
- 11. Strike off the top surface of concrete with a screeding and rolling motion of the tamping rod.
- 12. Clean overflow concrete away from the base of the mold.
- 13. Remove the mold from the concrete by raising it carefully in a vertical direction. Raise the mold 300 mm (12 in) in 5 ± 2 seconds by a steady upward lift with no lateral or torsional (twisting) motion being imparted to the concrete.
- 14. The entire operation from the start of the filling through removal of the mold shall be carried out without interruption and shall be completed within an elapsed time of $2\frac{1}{2}$ minutes. Immediately measure the slump.
- 15. Invert the slump mold and set it next to the specimen.
- 16. Lay the tamping rod across the mold so that it is over the test specimen.
- 17. Measure the distance between the bottom of the rod and the displaced original center of the top of the specimen to the nearest 5 mm (1/4 in).
 - **Note 2:** If a decided falling away or shearing off of concrete from one side or portion of the mass occurs, disregard the test and make a new test on another portion of the sample. 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 the plasticity and cohesiveness necessary for the slump test to be applicable.
- 18. Discard the tested sample.

Report

- · Results on forms approved by the agency
- Sample ID
- Slump to the nearest 5 mm (¼ in).

Performance Exam Checklist FOP for AASHTO T 119 Slump of Hydraulic Cement Concrete

Part	Participant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
Firs	t layer		
1.	Mold and floor or base plate dampened?		
2.	Mold held firmly against the base by standing on the two foot pieces? Mold not allowed to move in any way during filling?		
3.	Representative sample scooped into the mold, moving a scoop around the perimeter of the mold to evenly distribute the concrete as discharged?		
4.	Mold approximately one third (by volume), 67 mm (2½ in) deep?		
5.	Layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes?		
Sec	ond layer		
6.	Representative samples scooped into the mold, moving a scoop around the perimeter of the mold to evenly distribute the concrete as discharged?		
7.	Mold filled approximately two thirds (by volume), 155 mm (61/4 in), deep?		
8.	Layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes, penetrate approximately 25 mm (1 in) into the bottom layer?		
Thir	rd layer		
9.	Representative sample scooped into the mold, moving a scoop around the perimeter of the mold to evenly distribute the concrete as discharged??		
10.	Mold filled to just over the top of the mold?		
11.	Layer rodded throughout its depth 25 times with hemispherical end of rod, uniformly distributing strokes, penetrate approximately 25 mm (1 in) into the second layer?		
12.	Excess concrete kept above the mold at all times while rodding?		
13.	Concrete struck off level with top of mold using tamping rod?		
14.	Concrete removed from around the outside bottom of the mold?		
15.	Mold lifted upward 300 mm (12 in) in one smooth motion, without a lateral or twisting motion of the mold, in 5 ± 2 seconds?		
16.	Test performed from start of filling through removal of the mold within 2½ minutes?		
17.	Slump immediately measured to the nearest 5 mm (¼ in) from the top of the mold to the displaced original center of the top surface of the specimen?		

First Attempt: Pass	Fail	Second Attempt:	Pass	Fail
Signature of Examiner _			WA	QTC #:
This checklist is derived, American Concrete Instit		hted material print	ed in ACI	CP-1, published by the
Comments:				

FOP for AASHTO T 121

Density (Unit Weight), Yield, and Air Content (Gravimetric) of Concrete

Scope

This procedure covers the determination of density, or unit weight, of freshly mixed concrete in accordance with AASHTO T 121-17. It also provides formulas for calculating the volume of concrete produced from a mixture of known quantities of component materials, and provides a method for calculating cement content and cementitious material content – the mass of cement or cementitious material per unit volume of concrete. A procedure for calculating water/cement ratio is also covered.

Warning—Fresh Hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.

Apparatus

- Measure: May be the bowl portion of the air meter used for determining air content under the FOP for AASHTO T 152. Otherwise, it shall be a metal cylindrical container meeting the requirements of AASHTO T 121. The capacity and dimensions of the measure shall conform to those specified in Table 1.
- Balance or scale: Accurate to within 45 g (0.1 lb) or 0.3 percent of the test load, whichever is greater, at any point within the range of use.
- Tamping rod: 16 mm (% in) diameter and 400 mm (16 in) to 600 mm (24 in) long, having a hemispherical tip the same diameter as the rod. (Hemispherical means "half a sphere"; the tip is rounded like half of a ball.)
- Vibrator: 7000 vibrations per minute, 19 to 38 mm ($\frac{3}{4}$ to $\frac{1}{2}$ in) in diameter, and the length of the shaft shall be at least 610 mm (24 in).
- Scoop: a receptacle of appropriate size so that each representative increment of the concrete sample can be placed in the container without spillage.
- Strike-off plate: A flat rectangular metal plate at least 6 mm (¼ in) thick or a glass or acrylic plate at least 12 mm (½ in) thick, with a length and width at least 50 mm (2 in) 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 1.5 mm (½ in).
- Mallet: With a rubber or rawhide head having a mass of 0.57 ± 0.23 kg (1.25 ± 0.5 lb) for use with measures of 0.014 m³ (½ ft³) or less, or having a mass of 1.02 ± 0.23 kg (2.25 ± 0.5 lb) for use with measures of 0.028 m³ (1 ft³).

Nominal Maximum
Size of Coarse
Aggregate***
mm (in)

25
(1)
50
(2)
76
(3)

	Inside	Inside		hicknesses (in)
Capacity m ³ (ft ³)	Diameter mm (in)	Height mm (in)	Bottom	Wall
0.0071	203 ± 2.54	213 ± 2.54	5.1	3.0
(1/4)**	(8.0 ± 0.1)	(8.4 ± 0.1)	(0.20)	(0.12)
0.0142	254 ± 2.54	279 ± 2.54	5.1	3.0
(1/2)	(10.0 ± 0.1)	(11.0 ± 0.1)	(0.20)	(0.12)
0.0283	356 ± 2.54	284 ± 2.54	5.1	3.0
(1)	(14.0 ± 0.1)	(11.2 ± 0.1)	(0.20)	(0.12)

Table 1 Dimensions of Measures*

*Note: The indicated size of measure shall be for aggregates of nominal maximum size equal to or smaller
than that listed.

 $^{^{**}}$ Measure may be the base of the air meter used in the FOP for AASHTO T 152.

Procedure Selection

There are two methods of consolidating the concrete – rodding and vibration. If the slump is greater than 75 mm (3 in), consolidation is by rodding. When the slump is 25 to 75 mm (1 to 3 in), internal vibration or rodding 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 concrete with slump less than 25 mm (1 in), consolidate the sample by internal vibration. Do not consolidate self-consolidating concrete (SCC).

When using measures greater than 0.0142 m³ (½ ft³) see AASHTO T 121.

Procedure - Rodding

 Obtain the sample in accordance with the FOP for WAQTC TM 2. Testing may be performed in conjunction with the FOP for AASHTO T 152. When doing so, this FOP should be performed prior to the FOP for AASHTO T 152.

Note 1: If the two tests are being performed using the same sample, this test shall begin within five minutes of obtaining the sample.

- 2. Determine the mass of the dry empty measure.
- 3. Dampen the inside of the measure.
- 4. Use the scoop to fill the measure approximately ½ full with concrete. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 5. 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.
- 6. Tap around the perimeter of the measure smartly 10 to 15 times with the mallet to close voids and release trapped air.

^{***} Nominal maximum size: One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size.

- 7. Add the second layer, filling the measure about \(^2\) full. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 8. Consolidate this layer with 25 strokes of the tamping rod, penetrating about 25 mm (1 in) into the bottom layer.
- 9. Tap around the perimeter of the measure smartly 10 to 15 times with the mallet.
- 10. Add the final layer, slightly overfilling the measure. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 11. Consolidate this layer with 25 strokes of the tamping rod, penetrating about 25 mm (1 in) into the second layer.
- 12. Tap around the perimeter of the measure smartly 10 to 15 times with the mallet.
 - **Note 2:** The measure should be slightly over full, about 3 mm (½ in) above the rim. If there is a great excess of concrete, remove a portion with the scoop. If the measure 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.
- 13. Strike off by pressing the strike-off plate flat against the top surface, covering approximately \(^2\) of the measure. Withdraw the strike-off plate with a sawing motion to finish the \(^2\) originally covered. Cover the original \(^2\) again with the plate; finishing the remaining \(^1\) with a sawing motion (do not lift the plate; continue the sawing motion until the plate has cleared the surface of the measure). Final finishing may be accomplished with several strokes with the inclined edge of the strike-off plate. The surface should be smooth and free of voids.
- 14. Clean off all excess concrete from the exterior of the measure including the rim.
- 15. Determine and record the mass of the measure and the concrete.
- 16. If the air content of the concrete is to be determined, proceed to Rodding Procedure Step 13 of the FOP for AASHTO T 152.

Procedure - Internal Vibration

- 1. Perform Steps 1 through 3 of the rodding procedure.
- 2. Use the scoop to fill the measure approximately ½ full with concrete. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 3. Insert the vibrator at three different points in each layer. Do not let the vibrator touch the bottom or side of the measure.
 - **Note 3:** Remove the vibrator slowly, so that no air pockets are left in the material.
 - **Note 4:** 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.
- 4. Slightly overfill the measure. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 5. Insert the vibrator as in Step 3. Do not let the vibrator touch the side of the measure, but do penetrate the first layer approximately 25 mm (1 in).
- 6. Return to Step 13 of the rodding procedure and continue.

Procedure – Self Consolidating Concrete

- 1. Perform Steps 1 through 3 of the rodding procedure.
- 2. Use the scoop to slightly overfill the measure. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 3. Complete steps 13 thru 16 of the rodding procedure.

Calculations

Density

$$D = \frac{M_m}{V_m}$$

Where:

D = density of the concrete mix M_m = mass of concrete in measure V_m = volume of measure (Annex A)

Yield m³

$$Y_{m^3} = \frac{W}{D}$$

Where:

 Y_m^3 = yield (m³ of the batch of concrete) W = total mass of the batch of concrete

Yield yd3

$$Y_{ft^3} = \frac{W}{D} Y_{yd^3} = \frac{Y_{ft^3}}{27ft^3/yd^3}$$

Where:

Yft³ = yield (ft³ of the batch of concrete) Yyd³ = yield (yd³ of the batch of concrete) W = total mass of the batch of concrete D = density of the concrete mix

Note 5: The total mass, W, includes the masses of the cement, water, and aggregates in the concrete.

Cement Content

$$N = \frac{N_t}{Y}$$

Where:

N = actual cementitous material content per Ym³ or Yyd³ Nt = mass of cementitious material in the batch $Y = Y_m^3$ or Y_{vd}^3

Note 6: Specifications may require Portland Cement content and supplementary cementitious materials content.

Water Content

The mass of water in a batch of concrete is the sum of:

- · water added at batch plant
- · water added in transit
- water added at jobsite
- free water on coarse aggregate*
- free water on fine aggregate*
- liquid admixtures (if required by the agency)

This information is obtained from concrete batch tickets collected from the driver. Use the Table 2 to convert liquid measures.

Table 2 Liquid Conversion Factors

To Convert From	То	Multiply By
Liters, L	Kilograms, kg	1.0
Gallons, gal	Kilograms, kg	3.785
Gallons, gal	Pounds, Ib	8.34
Milliliters, mL	Kilograms, kg	0.001
Ounces, oz	Milliliters, mL	28.4
Ounces, oz	Kilograms, kg	0.0284
Ounces, oz	Pounds, Ib	0.0625
Pounds, lb	Kilograms, kg	0.4536

^{*}Mass of free water on aggregate

$$Free\ Water\ Mass = \textit{CA or FC Aggregate} - \frac{\textit{CA or FC Aggregate}}{1 + (\textit{Free Water Percentage}/100)}$$

Where:

Free Water Mass = on coarse or fine aggregate
FC or CA Aggregate = mass of coarse or fine aggregate
Free Water Percentage = percent of moisture of coarse or fine aggregate

Water/Cement Ratio

$$\frac{Water\ Content}{C}$$

Where:

Water Content = sum mass of water in the batch C = sum mass of cementitious materials

Example

Mass of concrete in measure (Mm) 16.290 kg (36.06 lb)

Volume of measure (Vm) 0.007079 m³ (0.2494 ft³)

From batch ticket:

Yards batched 4 yd³

Cement 950 kg (2094 lb)

Fly ash 180 kg (397 lb)

Coarse aggregate 3313 kg (7305 lb) Fine aggregate 2339 kg (5156 lb)

Water added at plant 295 L (78 gal)

Other

Water added in transit 0

Water added at jobsite 38 L (10 gal)

Total mass of the batch of concrete (W) 7115 kg (15,686 lb)

Moisture content of coarse aggregate 1.7 percent

Moisture content of coarse aggregate 5.9 percent

Density

$$D = \frac{M_m}{V_m}$$

$$D = \frac{16.920 \, kg}{0.007079 \, m^3} = 2390 \, kg/m^3 \, D = \frac{36.06 \, lb}{0.2494 \, ft^3} = 144.6 \, lb/ft^3$$

Given:

Mm = 16.920 kg (36.06 lb) Vm = 0.007079 m³ (0.2494 ft³) (Annex A)

Yield m³

$$Y_{m^3} = \frac{W}{D}$$

$$Y_{m^3} = \frac{7115 \, kg}{2390 \, kg/m^3} = 2.98 \, m^3$$

Given:

Total mass of the batch of concrete (W), kg = 7115 kg

Yield yd3

$$Y_{ft^3} = \frac{W}{D} \qquad Y_{yd^3} = \frac{Y_{ft^3}}{27ft^3/yd^3}$$

$$Y_{ft^3} = \frac{15,686 \, lb}{144.6 \, lb/ft^3} = 108.48 \, ft^3 \qquad Y_{yd^3} = \frac{108.48 \, ft^3}{27 \, ft^3/yd^3} = 4.02 \, yd^3$$

Given:

Total mass of the batch of concrete (W), lb = 15,686 lb

Cement Content

$$N = \frac{N_t}{Y}$$

$$N = \frac{950 \, kg + 180 \, kg}{2.98 \, m^3} = 379 \, kg/m^3 \, N = \frac{2094 \, lb + 397 \, lb}{4.02 \, yd^3} = 620 \, lb/yd^3$$

Given:

 N_t (cement) = 950 kg (2094 lb) N_t (flyash) = 180 kg (397 lb) $Y = Ym^3$ or Yyd^3

Note 6: Specifications may require Portland Cement content and supplementary cementitious materials content.

Free water

Free Water Mass = CA or FC Aggregate
$$-\frac{CA \text{ or FC Aggregate}}{1 + (Free Water Percentage/100)}$$

$$CA \text{ Free Water} = 3313 \text{ kg} - \frac{3313 \text{ kg}}{1 + (1.7/100)} = 55 \text{ kg}$$

$$CA \text{ Free Water} = 7305 \text{ lb} - \frac{7305 \text{ lb}}{1 + (1.7/100)} = 122 \text{ lb}$$

$$FA \text{ Free Water} = 2339 \text{ kg} - \frac{2339 \text{ kg}}{1 + (5.9/100)} = 130 \text{ kg}$$

$$FA \text{ Free Water} = 5156 \text{ lb} - \frac{5156 \text{ lb}}{1 + (5.9/100)} = 287 \text{ lb}$$

Given:

CA aggregate = 3313 kg (7305 lb) FC aggregate = 2339 kg (5156 lb) CA moisture content = 1.7 percent FC moisture content = 5.9 percent

Water Content

Sum of all water in the mix.

$$Water\ Content = [(78\ gal + 10\ gal) * 3.785\ kg/gal] + 55\ kg + 130\ kg = 518\ kg$$

$$Water\ Content = [(78\ gal + 10\ gal) * 8.34\ lb/gal] + 130\ lb + 287\ lb = 1151\ lb$$
 Given:

Water added at plant = 295 L (78 gal) Water added at the jobsite = 38 L (10 gal)

Water/Cement Ratio

$$W/C = \frac{518 \, kg}{950 \, kg + 180 \, kg} = 0.458 \quad W/C = \frac{1151 \, lb}{2094 \, lb + 397 \, lb} = 0.462$$

Report 0.46

Report

- Results on forms approved by the agency
- Sample ID
- Density (unit weight) to 1 kg/m³ (0.1 lb/ft³)
- Yield to 0.01 m³ (0.01 yd³)
- Cement content to 1 kg/m³ (1 lb/yd³)
- Cementitious material content to 1 kg/m³ (1 lb/yd³)
- Water/Cement ratio to 0.01

ANNEX A

Standardization of Measure

Standardization is a critical step to ensure accurate test results when using this apparatus. Failure to perform the standardization procedures as described herein will produce inaccurate or unreliable test results.

Apparatus

- Listed in the FOP for AASHTO T 121
 - Measure
 - Balance or scale
 - Strike-off plate
- Thermometer: Standardized liquid-in-glass, or electronic digital total immersion type, accurate to 0.5°C (1°F)

Procedure

- 1. Determine the mass of the dry measure and strike-off plate.
- 2. Fill the measure with water at a temperature between 16°C and 29°C (60°F and 85°F) and cover with the strike-off plate in such a way as to eliminate bubbles and excess water.
- Wipe the outside of the measure and cover plate dry, being careful not to lose any water from the measure.
- 4. Determine the mass of the measure, strike-off plate, and water in the measure.
- 5. Determine the mass of the water in the measure by subtracting the mass in Step 1 from the mass in Step 4.
- 6. Measure the temperature of the water and determine its density from Table A1, interpolating as necessary.
- 7. Calculate the volume of the measure, V_m, by dividing the mass of the water in the measure by the density of the water at the measured temperature.

Calculations

$$V_m = \frac{M}{D}$$

Where:

 $V_{\rm m}$ = volume of the mold M = mass of water in the mold

D = density of water at the measured temperature

Example

Mass of water in Measure = 7.062 kg (15.53 lb)

Density of water at 23°C (73.4°F) = 997.54 kg/m³ (62.274 lb/ft³)

Table A1 Unit Mass of Water 15°C to 30°C

°C	(°F)	kg/m ³	(lb/ft ³)	°C	(°F)	kg/m ³	(lb/ft ³)
15	(59.0)	999.10	(62.372)	23	(73.4)	997.54	(62.274)
15.6	(60.0)	999.01	(62.366)	23.9	(75.0)	997.32	(62.261)
16	(60.8)	998.94	(62.361)	24	(75.2)	997.29	(62.259)
17	(62.6)	998.77	(62.350)	25	(77.0)	997.03	(62.243)
18	(64.4)	998.60	(62.340)	26	(78.8)	996.77	(62.227)
18.3	(65.0)	998.54	(62.336)	26.7	(80.0)	996.59	(62.216)
19	(66.2)	998.40	(62.328)	27	(80.6)	996.50	(62.209)
20	(68.0)	998.20	(62.315)	28	(82.4)	996.23	(62.192)
21	(69.8)	997.99	(62.302)	29	(84.2)	995.95	(62.175)
21.1	(70.0)	997.97	(62.301)	29.4	(85.0)	995.83	(62.166)
22	(71.6)	997.77	(62.288)	30	(86.0)	995.65	(62.156)

Report

- Measure ID
- Date Standardized
- Temperature of the water
- Volume, $V_{\rm m}$, of the measure

Performance Exam Checklist

FOP for AASHTO T 121 Density (Unit Weight), Yield, and Air Content (Gravimetric) of Concrete

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Mass and volume of empty measure determined?		
Firs	t Layer		
2.	Dampened measure filled approximately one third full, moving a scoop around the perimeter of the measure to evenly distribute the concrete as discharged?		
3.	Layer rodded throughout its depth 25 times, without forcibly striking the bottom of the measure, with hemispherical end of rod, uniformly distributing strokes?		
4.	Perimeter of the measure tapped 10 to 15 times with the mallet after rodding?		
Sec	ond layer		
5.	Measure filled approximately two thirds full, moving a scoop around the perimeter of the measure to evenly distribute the concrete as discharged?		
6.	Layer rodded throughout its depth, just penetrating the previous layer (approximately 25 mm (1 in) 25 times with hemispherical end of rod, uniformly distributing strokes?	. ——	
7.	Perimeter of the measure tapped 10 to 15 times with the mallet after rodding?		
Thir	d layer		
8.	Measure slightly overfilled, moving a scoop around the perimeter of the measure to evenly distribute the concrete as discharged?		
9.	Layer rodded throughout its depth, just penetrating the previous layer (approximately 25 mm (1 in) 25 times with hemispherical end of rod, uniformly distributing strokes?	,	
10.	Perimeter of the measure tapped 10 to 15 times with the mallet after rodding each layer?		
11.	Any excess concrete removed using a trowel or a scoop, or small quantity of concrete added to correct a deficiency, after consolidation of final layer?		
12.	Strike-off plate placed flat on the measure covering approximately \(^2\) of the surface, then sawing action used to withdraw the strike-off plate across the previously covered surface?		
13.	Strike-off plate placed flat on the measure covering approximately \(^2\) of the surface, then sawing action used to advance the plate across the entire measure surface?		
14.	Strike off completed using the inclined edge of the plate creating a smooth surface?		

Procedure Element	Trial 1	Trial 2					
5. All excess concrete cleaned off and mass of full measure determined?							
16. Net mass calculated?							
17. Density calculated correctly?							
First Attempt: Pass Fail Second Attempt: Pass Fail							
Signature of Examiner WAQTC #:							
This checklist is derived, in part, from copyrighted material printed in ACI CP-1, publis American Concrete Institute.	hed by th	ne					

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 No. 4 sieve using a loose volume bucket.

2. Equipment

- a. A mechanical sieve shaker.
- b. Sieves Sieves conforming to the requirements of ASTM E11. 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, \pm 4 hours.
- b. Reduce the sample to testing size per the FOP for AASHTO R 76.
- c. Place the sample in the volume bucket and record the volume as the total volume.
- d. Shake the sample over the 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 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:

$$100 - \frac{\text{(Volume on sieve} \times 100)}{\text{Total Volume}} = \% \text{ passing}$$

Performance Exam Checklist WSDOT T 123 Method of Test for Bark Mulch

Part	ticipant Name		Exa	ım Date				
Pro	cedure Element					Yes	No	
1.	The tester has a copy	y of the curren	t procedure on hand?					
2.	current calibration/v	rerification tags	-	nd if requ	uired, has the			
3.	Bark mulch sample		_					
4.	Five (5) gallon buck	Five (5) gallon bucket calibrated in 1 gal. increments?						
5.	Sample quartered or split and placed in calibrated bucket?							
6.	Volume of sample in bucket recorded as total volume?							
7.	Sample screened in the shaker through 1½ in screen, breaker screens and No. 4 screen?							
8.	Do not over shake to prevent degrading of sample?							
9.	Remove 1½ in screen and damp material in calibrated bucket and record volume as volume on 1½ in screen?							
10.	Place all breaker scr as volume on No. 4		own to No. 4 screen in buck	et and red	cord volume			
11.	All calculations perf	formed correct	ly?					
12.	Report results?							
Firs	et Attempt: Pass	Fail	Second Attempt:	Pass	Fail			
Sig	nature of Examiner							

Comments:

WSDOT Test Method T 125

Determination of Fiber Length Percentages in Wood Strand Mulch

1. Scope

- 1.1. This test method covers the determination of the percentage, by mass, of fiber strands in a wood strand mulch sample meeting the 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.

2. Referenced Documents

- 2.1. AASHTO Standards:
 - M 231 Weighing Devices Used in the Testing of Materials
 - R 76 Reducing Samples of Aggregate to Testing Size

3. Summary of Test Method

3.1. A sample of wood strand mulch is separated into individual fiber strands and the length, width and thickness of each strand is measured. The fiber stands are then separated into two categories; Strands meeting specified requirements and Strands not meeting specified requirements. The percentage of wood fiber strand is then computed and compared to the requirements of the specification. (See Calculation below)

4. Apparatus

- 4.1. *Balance* shall have sufficient capacity, be readable to 0.1 percent of the sample mass, or better, and conform to the requirements of AASHTO M 231 for general-purpose balance required for the principle sample mass being tested.
- 4.2 *Measuring device* capable of reading to the nearest 1/16th inch (can either be one device or two separate devices)

5. Sampling

- 5.1 Split a bale of wood strand mulch into three approximately equal sections. From the interior face of each section obtain a minimum of 150 g of fiber strand, taking care not to damage the material.
- 5.2 Recombine the three 150 g samples and reduce the combined sample to a minimum sample size of 100g, in accordance with FOP for AASHTO R 76, Method B Quartering.

6. Sample Preparation

6.1. Air dry the sample to a Constant Mass as defined in AASHTO T 265.

7. Procedure

- 7.1. Spread the sample on a clean flat surface large enough to permit careful inspection of each strand. Measure the length, width and thickness of each strand in the 100g sample.
- 7.2. Compare the measurements of each strand to the specified requirements and separate the strands into two categories:

Strands meeting specified requirements

Strands not meeting specified requirements

7.3. Determine the total mass of each category.

8. Calculation

- 8.1. Report the following information:
 - 8.1.1. Calculate the percentage of fiber strand meeting the specified requirements to the nearest one percent as follows:

$$P = [(S)/(S+N)] \times 100$$

where:

P = percent of strands meeting the required specifications

S = mass of strands meeting required specifications

N = mass of strands not meeting required specifications

Performance Exam Checklist WSDOT Test Method 125

Par	cicipant Name		Exam Date			
Pro	cedure Element			Yes	No	
1.	The tester has a copy	of the curren	t procedure on hand?			
2.	All equipment is fun current calibration/ve	•	rding to the test procedure, and if required, he s present?	as the		
3.	Sample reduced to co	orrect size?				
4.	Sample dried and co-	oled, if necess	sary?			
5.	Sample properly measured?					
6.	Strands separated int categories?	to "meeting sp	pecification" and "not meeting specifications	"		
7.	Dry mass of each cat	tegory determ	ined to nearest 0.1 g?			
8.	Calculation performe	ed correctly?				
Firs	t Attempt: Pass	Fail	Second Attempt: Pass Fail			
Sig	nature of Examiner					
Con	ments:					

WSDOT Test Method T 126

Determination of Fiber Length Percentages in Hydraulically-Applied Erosion Control Products

1. Scope

- 1.1 This method covers the procedure for measuring the fiber length of a Hydraulically-applied Erosion Control Product (HECP).
- 1.2 HECP is engineered and processed to specific length and width dimensions to facilitate the hydraulic application process. When the correct percentages of fiber lengths and widths exist within the HECP for hydraulic planting, the result will be a properly layered, interlocking mulch to hold seed and moisture.
- 1.3 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. (Warning- HECP's may be dusty, a dust mask is recommended when working with these products.

2. Terminology

- 2.1 Hydraulically applied applied within a slurry, solution, or emulsion to the soil surface as a spray-on or dropped-on application through various means (e.g. nozzle, tower, aerially, etc.) formation of a compound by the combining of water with some other substance.
- 2.2 Hydraulically-applied Erosion Control Product (HECP) A manufactured, degradable, prepackaged fibrous material that is mixed with water and hydraulically applied as slurry, solution, or emulsion to reduce soil erosion and assist in the establishment and growth of vegetation.

3. Apparatus

- 3.1 Balance shall have sufficient capacity, readable to 0.1 percent of the sample mass, or better, and conform to the requirements of AASHTO M 231 for general-purpose balance required for the principle sample mass being tested.
- 3.2 Sieves Meeting the requirements of ASTM E11
- 3.3 Mechanical sieve shaker Meeting the requirements of AASHTO T 27

4. Sample Preparation

Obtain approximate 60 grams of material from a full, sealed bag or bale in the following manner:

Divide the bag or bale into thirds; top, middle and bottom of bag or bale.

Extract approximately 20 grams of material from the center of each portion (or the inside face of each ½ of a bale) taking care not to damage the extracted material.

Recombine the three samples and reduce the sample, in accordance with AASHTO 248 Method B, to approximately 15 grams. Place the sample in a large container and gently separate the compressed fiber.

5. Procedure

- 5.1 Select sieves with suitable openings to furnish data 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.
- 5.2 Place sieves in mechanical shaker and shake for a 5 minutes.
- 5.3 Determine the individual or cumulative mass retained on each sieve and the pan to the nearest 0.1g

6. Calculations

Calculate the percentage of fiber retained on each of the sieves using the following formula:

Percent Retained:

$$CPR = \frac{CMR}{M} \times 100$$

Where:

CPR = Cumulative Percent Retained

CMR= Cumulative Mass Retained

M = Total Sample mass

7. Report

Results shall be reported as follows:

- 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

WSDOT Test Method T 127

Preparation of Leachate Sample for Testing Toxicity of HECP Effluents

1. Scope

This test method outlines the procedure for collecting leachate from a HECP sample.

2. References

2.1 EPA-821-R-02-012 Methods for Measuring Acute Toxicity of Effluents

3. Preparation of the HECP Sample

- 3.1 Lay cheesecloth over clean topsoil
- 3.2 Apply the HECP to the cheesecloth at the following coverage:
 - 3.2.1 HECP Long-Term or Moderate-Term Mulch 3,500 lbs per acre in two lifts with no more than 2000# per acre in any one lift.
 - 3.2.2 HECP Short-Term Mulch 2000# per acre minimum, or manufacturer's recommended rate may be used.
- 3.3 Allow material to cure for 48 hours
- 3.4 Pull cheesecloth up with mat of HECP, brush off any soil
- 3.5 Cut HECP into squares and provide 1.02 pounds of HECP to the laboratory.

4. Preparation of the Leachate (done by the laboratory)

- 4.1 Allow hydromulch product to soak in water for one hour
- 4.2 Remove hydromulch sheet from water
- 4.3 Filter water through a 60 micron filter water may be pre-filtered through a 35 mesh and then a 120 mesh before using the 230 mesh/63 micron filter.

Note: Leachate not immediately used for testing should be stored at 4° C in the dark until use.

4.4 Test leachate according to EPA-821-R-02-012 Methods for Measuring Acute Toxicity of Effluents.

WSDOT SOP 128

Sampling for Aggregate Source Approval

1. Scope

This method describes the procedure for sampling pits and quarries for Aggregate Source Approval (ASA).

2. Significance and Use

There are two methods for initiating the process for an Aggregate Source Approval:

- a. The source owner request approval, pays for the sampling and testing, and coordinates this through the State Materials Laboratory who coordinates with the Regions. Sample is obtained by the Region Independent Assurance Inspector (IAI) or a delegate of the Region Materials Engineer.
- b. The aggregate source is sampled and tested as part of a WSDOT project, in which case the WSDOT project pays for the sampling and testing costs which may or may not be coordinated with the ASA process at the State Materials Laboratory. Sample is obtained by the IAI or a delegate of the Region Materials Engineer.

3. Safety

All WSDOT employee required to sample from a pit or quarry will contact the pit/quarry owner or their designated representative prior to arrival at the site and arrange for an escort into the sampling site.

All WSDOT employees will be accompanied by the pit/quarry owner or their representative during the sampling process.

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.

4. Sampling

All samples will be obtained in accordance with WSDOT FOP for AASHTO T 2.

Stockpiles produced for ASA sampling must contain a minimum of 10 tons of material. The material in the stockpile shall be of the same quality as the final product.

Sampling location and size of sample is listed in Table 1.

Aggregate Type	Sampling Site	Size of Sample in lbs	Notes
Concrete Coarse	Stockpile	50-100	Material must be clean and washed
Concrete Fine	Stockpile	30-40	Material must be clean and washed
Crushed Surfacing /Mineral Aggregate	Stockpile	80-100	For quality tests on crushed materials submit approximately 80 lbs of 1¼" minus material. Samples obtained for quarry spalls may not be used for quality tests for crushed materials.
Quarry Spalls	Face of pit , transport unit or stockpile	50-80	No rock larger than 4" in diameter.
All other Aggregate Types	Face of pit , transport unit or stockpile	50-80	No rock larger than 4" in diameter.

Table 1

6. Report

The Regional Materials Engineer's (RME) representative will record the following information in an Inspector's Daily Report (IDR) DOT Form 422-004A:

- Name of Source Owner's Representative accompanying the RME representative during sampling process.
- Time and Date of sampling
- Location where the sample is taken (stockpile/pit/face)
- Amount of sample (pounds and number of bags)
- Any concerns or specific request the Owner's representative may have.

The RME's representative shall take pictures of the following items; a wide view of the mining operation, the sampling location in the pit or quarry, a close-up of the material in the stockpile being sampled (when applicable), and a close-up of the material sampled.

The IDR information and pictures will be e-mailed to the State ASA Engineer.

WSDOT Errata to FOP for AASHTO T 152

Air Content of Freshly Mixed Concrete by the Pressure Method

WAQTC FOP for AASHTO T 152 has been adopted by WSDOT with the following changes:

Apparatus

• Vibrator:

Replace with details below:

The vibrator frequency shall be at least 9000 vibrations per minute, 19 to 38 mm (0.75 to 1.50 in) in diameter, at least 75 mm (3 in) longer than the section being vibrated for use with low slump concrete

FOP for AASHTO T 152

Air Content of Freshly Mixed Concrete by the Pressure Method

Scope

This procedure covers determination of the air content in freshly mixed Portland Cement Concrete containing dense aggregates in accordance with AASHTO T 152-17, Type B meter. It is not for use with lightweight or highly porous aggregates. This procedure includes standardization of the Type B air meter gauge, Annex A.

Warning—Fresh Hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.

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 standardization only)
- Tamping rod: 16 mm (% in) diameter and 400 mm (16 in) to 600 mm (24 in) long, having a hemispherical tip the same diameter as the rod. (Hemispherical means "half a sphere"; the tip is rounded like half of a ball.)
- Vibrator: 9000 vibrations per minute, 19 to 38 mm (0.75 to 1.50 in) in diameter, at least 75 mm (3 in) longer than the section being vibrated for use with low slump concrete
- Scoop: a receptacle of appropriate size so that each representative increment of the concrete sample can be placed in the container without spillage.
- Container for water: rubber syringe (may also be a squeeze bottle)
- Strike-off bar: Approximately 300 mm x 22 mm x 3 mm (12 in x \(^3\)4 in × \(^1\)8 in)
- Strike-off plate: A flat rectangular metal plate at least 6 mm (¼ in) thick or a glass or acrylic plate at least 12 mm (½ in) thick, with a length and width at least 50 mm (2 in) 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 1.5 mm (¼ in).

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 0.57 \pm 0.23 kg (1.25 \pm 0.5 lb)

Procedure Selection

There are two methods of consolidating the concrete – rodding and vibration. If the slump is greater than 75 mm (3 in), consolidation is by rodding. When the slump is 25 to 75 mm (1 to 3 in), internal vibration or rodding 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 concrete with slumps less than 25 mm (1 in), consolidate the sample by internal vibration. Do not consolidate self-consolidating concrete (SCC).

Procedure - Rodding

- Obtain the sample in accordance with the FOP for WAQTC TM 2. If the concrete mixture contains aggregate retained on the 37.5mm (1½ in) sieve, the aggregate must be removed in accordance with the Wet Sieving portion of the FOP for WAQTC TM 2.
 - **Note 2:** Testing shall begin within five minutes of obtaining the sample.
- 2. Dampen the inside of the air meter measure and place on a firm level surface.
- 3. Use the scoop to fill the measure approximately ½ full with concrete. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 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 around the perimeter of the measure smartly 10 to 15 times with the mallet to close voids and release trapped air.
- 6. Add the second layer, filling the measure about \% full. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 7. Consolidate this layer with 25 strokes of the tamping rod, penetrating about 25 mm (1 in) into the bottom layer.
- 8. Tap around the perimeter of the measure smartly 10 to 15 times with the mallet.
- 9. Add the final layer, slightly overfilling the measure. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 10. Consolidate this layer with 25 strokes of the tamping rod, penetrating about 25 mm (1 in) into the second layer.
- 11. Tap around the perimeter of the measure smartly 10 to 15 times with the mallet.
 - **Note 3:** The measure should be slightly over full, about 3 mm (1/8 in) above the rim. If there is a great excess of concrete, remove a portion with the trowel or scoop. If the measure 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 measure just full. The surface should be smooth and free of voids.
- 13. Clean the top flange of the measure 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 measure.
- 16. Inject water through a petcock on the cover until water emerges from the petcock on the other side.
- 17. Incline slightly and gently rock the air meter 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 determined for the gauge. 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. Close the air bleeder valve.
- 20. Close both petcocks.
- 21. Open the main air valve.
- 22. Tap around the perimeter of the measure smartly with the mallet.
- 23. With the main air valve open, lightly tap the gauge to settle the needle, and then read the air content to the nearest 0.1 percent.
- 24. Release or close the main air valve.
- 25. Open both petcocks to release pressure, remove the concrete, and thoroughly clean the cover and measure 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 37.5 mm (1½ in) or larger is present, aggregate must be removed in accordance with the Wet Sieving portion of the FOP for WAQTC TM 2.
- 2. Dampen the inside of the air meter measure and place on a firm level surface.
- Use the scoop to fill the measure approximately 1/2 full with concrete. Evenly
 distribute the concrete in a circular motion around the inner perimeter of the
 measure.
- 4. Insert the vibrator at three different points. Do not let the vibrator touch the bottom or side of the measure. Remove the vibrator slowly, so that no air pockets are left in the material. 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. Use the scoop to fill the measure a bit over full. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 6. Insert the vibrator as in Step 4. Do not let the vibrator touch the side of the measure, and penetrate the first layer approximately 25 mm (1 in). Remove the vibrator slowly, so that no air pockets are left in the material. 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.
- 7. Return to Step 12 of the rodding procedure and continue.

Procedure – Self Consolidating Concrete

- 1. Obtain the sample in accordance with the FOP for WAQTC TM 2.
- 2. Dampen the inside of the air meter measure and place on a firm level surface.
- 3. Use the scoop to slightly overfill the measure. Evenly distribute the concrete in a circular motion around the inner perimeter of the measure.
- 4. Complete Steps 12 thru 26 of the rodding procedure.

Report

- Results on forms approved by the agency
- Sample ID
- Percent of air to the nearest 0.1 percent.
- Some agencies require an aggregate correction factor in order to determine total percent of entrained air.

Total percent entrained air = Gauge reading – aggregate correction factor from mix design

(See AASHTO T 152 for more information.)

ANNEX A—Standardization of Air Meter Gauge

Standardization is a critical step to ensure accurate test results when using this apparatus. Failure to perform the standardization procedures as described below will produce inaccurate or unreliable test results.

Standardization shall be performed at a minimum of once every three months. Record the date of the standardization, the standardization results, and the name of the technician performing the standardization in the log book kept with each air meter.

There are two methods for standardizing the air meter, mass or volume, both are covered below.

- 1. Screw the short piece of straight tubing into the threaded petcock hole on the underside of the cover.
- 2. Determine and record the mass of the dry, empty air meter measure and cover assembly (mass method only).
- 3. Fill the measure nearly full with water.
- 4. Clamp the cover on the measure with the tube extending down into the water. Mark the petcock with the tube attached for future reference.
- 5. 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.
- 6. Wipe off the air meter measure and cover assembly; determine and record the mass of the filled unit (mass method only).
- 7. 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.
- 8. Close both petcocks and immediately open the main air valve exhausting air into the measure. 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.
- 9. Determine which petcock has the straight tube attached to it. Attach the curved tube to external portion of the same petcock.
- 10. 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 measure and drain the water in the curved tube back into the measure. 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 standardization vessel is level full.

Note A1: Many air meters are supplied with a standardization vessel(s) of known volume that are used for this purpose. Standardization vessel must be protected from crushing or denting. If an external standardization 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.

- 11. 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.
- 12. Close both petcocks and immediately open the main air valve exhausting air into the measure. Wait a few seconds until the meter needle is stabilized. The gauge should now read 5.0 ± 0.1 percent. If the gauge is outside that range, the meter needs adjustment. The adjustment could involve adjusting the starting point so that the gauge reads 5.0 ± 0.1 percent when this standardization is run, or could involve moving the gauge needle to read 5.0 percent. Any adjustment should comply with the manufacturer's recommendations.
- 13. 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.
- 14. If an internal standardization vessel is used, follow steps 1 through 8 to set initial reading.
- 15. Release pressure from the measure and remove cover. Place the internal standardization vessel into the measure. This will displace 5 percent of the water in the measure. (See AASHTO T 152 for more information on internal standardization vessels.)
- 16. Place the cover back on the measure and add water through the petcock until all the air has been expelled.
- 17. 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.
- 18. Close both petcocks and immediately open the main air valve exhausting air into the measure. Wait a few seconds until the meter needle stabilizes. The gauge should now read 5 percent.
- 19. Remove the extension tubing from threaded petcock hole in the underside of the cover before starting the test procedure.

Report

- · Air Meter ID
- Date Standardized
- Initial Pressure (IP)

Performance Exam Checklist

FOP for AASHTO T 152

Air Content of Freshly Mixed Concrete by the Pressure Method

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Representative sample selected?		
Firs	t Layer		
2.	Dampened measure filled approximately one third full, moving a scoop around the perimeter of the measure to evenly distribute the concrete as discharged?		
3.	Layer rodded throughout its depth 25 times, without forcibly striking the bottom of the measure, with hemispherical end of rod, uniformly distributing strokes?		
4.	Perimeter of the measure tapped 10 to 15 times with the mallet after rodding?		
Sec	ond layer		
5.	Measure filled approximately two thirds full, moving a scoop around the perimeter of the measure to evenly distribute the concrete as discharged?		
6.	Layer rodded throughout its depth, just penetrating the previous layer (approximately 25 mm (1 in) 25 times with hemispherical end of rod, uniformly distributing strokes?		
7.	Perimeter of the measure tapped 10 to 15 times with the mallet after rodding?		
Thir	d layer		
8.	Measure slightly overfilled, moving a scoop around the perimeter of the measure to evenly distribute the concrete as discharged?		
9.	Layer rodded throughout its depth, just penetrating the previous layer (approximately 25 mm (1 in)) 25 times with hemispherical end of rod, uniformly distributing strokes?		
10.	Perimeter of the measure tapped 10 to 15 times with the mallet after rodding each layer?		
11.	Concrete struck off level with top of the measure using the bar or strike-off plate and rim cleaned off?		
12.	Top flange of base cleaned?		
Usir	ng a Type B Meter:		
13.	Both petcocks open?		
14.	Air valve closed between air chamber and the measure?		
15.	Inside of cover cleaned and moistened before clamping to base?		
16.	Water injected through petcock until it flows out the other petcock?		
17.	Water injection into the petcock continued while jarring and or rocking the meter to insure all air is expelled?		

Pro	cedure Element	Trial 1	Trial 2
18.	Air pumped up to just past initial pressure line?		
19.	A few seconds allowed for the compressed air to stabilize?		
20.	Gauge adjusted to the initial pressure?		
21.	Both petcocks closed?		
22.	Air valve opened between chamber and measure?		
23.	The outside of measure tapped smartly with the mallet?		
24.	With the main air valve open, gauge lightly tapped and air percentage read to the nearest 0.1 percent?		
25.	Air valve released or closed and then petcocks opened to release pressure before removing the cover?		
26.	Aggregate correction factor applied if required?		
27.	Air content recorded to 0.1 percent?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
	checklist is derived, in part, from copyrighted material printed in ACI CP-1, publis rican Concrete Institute.	hed by th	ne

Comments:

WSDOT Errata to FOP for AASHTO T 166

Bulk Specific Gravity (G_{mb}) of Compacted Asphalt Mixtures Using Saturated Surface-Dry Specimens

WAQTC FOP for AASHTO T 166 has been adopted by WSDOT with the following changes:

Test Specimens

Include items below:

Transportation of warm laboratory molded specimens is not recommended before they have cooled to room temperature. If however, a specimen must be transported prior to reaching room temperature the following guidelines should be used to transport the specimen:

- 1. Place the specimen in a container that has a flat bottom surface to prevent deformation of the bottom of the specimen. *Note*: A flat piece of wood, rigid aluminum or reinforced cardboard may be used to create a flat surface in an HMA sample box.
- 2. Make sure the specimen is not deformed in handling.
- 3. Do not stack anything on top of the specimen container.
- 4. Transport the container in the cab of the vehicle or secure it in the vehicle bed to prevent movement during transit.

Procedure - Method A (Suspension)

Include step h. below:

1.	Dry the specimen to constant mass, if required.
	a.
	b.
	C.

e. f.

d.

g.

h. As an alternative to oven drying to constant mass, drying the sample according to AASHTO R 79 may be used.

Replace step 2 with below:

2. Cool the specimen in air for a minimum of 15 hours and a maximum of 24 hours to 25 ± 5 °C (77 \pm 9°F), and determine and record the dry mass to the nearest 0.1 g. Designate this mass as "A."

FOP for AASHTO T 166

Bulk Specific Gravity (G_{mb}) of Compacted Asphalt Mixtures Using Saturated Surface-Dry Specimens

Scope

This procedure covers the determination of bulk specific gravity (G_{mb}) of compacted asphalt mixtures using three methods – A, B, and C – in accordance with AASHTO T 166-16. This FOP is for use on specimens not having open or interconnecting voids or absorbing more than 2.00 percent water by volume, or both. When specimens have open or interconnecting voids or absorbing more than 2.00 percent water by volume, or both, AASHTO T 275 or AASHTO T 331 should be performed.

Overview

Method A: Suspension

• Method B: Volumeter

• Method C: Rapid test for A or B

Test Specimens

Test specimens may be either laboratory-molded or from asphalt mixture pavement. For specimens it is recommended that the diameter be equal to four times the maximum size of the aggregate and the thickness be at least one and one half times the maximum size.

Test specimens from asphalt mixture pavement will be sampled according to AASHTO R 67.

Terminology

Constant Mass: The state at which a mass does not change more than a given percent, after additional drying for a defined time interval, at a required temperature.

Apparatus - Method A (Suspension)

Balance or scale: 5 kg capacity, readable to 0.1 g, and fitted with a suitable suspension apparatus and holder to permit weighing the specimen while suspended in water, conforming to AASHTO M 231.

- Suspension apparatus: Wire of the smallest practical size and constructed to permit the container to be fully immersed.
- Water bath: For immersing the specimen in water while suspended under the balance or scale, and equipped with an overflow outlet for maintaining a constant water level.
- Towel: Damp cloth towel used for surface drying specimens.
- Oven: Capable of maintaining a temperature of 110 ± 5 °C (230 ± 9 °F) for drying the specimens to a constant mass.
- Pan: Pan or other suitable container of known mass, large enough to hold a sample for drying in oven.
- Thermometer: Having a range of 19 to 27°C (66 to 80°F), graduated in 0.1°C (0.2°F) subdivisions.

Procedure - Method A (Suspension)

Recently molded laboratory samples that have not been exposed to moisture do not need drying.

- 1. Dry the specimen to constant mass, if required.
 - a. Initially dry overnight at $52 \pm 3^{\circ}$ C ($125 \pm 5^{\circ}$ F).
 - b. Determine and record the mass of the specimen (M_p).
 - c. Return the specimen to the oven for at least 2 hours.
 - d. Determine and record the mass of the specimen (M_n).
 - e. Determine percent change by subtracting the new mass determination (M_n) from the previous mass determination (M_p) divide by the previous mass determination (M_p) multiply by 100.
 - f. Continue drying until there is less than 0.05 percent change in specimen mass after 2-hour drying intervals (constant mass).
 - g. Constant mass has been achieved, sample is defined as dry.
 - **Note 1:** To expedite the procedure, steps 1 and 2 may be performed last. To further expedite the process, see Method C.
- 2. Cool the specimen in air to $25 \pm 5^{\circ}$ C (77 $\pm 9^{\circ}$ F), and determine and record the dry mass to the nearest 0.1 g. Designate this mass as "A."
- 3. Fill the water bath to overflow level with water at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F) and allow the water to stabilize.
- 4. Zero or tare the balance with the immersion apparatus attached, ensuring that the device is not touching the sides or the bottom of the water bath.
- 5. Immerse the specimen shaking to remove the air bubbles. Place the specimen on its side in the suspension apparatus. Leave it immersed for 4 ± 1 minutes.
- 6. Determine and record the submerged weight to the nearest 0.1 g. Designate this submerged weight as "C."
- 7. Remove the sample from the water and quickly surface dry with a damp cloth towel within 5 seconds.
- 8. Zero or tare the balance.
- 9. Immediately determine and record the mass of the SSD specimen to nearest 0.1 g. Designate this mass as "B." Any water that seeps from the specimen during the mass determination is considered part of the saturated specimen. Do not to exceed 15 seconds performing Steps 7 through 9.

Calculations - Method A (Suspension)

Constant Mass:

Calculate constant mass using the following formula:

$$\%Change = \frac{M_p - M_n}{M_p} \times 100$$

Where:

 M_p = previous mass measurement, g M_n = new mass measurement, g

Bulk specific gravity (G_{mb}) and percent water absorbed:

$$G_{mb} = \frac{A}{B - C}$$

Percent Water Absorbed (by volume) =
$$\frac{B-A}{B-C} \times 100$$

Where:

A = Mass of dry specimen in air, g
B = Mass of SSD specimen in air, g

C = Weight of specimen in water at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F), g

Example:

$$G_{mb} = \frac{4833.6 \ g}{4842.4 \ g - 2881.3 \ g} = 2.465$$

% Water Absorbed (by volume) =
$$\frac{4842.4 g - 4833.6 g}{4842.4 g - 2881.3 g} \times 100 = 0.45\%$$

Apparatus - Method B (Volumeter)

- Balance or scale: 5 kg capacity, readable to 0.1 g and conforming to AASHTO M 231.
- Water bath: Thermostatically controlled to 25 ± 0.5°C (77 ± 0.9°F).
- Thermometer: Range of 19 to 27°C (66 to 80°F), and graduated in 0.1°C (0.2°F) subdivisions.
- Volumeter: Calibrated to 1200 mL or appropriate capacity for test sample and having a tapered lid with a capillary bore.
- Oven: Capable of maintaining a temperature of 110 ± 5 °C (230 ± 9 °F) for drying the specimens to a constant mass.
- Pan: Pan or other suitable container of known mass, large enough to hold a sample for drying in oven.
- Towel: Damp cloth towel used for surface drying specimens.

Procedure - Method B (Volumeter)

Recently molded laboratory samples that have not been exposed to moisture do not need drying.

- 1. Dry the specimen to constant mass, if required.
 - a. Initially dry overnight at 52 ± 3 °C (125 ± 5 °F).
 - b. Determine and record the mass of the specimen (M_p).
 - c. Return the specimen to the oven for at least 2 hours.
 - d. Determine and record the mass of the specimen (M_p).
 - e. Determine percent change by subtracting the new mass determination (M_n) from the previous mass determination (M_p) divide by the previous mass determination (M_p) multiply by 100.
 - f. Continue drying until there is less than 0.05 percent change in specimen mass after 2-hour drying intervals (constant mass).
 - g. Constant mass has been achieved, sample is defined as dry.
 - **Note 1:** To expedite the procedure, steps 1 and 2 may be performed last. To further expedite the process, see Method C.
- 2. Cool the specimen in air to $25 \pm 5^{\circ}$ C (77 $\pm 9^{\circ}$ F), and determine and record the dry mass to the nearest 0.1 g. Designate this mass as "A."
- 3. Immerse the specimen in the temperature-controlled water bath for at least 10 minutes.
- 4. Fill the volumeter with distilled water at $25 \pm 1^{\circ}$ C ($77 \pm 1.8^{\circ}$ F) making sure some water escapes through the capillary bore of the tapered lid. Wipe the volumeter dry. Determine the mass of the volumeter to the nearest 0.1 g. Designate this mass as "D."
- 5. At the end of the ten minute period, remove the specimen from the water bath and quickly surface dry with a damp cloth towel within 5 seconds.
- 6. Immediately determine and record the mass of the SSD specimen to the nearest 0.1g.
- 7. Designate this mass as "B." Any water that seeps from the specimen during the mass determination is considered part of the saturated specimen.
- 8. Place the specimen in the volumeter and let stand 60 seconds.
- 9. Bring the temperature of the water to $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F) and cover the volumeter, making sure some water escapes through the capillary bore of the tapered lid.
- 10. Wipe the volumeter dry.
- 11. Determine and record the mass of the volumeter and specimen to the nearest 0.1 g. Designate this mass as "E."
 - **Note 2:** Method B is not acceptable for use with specimens that have more than 6 percent air voids.

Calculations - Method B (Volumeter)

Constant Mass:

Calculate constant mass using the following formula:

$$\%Change = \frac{M_p - M_n}{M_n} \times 100$$

Where:

 M_p = previous mass measurement, g M_n = new mass measurement, g

Bulk specific gravity (Gmb) and percent water absorbed:

$$G_{mb} = \frac{A}{B + D - E}$$

Percent Water Absorbed (by volume) =
$$\frac{B-A}{B+D-E} \times 100$$

Where:

A = Mass of dry specimen in air, g B = Mass of SSD specimen in air, g D = Mass of volumeter filled with water at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F), g

E = Mass of volumeter filled with specimen and water, g

Example:

$$G_{mb} = \frac{4833.6 \ g}{4842.4 \ g + 2924.4 \ g - 5806.0 \ g} = 2.465$$

% Water Absorbed (by volume) =
$$\frac{4842.4 \ g - 4833.6 \ g}{4842.4 \ g + 2924.4 \ g - 5806.0 \ g} \times 100 = 0.45\%$$

Method C (Rapid Test for Method A or B)

See Methods A or B.

Note 3: This procedure can be used for specimens that are not required to be saved and contain substantial amounts of moisture. Cores can be tested the same day as obtained by this method.

Procedure - Method C (Rapid Test for Method A or B)

- 1. Start on Step 3 of Method A or B, and complete that procedure, then determine dry mass, "A," as follows.
- 2. Determine and record mass of a large, flat-bottom container.
- 3. Place the specimen in the container.
- 4. Place in an oven at a minimum of 105°C (221°F). Do not exceed the Job Mix Formula mixing temperature.
- 5. Dry until the specimen can be easily separated into fine aggregate particles that are not larger than 6.3 mm ($\frac{1}{4}$ in).
- 6. Determine and record the mass of the specimen (M_p) .
- 7. Return the specimen to the oven for at least 2 hours.
- 8. Determine and record the mass of the specimen (M_p) .
- 9. Determine percent change by subtracting the new mass determination (M_n) from the previous mass determination (M_p) divide by the previous mass determination (M_n) multiply by 100.
- 10. Continue drying until there is less than 0.05 percent change in specimen mass after 2-hour drying intervals (constant mass).
- 11. Constant mass has been achieved, sample is defined as dry.
- 12. Cool in air to $25 \pm 5^{\circ}$ C (77 $\pm 9^{\circ}$ F).
- 13. Determine and record the mass of the container and dry specimen to the nearest 0.1 g.
- 14. Determine and record the mass of the dry specimen to the nearest 0.1 g by subtracting the mass of the container from the mass determined in Step 13. Designate this mass as "A."

Calculations - Method C (Rapid Test for Method A or B)

Complete the calculations as outlined in Methods A or B, as appropriate.

Report

- Results on forms approved by the agency
- Sample ID
- G_{mb} to 0.001
- Absorption to 0.01 percent
- Method performed.

Performance Exam Checklist

FOP for AASHTO T 166

Bulk Specific Gravity (G_{mb}) of Compacted Asphalt Mixtures Using Saturated Surface-Dry Specimens

Par	Participant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
Me	thod A:		
1.	Mass of dry sample in air determined.		
	a. Dried overnight at $52 \pm 3^{\circ}$ C ($125 \pm 5^{\circ}$ F) and at successive 2-hour intervals to constant mass?		
	b. Cooled in air to 25 ± 5 °C (77 ± 9°F)?		
	c. Dry mass determined to 0.1g?		
2.	Water at the overflow?		
3.	Balance zeroed?		
4.	Immersed weight determined.		
	a. Water at 25 ± 1°C (77 ± 1.8°F)?		
	b. Immersed, shaken, on side, for 4 ± 1 minutes?		
	c. Immersed weight determined to 0.1g?		
5.	Sample rapidly surface dried with damp towel and saturated surface dry (SSD) mass determined to 0.1 g (entire operation performed within 15 seconds)?		
6.	G _{mb} calculated to 0.001?		
7.	Absorption calculated to 0.01 percent		
Me	thod B:		
1.	Specimen dried, cooled, and mass determined as in Method A?		
2.	Saturated surface dry (SSD) mass determined to 0.1g. a. Immersed at least 10 minutes at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F)?		
	b. Sample rapidly dried with damp towel?		
	c. Specimen mass determined to 0.1 g?		
	d. Any water that seeps from specimen included in mass?		
3.	Mass of volumeter filled with distilled water at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F) determined?		
4.	SSD specimen placed into volumeter and let stand for 1 minute?		
5.	Temperature of water brought to $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F) and volumeter covered, allowing some water to escape through the capillary bore of the tapered lid?		

Pro	cedure Element	Trial 1	Trial 2
6.	Volumeter wiped dry, and mass of volumeter and contents determined?		
7.	G _{mb} calculated to 0.001?		
8.	Absorption calculated to 0.01 percent?		
Met	hod C/A:		
1.	Immersed weight determined.		
	a. Water at $25 \pm 1^{\circ}$ C (77 ± 1.8°F)?		
	b. Immersed, shaken, on side, for 4 ± 1 minutes?		
	c. Immersed weight determined to 0.1 g?		
2.	Sample rapidly surface dried with damp cloth (within 5 seconds)?		
3.	Saturated surface dry mass determined to 0.1 g?		
4.	Dry mass determined by:		
	a. Heating in oven at a minimum of 105°C (221°F)?		
	b. Breaking down to 6.3 mm (¼ in) particles?		
	c. Drying in oven to constant mass (change less than 0.05 percent in 2 hours of additional drying)?		
	d. Cooled in air to 25 ± 5 °C (77 \pm 9°F) and mass determined to 0.1 g?		
5.	G _{mb} calculated to 0.001?		
6.	Absorption calculated to 0.01?		
Met	hod C/B:		
1.	Saturated surface dry (SSD) mass determined to 0.1g. a. Immersed at least 10 minutes at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F)?		
	b. Sample rapidly dried with damp towel (within 5 seconds)?		
	c. Specimen mass determined to 0.1g?		
	d. Any water that seeps from specimen included in mass?		
2.	Mass of volumeter filled with distilled water at $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F) determined to 0.1 g?		
3.	SSD specimen placed into volumeter and let stand for 1 minute?		
4.	Temperature of water brought to $25 \pm 1^{\circ}$ C (77 $\pm 1.8^{\circ}$ F) and volumeter covered, allowing some water to escape through the capillary pore of the tapered lid?		
5.	Volumeter wiped dry, and mass of volumeter and contents determined to 0.1 g?		

Procedure Element		Trial 1	Trial 2
6.	Dry mass determined by:		
	a. Warming in oven at a minimum of 105°C (221°F)?		
	b. Breaking down to 6.3 mm (¼ in) particles?		
	c. Drying in oven to constant mass (change less than 0.05 percent in 2 hours of additional drying)?		
	d. Cooled in air to 25 ± 5 °C (77 ± 9 °F) and mass determined to 0.1 g?		·
7.	G _{mb} calculated to 0.001?		
8.	Absorption calculated to 0.01 percent?		
First	Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ture of Examiner WAQTC #:		

Comments:

WSDOT Errata to FOP for AASHTO T 168

Sampling of Bituminous Paving Mixtures

WAQTC FOP for AASHTO T 168 has been adopted by WSDOT with the following changes:

Sample Size

For Acceptance sampling and testing only: WSDOT requires a minimum of two times the amount required for testing. This should be 60 lbs.

For Acceptance and Conformation sampling and testing: WSDOT requires a minimum of four times the amount required for testing. This should be approximately 120 lbs. (See WSDOT *Construction Manual Section 9-3.7* for Conformation sampling frequency)

Sampling

General

Include the steps below:

- Immediately upon obtaining a sample, using a verified thermometer, check and record temperature of the sample.
- The material shall be tested to determine variations. The supplier/contractor shall sample the HMA mixture in the presence of the Project Engineer.
 The supplier/contractor shall provide one of the following for safe and representative sampling:
 - a. A mechanical sampling device installed between the discharge of the silo and the truck transport that is approved by the Regional Materials Engineer.
 - b. Platforms or devices to enable sampling from the truck transport without entering the truck transport for sampling HMA.

Attached Sampling Devices

Sampling from Roadway Prior to Compaction (Plate Method)

Method 1 - Obtaining a Sample on Untreated Base: - *Method not recognized by WSDOT.*

Method 2 - Obtaining a Sample on Asphalt Surface: - Method not recognized by WSDOT.

FOP for AASHTO T 168

Sampling of Bituminous Paving Mixtures

Scope

This procedure covers the sampling of bituminous paving mixtures from HMA plants, haul units, and roadways in accordance with AASHTO T 168-03. Sampling is as important as testing, and every precaution must be taken to obtain a truly representative sample.

Apparatus

- Shovel
- Sample containers: such as cardboard boxes, metal cans, stainless steel bowls, or other agency-approved containers
- Scoops, trowels, or other equipment to obtain mix
- Sampling plate: Thick metal plate, minimum 8 gauge, sized to accommodate sample requirements, 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 in diameter should be provided in each corner.
- Cookie cutter sampling device: Formed steel angle with two 100 mm by 150 mm by 9 mm (4 in by 6 in by 3/8 in) handles, sized to accommodate sample requirements.
 Minimum 2 in smaller than the sampling plate when used together.
 - **Example:** Sampling plate 380 mm (15 in) square and a cookie cutter sampling device 330 mm (13 in) square.
- Mechanical sampling device

Sample Size

Sample size depends on the test methods specified by the agency for acceptance. Check agency requirement for the size required.

Sampling

General

- The material shall be tested to determine variations. The supplier/contractor shall provide equipment for safe and appropriate sampling, including sampling devices on plants when required.
- For dense graded mixture samples use cardboard boxes, stainless steel bowls or other agency-approved containers.
- For hot open graded mixture samples use 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.

Attached Sampling Devices

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.

- 1. Lightly coat the container attached to the sampling device with an agencyapproved release agent or preheat it, or both, to approximately the same discharge temperature of the mix.
- 2. Pass the container twice through the material perpendicularly without overfilling the container.
- 3. Repeat until proper sample size has been obtained.
- 4. Transfer the HMA to an agency-approved container without loss of material.

Sampling from Haul Units

- 1. Visually divide the haul unit into approximately four equal quadrants.
- 2. Identify one sampling location in each quadrant.
- 3. Dig down and remove approximately 0.3 m (1 ft) of material to avoid surface segregation. Obtain each increment from below this level.
- 4. Combine the increments to form a sample of the required size.

Sampling from Roadway Prior to Compaction (Plate Method)

Plate method using the "cookie cutter" sampling device.

There are two conditions that will be encountered when sampling hot mix asphalt (HMA) from the roadway prior to compaction. The two conditions are:

- Laying HMA on grade or untreated base material requires Method 1.
- Laying HMA on existing asphalt or laying a second lift of HMA requires Method 2.

SAFETY

Sampling is performed behind the paving machine and in front of the breakdown roller. For safety, the roller must remain at least 3 m (10 ft) behind the sampling operation until the sample has been taken and the hole filled with loose HMA.

Method 1 requires a plate to be placed in the roadway in front of the paving operation and therefore there is always concern with moving, operating equipment. It is safest to stop the paving train while a plate is installed in front of the paver. When this is not possible the following safety rules must be followed.

1. The plate placing operation must be at least 3 m (10 ft) in front of the paver or pickup device. The technician placing the plate must have eye contact and communication with the paving machine operator. If eye contact cannot be maintained at all time, a third person must be present to provide communication between the operator and the technician.

2. No technician is to be between the asphalt supply trucks and the paving machine. The exception to this rule is if the supply truck is moving forward creating a windrow, in which case the technician must be at least 3 m (10 ft) behind the truck.

If at any time the Engineer feels that the sampling technique is creating an unsafe condition, the operation is to be halted until it is made safe or the paving operation will be stopped while the plate is being placed.

Method 1 - Obtaining a Sample on Untreated Base:

- 1. Following the safety rules detailed above, the technician is to:
 - a. Smooth out a location in front of the paver at least 0.5 m (2 ft) inside the edge of the mat.
 - b. Lay the plate down diagonally with the direction of travel, keeping it flat and tight to the base with the lead corner facing the paving machine.
- 2. Secure the plate in place by driving a nail through the hole in the lead corner of the plate.
- 3. Pull the wire, attached to the outside corner of the plate, taut past the edge of the HMA mat and secure with a nail.
- 4. Let the paving operation proceed over the plate and wire. Immediately proceed with the sampling.
- 5. Using the exposed end of the wire, pull the wire up through the fresh HMA to locate the corner of the plate. Place the "cookie cutter" sample device, just inside the end of the wire; align the cutter over the plate. Press "cookie cutter" device down through the HMA to the plate.
- 6. Using a small square tipped shovel or scoop, or both, carefully remove all the HMA from inside of the cutter and place in a sample container. 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.
- 7. Remove the sample cutter and the plate from the roadway. The hole made from the sampling must be filled by the contractor with loose HMA.

Method 2 - Obtaining a Sample on Asphalt Surface:

- 1. After the paving machine has passed the sampling point, immediately place the "cookie cutter" sampling device on the location to be sampled. Push the cutter down through the HMA until it is flat against the underlying asphalt mat.
- 2. Using a small square tipped shovel or scoop, or both, carefully remove all the HMA from inside of the cutter and place in a sample container. The hole made from the sampling must be filled by the contractor with loose HMA.

Identification and Shipping

- 1. Identify sample containers as required by the agency.
- 2. Ship samples in containers that will prevent loss, contamination, or damage.

Report

- On forms approved by the agency
- Sample ID
- Date
- Time
- Location
- Quantity represented

Performance Exam Checklist FOP For AASHTO T 168 SAMPLING BITUMINOUS PAVING MIXTURES

Par	ticipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Was sample taken with an attached sampling device correctly?		
	a. Container coated or preheated or both?		
	b. Sampling device passed through stream twice perpendicular to material?		
	c. Sampling device not over filled?		
2.	Samples from truck transports taken from four quadrants at required depth of 300 mm (12 in)?		
3.	Samples from roadway taken correctly with plate(s).		
	a. When on untreated base plate placed well in front of paver?		
	b. Wire pulled to locate plate corner?		
	c. Cookie cutter placed on asphalt and pushed through to plate?		
	d. All material removed from inside the cutter?		
4.	Sample placed in appropriate container.		
5.	Sample size meets agency requirements?		
6.	Sample identified as required?		
Firs	st Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	ature of Examiner WAQTC #:		
Com	nments:		

WSDOT Errata to FOP for AASHTO T 176

Plastic Fines in Graded Aggregates and Soils by the Use of the Sand Equivalent Test

WAQTC FOP for AASHTO T 176 has been adopted by WSDOT with the following changes:

Sample Preparation

Replace step 7 with below:

7. WSDOT requires two samples.

Include step 8 below:

8. Dry the test sample in an oven in accordance with FOP for AASHTO T 255. The oven temperature shall not exceed 350°F (177°C). Cool to room temperature before testing. It is acceptable to place the test sample in a larger container to aid drying.

Procedure

- 6. After loosening the material from the bottom of the cylinder, shake the cylinder and contents by any one of the following methods:
 - c. Hand Method Method not recognized by WSDOT.
- 10. Clay and sand readings:

Replace step d with below:

d. If two Sand Equivalent (SE) samples are run on the same material and the second varies by more than \pm 8, based on the first cylinder result, additional tests shall be run.

FOP for AASHTO T 176

Plastic Fines in Graded Aggregates and Soils by the Use of the Sand Equivalent Test

Scope

This procedure covers the determination of plastic fines in accordance with AASHTO T 176-08. It serves as a rapid test to show the relative proportion of fine dust or clay-like materials in fine aggregates (FA) and soils.

Apparatus

See AASHTO T 176 for a detailed listing of sand equivalent apparatus. Note that the siphon tube and blow tube may be glass or stainless steel as well as copper.

- · Graduated plastic cylinder.
- Rubber stopper.
- Irrigator tube.
- Weighted foot assembly: Having a mass of 1000 ± 5g. There are two models of the
 weighted foot assembly. The older model has a guide cap that fits over the upper
 end of the graduated cylinder and centers the rod in the cylinder. It is read using a
 slot in the centering screws. The newer model has a sand-reading indicator 254 mm
 (10 in) above this point and is preferred for testing clay-like materials.
- · Bottle: clean, glass or plastic, of sufficient size to hold working solution
- Siphon assembly: The siphon assembly will be fitted to a 4 L (1 gal) bottle of
 working calcium chloride solution placed on a shelf 915 ± 25 mm (36 ± 1 in) above
 the work surface.
- Measuring can: With a capacity of 85 ± 5 mL (3 oz).
- Funnel: With a wide-mouth for transferring sample into the graduated cylinder.
- Quartering cloth: 600 mm (2 ft) square nonabsorbent cloth, such as plastic or oilcloth.
- Mechanical splitter: See the FOP for AASHTO R 76.
- Strike-off bar: A straightedge or spatula.
- Clock or watch reading in minutes and seconds.
- Manually-operated sand equivalent shaker: Capable of producing an oscillating motion at a rate of 100 complete cycles in 45 \pm 5 seconds, with a hand assisted half stroke length of 127 \pm 5 mm (5 \pm 0.2 in). It may be held stable by hand during the shaking operation. It is recommended that this shaker be fastened securely to a firm and level mount, by bolts or clamps, if a large number of determinations are to be made.
- Mechanical shaker: See AASHTO T 176 for equipment and procedure.
- Oven: Capable of maintaining a temperature of 110 ± 5°C (230 ± 9°F).
- Thermometer: Calibrated liquid-in-glass or electronic digital type designed for total immersion and accurate to 0.1°C (0.2°F).

Materials

- Stock calcium chloride solution: Obtain commercially prepared calcium chloride stock solution meeting AASHTO requirements.
- Working calcium chloride solution: Dilute one 3 oz. measuring can (85 ± 5 mL) of stock calcium chloride solution with 3.8 L (1 gal) distilled or demineralized water. Thoroughly mix the solution by filling the bottle with 2 L (½ gal) of water. Add the stock solution and agitate vigorously for 1 to 2 minutes. Add the remainder of the water, approximately 2 L (½ gal). Repeat the agitation process. The shelf life of the working solution is approximately 30 days. Label working solution with the date mixed. Discard working solutions more than 30 days old.

Note 1: The graduated cylinder filled to 4.4 in contains 88 mL and may be used to measure the stock solution.

Note 2: Tap water may be used if it is proven to be non-detrimental to the test and if it is allowed by the agency.

Control

The temperature of the working solution should be maintained at $22 \pm 3^{\circ}\text{C}$ ($72 \pm 5^{\circ}\text{F}$) during the performance of the test. If field conditions preclude the maintenance of the temperature range, reference samples should be submitted to the Central/Regional Laboratory, as required by the agency, where proper temperature control is possible. Samples that meet the minimum sand equivalent requirement at a working solution temperature outside of the temperature range need not be subject to reference testing.

Sample Preparation

- 1. Obtain the sample in accordance with the FOP for AASHTO T 2 and reduce in accordance with the FOP for AASHTO R 76.
- 2. Prepare sand equivalent test samples from the material passing the 4.75 mm (No. 4) sieve. If the material is in clods, break it up and re-screen it over a 4.75 mm (No. 4) sieve. All fines shall be cleaned from particles retained on the 4.75 mm (No. 4) sieve and included with the material passing that sieve.
- 3. Split or quarter 1000 to 1500 g of material from the portion passing the 4.75 mm (No. 4) sieve. Use extreme care to obtain a truly representative portion of the original sample.
 - **Note 3:** 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.
 - **Note 4:** All tests, including reference tests, will be performed utilizing Alternative Method No. 2 as described in AASHTO T 176, unless otherwise specified.
- 4. The sample must have the proper moisture content 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 content has been obtained.
 - **Note 5:** Clean sands having little 75 μ m (No. 200), such as sand for Portland Cement Concrete (PCC), may not form a cast.

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. When the moisture content is altered to provide the required cast, the altered sample should be placed in a pan, covered with a lid or with a damp cloth that does not touch the material, and allowed to stand for a minimum of 15 minutes. Samples that have been sieved without being air-dried and still retain enough natural moisture are exempted from this requirement.

If the material shows any free water, it is too wet to test and must be drained and air dried. Mix frequently to ensure uniformity. This drying process should continue until squeezing provides the required cast.

- 5. Place the sample on the quartering cloth and mix by alternately lifting each corner of the cloth and pulling it over the sample toward the diagonally opposite corner, being careful to keep the top of the cloth parallel to the bottom, thus 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.
- 6. Fill the measuring can by pushing it through the base of the pile while exerting pressure with the hand against the pile on the side opposite the measuring can. As the can is moved through 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 placing the maximum amount in the can. Strike off the can level full with the straightedge or spatula.
- 7. When required, repeat steps 5 and 6 to obtain additional samples.

Procedure

- 1. Start the siphon by forcing air into the top of the solution bottle through the tube while the pinch clamp is open. Siphon $101.6 \pm 2.5 \text{ mm}$ (4 $\pm 0.1 \text{ in}$) of working calcium chloride solution into the plastic cylinder.
- 2. Pour the prepared test sample from the measuring can into the plastic cylinder, using the funnel to avoid spilling.
- 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.
- 4. Allow the wetted sample to stand undisturbed for 10 ± 1 minutes.
- 5. At the end of the 10-minute period, stopper the cylinder and loosen the material from the bottom by simultaneously partially inverting and shaking the cylinder.
- 6. After loosening the material from the bottom of the cylinder, shake the cylinder and contents by any one of the following methods:
 - a. Mechanical Method Place the stoppered cylinder in the mechanical shaker, set the timer, and allow the machine to shake the cylinder and contents for 45 ± 1 seconds.

Caution: Agencies may require additional operator qualifications for the next two methods.

b. Manually-operated Shaker Method – Secure the stoppered cylinder in the three spring clamps on the carriage of the manually-operated sand equivalent shaker and set 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 strap.

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.

Proper shaking action is accomplished when the tip of the pointer 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 shaking for 100 strokes.

- c. Hand Method Hold the cylinder in a horizontal position 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 229 mm \pm 25 mm (9 \pm 1 in). 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.
- 7. Set the cylinder upright on the work table and remove the stopper.
- 8. 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. Work the irrigator tube to the bottom of the cylinder as quickly as possible, since it becomes more difficult to do this as the washing proceeds. This flushes the fine material into suspension above the coarser sand particles.

Continue to apply a stabbing and twisting action while flushing the fines upward until the cylinder is filled to the 381 mm (15 in) mark. Then raise the irrigator slowly without shutting off the flow so that the liquid level is maintained at about 381 mm (15 in) while the irrigator is being withdrawn. Regulate the flow just before the irrigator is entirely withdrawn and adjust the final level to 381 mm (15 in).

Note 6: Occasionally the holes in the tip of the irrigator tube may become clogged by a particle of sand. If the obstruction cannot be freed by any other method, use a pin or other sharp object to force it out, using extreme care not to enlarge the size of the opening. Also, keep the tip sharp as an aid to penetrating the sample.

- 9. Allow the cylinder and contents to stand undisturbed for 20 minutes ± 15 seconds. Start timing immediately after withdrawing the irrigator tube.
 - **Note 7:** Any vibration or movement of the cylinder during this time will interfere with the normal settling rate of the suspended clay and will cause an erroneous result.

10. Clay and sand readings:

- a. 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.
 - **Note 8:** If no clear line of demarcation has formed at the end of the 20-minute sedimentation period, allow the sample to stand undisturbed until a clay 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, rerun the test using three individual samples of the same material. Read and record the clay column height of the sample requiring the shortest sedimentation period only. Once a sedimentation time has been established, subsequent tests will be run using that time. The time will be recorded along with the test results on all reports.
- b. After the clay reading has been taken, place the weighted foot assembly over the cylinder and gently lower the assembly until it comes to rest on the sand. Do not allow the indicator to hit the mouth of the cylinder as the assembly is being lowered. Subtract 254 mm (10 in) from the level indicated by the extreme top edge of the indicator and record this value as the sand reading.
- c. If clay or sand readings fall between 2.5 mm (0.1 in) graduations, record the level of the higher graduation as the reading. For example, a clay reading that appears to be 7.95 would be recorded as 8.0; a sand reading that appears to be 3.22 would be recorded as 3.3.
- d. If two Sand Equivalent (SE) samples are run on the same material and the second varies by more than \pm 4, based on the first cylinder result, additional tests shall be run.
- e. If three or more Sand Equivalent (SE) samples are run on the same material, average the results. If an individual result varies by more than \pm 4, based on the average result, additional tests shall be run.

Calculations

Calculate the SE to the nearest 0.1 using the following formula:

$$SE = \frac{Sand\ Reading}{Clay\ Reading} \times 100$$

Example

Sand Reading = 3.3
Clay Reading = 8.0
$$SE = \frac{3.3}{8.0} \times 100 = 41.25 \ or \ 41.3 \qquad \text{Report } 42$$

Note 9: This example reflects the use of equipment made with English units. At this time, equipment made with metric units is not available.

Report the SE as the next higher whole number. In the example above, the 41.3 would be reported as 42. An SE of 41.0 would be reported as 41.

When averaging two or more samples, raise each calculated SE value to the next higher whole number (reported value) before averaging.

Example

calculated value 1 = 41.3

calculated value 2 = 42.8

These values are reported as 42 and 43, respectively.

Average the two reported values:

Average
$$SE = \frac{42 + 43}{2} = 42.5$$
 Report 43

If the average value is not a whole number, raise it to the next higher whole number.

Report

- · Results on forms approved by the agency
- Sample ID
- · Results to the whole number
- · Sedimentation time if over 20 minutes

Performance Exam Checklist

FOP for AASHTO T 176

Plastic Fines in Graded Aggregates and Soils by the Use of the Sand Equivalent Test

Part	icipant Name Exam Date		
Reco	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Proc	cedure Element	Trial 1	Trial 2
Sam	ple Preparation		
1.	Sample passed through 4.75 mm (No. 4) sieve?		
2.	Material in clods broken up and re-screened?		
3.	Split or quarter 1,000 to 1,500g of material passing the 4.75 mm (No. 4) sieve? Note: If necessary, the material may be dampened before splitting to avoid segregation or loss of fines.		
4.	No fines lost?		
5.	Working solution dated?		
6.	Temperature of working solution $22 \pm 3^{\circ}C$ ($72 \pm 5^{\circ}F$)?		
7.	Working calcium chloride solution 915 \pm 25 mm (36 \pm 1in) above the work surface?		
8.	$101.6 \pm 2.5 \text{ mm}$ (4 ± 0.1in) working calcium chloride solution siphoned into cylinder?		
9.	Material checked for moisture condition by tightly squeezing small portion in palm of hand and forming a cast?		
10.	Sample at proper water content?		
	a. If too dry (cast crumbles easily) water added, re-mixed, covered, and allowed to stand for at least 15 minutes?		
	b. If too wet (shows free water) sample drained, air dried and mixed frequently?		
11.	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?		
12.	Is material thoroughly mixed?		
13.	When material appears to be homogeneous, mixing finished with sample in a pile near center of cloth?		
14.	Fill the 85 mL (3 oz) tin by pushing through base of pile with other hand on opposite side of pile?		
15.	Material fills tin to overflowing?		
16.	Material compacted into tin with palm of hand?		
17.	Tin struck off level full with spatula or straightedge?		
18.	Prepared sample funneled into cylinder with no loss of fines?		

Pro	cedure Element	_ Trial 1	Trial 2
19.	Bottom of cylinder tapped sharply on heel of hand several times to release air bubbles?		
20.	Wetted sample allowed to stand undisturbed for 10 min. ± 1 min?		
21.	Cylinder stoppered and material loosened from bottom by shaking?		
22.	Stoppered cylinder placed properly in mechanical shaker and cylinder shaken 45 ± 1 seconds?		
23.	Following shaking, cylinder set vertical on work surface and stopper removed?		
24.	Irrigator tube inserted in cylinder and material rinsed from cylinder walls as irrigator is lowered?		
25.	Irrigator tube forced through material to bottom of cylinder by gentle stabbing and twisting action?		
26.	Stabbing and twisting motion applied until cylinder filled to 381 mm (15 in) mark?		
27.	Liquid raised and maintained at 381 mm (15 in) mark while irrigator is being withdrawn?		
28.	Liquid at the 381 mm (15 in) mark?		
29.	Contents let stand 20 minutes ± 15 seconds?		
30.	Timing started immediately after withdrawal of irrigator?		
31.	No vibration or disturbance of the sample?		
32.	Readings taken at 20 minutes or up to 30 minutes, when a definite line appears?		
33.	Clay level correctly read, rounded, and recorded?		
34.	Weighted foot assembly lowered into cylinder without hitting mouth of cylinder?		
35.	Sand level correctly read, rounded, and recorded?		
36.	Calculations performed correctly?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
Com	ments:		

MOISTURE-DENSITY RELATIONS OF SOILS:

FOP for AASHTO T 99
USING A 2.5 kg (5.5 lb) RAMMER AND A 305 mm (12 in) DROP

FOP for AASHTO T 180
USING A 4.54 kg (10 lb) RAMMER AND A 457 mm (18 in) DROP

Scope

This procedure covers the determination of the moisture-density relations of soils and soil-aggregate mixtures in accordance with two similar test methods:

- AASHTO T 99-17: Methods A, B, C, and D
- AASHTO T 180-17: Methods A, B, C, and D

This test method applies to soil mixtures having 40 percent or less retained on the 4.75 mm (No. 4) sieve for methods A or B, or, 30 percent or less retained on the 19 mm (¾ in) with methods C or D. The retained material is defined as oversize (coarse) material. If no minimum percentage is specified, 5 percent will be used. Samples that contain oversize (coarse) material that meet percent retained criteria should be corrected by using Annex A, Correction of Maximum Dry Density and Optimum Moisture for Oversized Particles. Samples of soil or soil-aggregate mixture are prepared at several moisture contents and compacted into molds of specified size, using manual or mechanical rammers that deliver a specified quantity of compactive energy. The moist masses of the compacted samples are multiplied by the appropriate factor to determine wet density values. Moisture contents of the compacted samples are determined and used to obtain the dry density values of the same samples. Maximum dry density and optimum moisture content for the soil or soil-aggregate mixture is determined by plotting the relationship between dry density and moisture content.

Apparatus

- Mold Cylindrical mold made of metal with the dimensions shown in Table 1 or Table 2. If permitted by the agency, the mold may be of the "split" type, consisting of two half-round sections, which can be securely locked in place to form a cylinder. Determine the mold volume according to *Annex B*, *Standardization of the Mold*.
- Mold assembly Mold, base plate, and a detachable collar.
- Rammer –Manually or mechanically-operated rammers as detailed in Table 1 or Table 2. A manually-operated rammer shall be equipped with a guide sleeve to control the path and height of drop. The guide sleeve shall have at least four vent holes no smaller than 9.5 mm (¾ in) in diameter, spaced approximately 90 degrees apart and approximately 19 mm (¾ in) from each end. A mechanically-operated rammer will uniformly distribute blows over the sample and will be calibrated with several soil types, and be adjusted, if necessary, to give the same moisture-density results as with the manually operated rammer. For additional information concerning calibration, see the FOP for AASHTO T 99 and T 180.
- Sample extruder A jack, lever frame, or other device for extruding compacted specimens from the mold quickly and with little disturbance.

• Balance(s) or scale(s) of the capacity and sensitivity required for the procedure used by the agency.

A balance or scale with a capacity of 11.5 kg (25 lb) and a sensitivity of 1 g for obtaining the sample, meeting the requirements of AASHTO M 231, Class G 5.

A balance or scale with a capacity of 2 kg and a sensitivity of 0.1 g is used for moisture content determinations done under both procedures, meeting the requirements of AASHTO M 231, Class G 2.

- Drying apparatus A thermostatically controlled drying oven, capable of maintaining a temperature of 110 ± 5°C (230 ± 9°F) for drying moisture content samples in accordance with the FOP for AASHTO T 255/T 265.
- Straightedge A steel straightedge at least 250 mm (10 in) long, with one beveled edge and at least one surface plane within 0.1 percent of its length, used for final trimming.
- Sieve(s) 4.75 mm (No. 4) and/or 19.0 mm (¾ in), meeting the requirements of FOP for AASHTO T 27/T 11.
- Mixing tools Miscellaneous tools such as a mixing pan, spoon, trowel, spatula, etc., or a suitable mechanical device, for mixing the sample with water.
- Containers with close-fitting lids to prevent gain or loss of moisture in the sample.

Table 1 Comparison of Apparatus, Sample, and Procedure – Metric

	T 99	T 180	
Mold Volume, m ³	Methods A, C: 0.000943 ± 0.000014	Methods A, C: 0.000943 ± 0.000014	
	Methods B, D: 0.002124 ± 0.00025	Methods B, D: 0.002124 ± 0.000025	
Mold Diameter, mm	Methods A, C: 101.60 ± 0.40	Methods A, C: 101.60 ± 0.4	
	Methods B, D: 152.40 ± 0.70	Methods B, D: 152.40 ± 0.70	
Mold Height, mm	116.40 ± 0.50	116.40 ± 0.50	
Detachable Collar Height, mm	50.80 ± 0.64	50.80 ± 0.64	
Rammer Diameter, mm	50.80 ± 0.25	50.80 ± 0.25	
Rammer Mass, kg	2.495 ± 0.009	4.536 ± 0.009	
Rammer Drop, mm	305	457	
Layers	3	5	
Blows per Layer	Methods A, C: 25	Methods A, C: 25	
	Methods B, D: 56	Methods B, D: 56	
Material Size, mm	Methods A, B: 4.75 minus	Methods A, B: 4.75 minus	
	Methods C, D: 19.0 minus	Methods C, D: 19.0 minus	
Test Sample Size, kg	Method A: 3 Method B: 7		
	Method C: 5 (1) Method D: 11(1)		
Energy, kN-m/m ³	592	2,693	

⁽¹⁾ This may not be a large enough sample depending on your nominal maximum size for moisture content samples.

	T 99 T 180					
Mold Volume, ft ³	Methods A, C: 0.0333 ± 0.0005	Methods A, C: 0.0333 ± 0.0005				
	Methods B, D: 0.07500 ± 0.0009	Methods B, D: 0.07500 ± 0.0009				
Mold Diameter, in	Methods A, C: 4.000 ± 0.016	Methods A, C: 4.000 ± 0.016				
	Methods B, D: 6.000 ± 0.026	Methods B, D: 6.000 ± 0.026				
Mold Height, in	4.584 ± 0.018	4.584 ± 0.018				
Detachable Collar Height, in	2.000 ± 0.025	2.000 ± 0.025				
Rammer Diameter, in	2.000 ± 0.025	2.000 ± 0.025				
Rammer Mass, Ib	5.5 ± 0.02	10 ± 0.02				
Rammer Drop, in	12	18				
Layers	3	5				
Blows per Layer	Methods A, C: 25	Methods A, C: 25				
	Methods B, D: 56	Methods B, D: 56				
Material Size, in	Methods A, B: No. 4 minus	Methods A, B: No.4 minus				
	Methods C, D: ¾ minus	Methods C, D: ¾ minus				
Test Sample Size, lb	Method A: 7 Method B: 16					
	Method C: 12 ₍₁₎ Method D: 25 ₍₁₎					
Energy, lb-ft/ft ³	12,375	56,250				

Table 2 Comparison of Apparatus, Sample, and Procedure – English

Sample

If the sample is damp, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus maintained at a temperature not exceeding 60°C (140°F). Thoroughly break up aggregations in a manner that avoids reducing the natural size of individual particles.

Obtain a representative test sample of the mass required by the agency by passing the material through the sieve required by the agency. See Table 1 or Table 2 for test sample mass and material size requirements.

In instances where the material is prone to degradation, i.e., granular material, a compaction sample with differing moisture contents should be prepared for each point.

Note 1: Both T 99 and T 180 have four methods (A, B, C, D) that require different masses and employ different sieves.

Note 2: If the sample is plastic (clay types), it should stand for a minimum of 12 hours after the addition of water to allow the moisture to be absorbed. In this case, several samples at different moisture contents should be prepared, put in sealed containers and tested the next day.

⁽¹⁾ This may not be a large enough sample depending on your nominal maximum size for moisture content samples.

Procedure

During compaction, rest the mold firmly on a dense, uniform, rigid, and stable foundation or base. This base shall remain stationary during the compaction process.

- 1. Determine the mass of the clean, dry mold. Include the base plate, but exclude the extension collar. Record the mass to the nearest 1 g (0.005 lb).
- 2. Thoroughly mix the selected representative sample with sufficient water to dampen it to approximately 4 to 8 percentage points below optimum moisture content. See Note 2. For many materials, this condition can be identified by forming a cast by hand.
- 3. Form a specimen by compacting the prepared soil in the mold assembly in approximately equal layers. For each layer:
 - a. Spread the loose material uniformly in the mold.
 - **Note 3:** It is recommended to cover the remaining material with a non-absorbent sheet or damp cloth to minimize loss of moisture.
 - b. Lightly tamp the loose material with the manual rammer or other similar device, this establishes a firm surface.
 - c. Compact each layer with uniformly distributed blows from the rammer. See Table 1 for mold size, number of layers, number of blows, and rammer specification for the various test methods. Use the method specified by the agency.
 - d. Trim down material that has not been compacted and remains adjacent to the walls of the mold and extends above the compacted surface.
- 4. Remove the extension collar. Avoid shearing off the sample below the top of the mold. The material compacted in the mold should not be over 6 mm (¼ in) above the top of the mold once the collar has been removed.
- 5. Trim the compacted soil even with the top of the mold with the beveled side of the straightedge.
- 6. Determine and record the mass of the mold, base plate, and wet soil to the nearest 1 g (0.005 lb) or better.
- 7. Determine and record the wet mass (M_w) of the sample by subtracting the mass in Step 1 from the mass in Step 6.
- 8. Calculate the wet density, in kg/m³ (lb/ft³), by dividing the wet mass by the measured volume (V_m).
- 9. Extrude the material from the mold. For soils and soil-aggregate mixtures, slice vertically through the center and take a representative moisture content sample from one of the cut faces, ensuring that all layers are represented. For granular materials, a vertical face will not exist. Take a representative sample. This sample must meet the sample size requirements of the test method used to determine moisture content.

Note 4: When developing a 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 from the mixing bowl may be preferred in order to determine the amount of moisture available for compaction.

- 10. Determine and record the moisture content of the sample in accordance with the FOP for AASHTO T 255/T 265.
- 11. Thoroughly break up the remaining portion of the molded specimen until it will again pass through the sieve, as judged by eye, and add to the remaining portion of the sample being tested. See Note 2.
- 12. Add sufficient water to increase the moisture content of the remaining soil by 1 to 2 percentage points and repeat steps 3 through 11.
- 13. Continue determinations until there is either a decrease or no change in the wet mass. There will be a minimum of three points on the dry side of the curve and two points on the wet side. For non-cohesive, drainable soils, one point on the wet side is sufficient.

Calculations

Wet Density

$$D_w = \frac{M_w}{V_m}$$

Where:

 $D_w = \text{wet density, kg/m}^3 \text{ (lb/ft}^3)$

 $M_{\rm w}$ = wet mass

 $V_{\rm m}$ = volume of the mold, Annex B

Dry Density

$$D_d = \left(\frac{D_w}{w + 100}\right) \times 100 \quad or \quad D_d = \frac{D_w}{\left(\frac{W}{100}\right) + 1}$$

Where:

D_d = dry density, kg/m³ (lb/ft³) w = moisture content, as a percentage

Example for 4-inch mold, Methods A or C

Wet mass, $M_{w} = 1.944 \text{ kg} (4.25 \text{ lb})$

Moisture content, w = 11.3 percent

Measured volume of the mold, $V_m = 0.000946 \text{ m}^3 (0.0334 \text{ ft}^3)$

Wet Density

$$D_w = \frac{1.944 \, kg}{0.000946 \, m^3} = 2055 \, kg/m^3 \quad D_w = \frac{4.25 \, lb}{0.0334 \, ft^3} = 127.2 \, lb/ft^3$$

Dry Density

$$D_d = \left(\frac{2055 \, kg/m^3}{11.3 + 100}\right) \times 100 = 1846 \, kg/m^3 \quad D_d = \left(\frac{127.2 \, lb/ft^3}{11.3 + 100}\right) \times 100 = 114.3 \, lb/ft^3$$

Or

$$D_d = \left(\frac{2055 \, kg/m^3}{\frac{11.3}{100} + 1}\right) = 1846 \, kg/m^3 \quad D_d = \left(\frac{127.2 \, lb/ft^3}{\frac{11.3}{100} + 1}\right) = 114.3 \, lb/ft^3$$

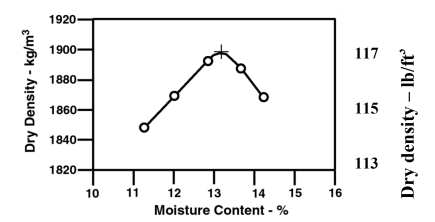
Moisture-Density Curve Development

When dry density is plotted on the vertical axis versus moisture content on the horizontal axis and the points are connected with a smooth line, a moisture-density curve is developed. The coordinates of the peak of the curve are the maximum dry density, or just "maximum density," and the "optimum moisture content" of the soil.

Example

Given the following dry density and corresponding moisture content values develop a moisture-density relations curve and determine maximum dry density and optimum moisture content.

Dry D	Moisture	
kg/m³	lb/ft ³	Content, %
1846	114.3	11.3
1868	115.7	12.1
1887	116.9	12.8
1884	116.7	13.6
1871	115.9	14.2



In this case, the curve has its peak at:

Maximum dry density = $1890 \text{ kg/m}^3 (117.0 \text{ lb/ft}^3)$

Optimum moisture content = 13.2 percent

Note that both values are approximate, since they are based on sketching the curve to fit the points.

Report

- Results on forms approved by the agency
- Sample ID
- Maximum dry density to the closest 1 kg/m³ (0.1 lb/ft³)
- Optimum moisture content to the closest 0.1 percent

ANNEX A

Correction of Maximum DRY Density and Optimum Moisture for Oversized Particles

This section corrects the maximum dry density and moisture content of the material retained on the 4.75 mm (No. 4) sieve, Methods A and B; or the material retained on the 19 mm (¾ in) sieve, Methods C and D. The maximum dry density, corrected for oversized particles and total moisture content, are compared with the field-dry density and field moisture content.

This correction can be applied to the sample on which the maximum dry density is performed. A correction may not be practical for soils with only a small percentage of oversize material. The agency shall specify a minimum percentage below which the method is not needed. If not specified, this method applies when more than 5 percent by weight of oversize particles is present.

Bulk specific gravity (G_{sb}) of the oversized particles is required to determine the corrected maximum dry density. Use the bulk specific gravity as determined using the FOP for AASHTO T 85 in the calculations. For construction activities, an agency established value or specific gravity of 2.600 may be used.

This correction can also be applied to the sample obtained from the field while performing in-place density.

- Use the sample from this procedure or a sample obtained according to the FOP for AASHTO T 310.
- 2. Sieve the sample on the 4.75 mm (No. 4) sieve for Methods A and B or the 19 mm (¾ in) sieve, Methods C and D.
- 3. Determine the dry mass of the oversized and fine fractions (M_{DC} and M_{DF}) by one of the following:
 - a. Dry the fractions, fine and oversized, in air or by use of a drying apparatus that is maintained at a temperature not exceeding 60°C (140°F).
 - b. Calculate the dry masses using the moisture samples.

To determine the dry mass of the fractions using moisture samples.

- 1. Determine the moist mass of both fractions, fine (M_{Mf}) and oversized (M_{Mc}) :
- 2. Obtain moisture samples from the fine and oversized material.
- 3. Determine the moisture content of the fine particles (MC_f) and oversized particles (MC_C) of the material by FOP for AASHTO T 255/T 265 or agency approved method.
- 4. Calculate the dry mass of the oversize and fine particles.

$$M_D = \frac{M_m}{1 + MC}$$

Where:

 M_D = mass of dry material (fine or oversize particles)

 M_m = mass of moist material (fine or oversize particles)

MC = moisture content of respective fine or oversized, expressed as a decimal

5. Calculate the percentage of the fine (P_f) and oversized (P_c) particles by dry weight of the total sample as follows: See Note 2.

$$P_f = \frac{100 \times M_{DF}}{M_{DF} + M_{DC}} \qquad \frac{100 \times 15.4 \ lb}{15.4 \ lbs + 5.7 \ lb} = 73\% \qquad \frac{100 \times 6.985 \ kg}{6.985 \ kg + 2.602 \ kg} = 73\%$$

And

$$P_c = \frac{100 \times M_{DC}}{M_{DF} + M_{DC}} \qquad \frac{100 \times 5.7 \ lb}{15.4 \ lbs + 5.7 \ lb} = 27\% \qquad \frac{100 \times 2.585 kg}{6.985 \ kg + 2.585 \ kg} = 27\%$$

Or for P_c :

$$P_c = 100 - P_f$$

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 fine particles

 M_{DC} = mass of oversize particles

Optimum Moisture Correction Equation

1. Calculate the corrected moisture content as follows:

$$MC_T = \frac{\left(MC_F \times P_f\right) + \left(MC_c \times P_c\right)}{100} \qquad \frac{\left(13.2\% \times 73.0\%\right) + \left(2.1\% \times 27.0\%\right)}{100} = 10.2\%$$

Where:

MC_T = corrected moisture content of combined fines and oversized particles, expressed as a percent moisture

 MC_f = moisture content of fine particles, as a percent moisture MC_C = moisture content of oversized particles, as a percent moisture

Note 1: Moisture content of oversize material can be assumed to be two (2) percent for most construction applications.

Note 2: In some field applications agencies will allow the percentages of oversize and fine materials to be determined with the materials in the wet state.

Density Correction Equation

1. Calculate the corrected dry density of the total sample (combined fine and oversized particles) as follows:

$$D_d = \frac{100\%}{\left[\left(\frac{P_f}{D_f} \right) + \left(\frac{P_c}{k} \right) \right]}$$

Where:

 D_d = corrected total dry density (combined fine and oversized particles) kg/m³ (lb/ ft^3)

 D_f = dry density of the fine particles kg/m³ (lb/ft³), determined in the lab

P_c = percent of oversize particles, of sieve used, by weight.

 P_f = percent of fine particles, of sieve used, by weight.

 $k = Metric: 1,000 * Bulk Specific Gravity (<math>G_{sb}$) (oven dry basis) of coarse particles (kg/m^3).

k = English: 62.4 * Bulk Specific Gravity (G_{sb}) (oven dry basis) of coarse particles (Ib/ft^3)

Note 3: If the specific gravity is known, then this value will be used in the calculation. For most construction activities the specific gravity for aggregate may be assumed to be 2.600.

Calculation

Example

Metric:

Maximum laboratory dry density (D_f): 1890 kg/m³ Percent coarse particles (P_c): 27 percent

Percent fine particles (P_f): 73 percent

Mass per volume coarse particles (k): $(2.697) (1000) = 2697 \text{ kg/m}^3$

$$D_d = \frac{100\%}{\left[\left(\frac{P_f}{D_f} \right) + \left(\frac{P_c}{k} \right) \right]}$$

$$D_d = \frac{100\%}{\left[\left(\frac{73\%}{1890 \, kg/m^3} \right) + \left(\frac{27\%}{2697 \, kg/m^3} \right) \right]}$$

$$D_d = \frac{100\%}{[0.03862 \, kg/m^3 + 0.01001 \, kg/m^3]}$$

$$D_d = 2056.3 \, kg/m^3 \, report \, 2056 \, kg/m^3$$

English:

Maximum laboratory dry density (D_f): 117.0 lb/ft³

Percent coarse particles (P_c): 27 percent

Percent fine particles (P_f) : 73 percent

Mass per volume of coarse particles (k): $(2.697) (62.4) = 168.3 \text{ lb/ft}^3$

$$D_{d} = \frac{100\%}{\left[\left(\frac{P_{f}}{D_{f}}\right) + \left(\frac{P_{c}}{k}\right)\right]}$$

$$D_{d} = \frac{100\%}{\left[\left(\frac{73\%}{117.0 \, lb/ft^{3}}\right) + \left(\frac{27\%}{168.3 \, lb/ft^{3}}\right)\right]}$$

$$D_{d} = \frac{100\%}{\left[0.6239 \, lb/ft^{3} + 0.1604 \, lb/ft^{3}\right]}$$

$$D_{d} = \frac{100\%}{0.7843 \, lb/ft^{3}}$$

 $D_d = 127.50 \ lb/ft^3 \ Report 127.5 \ lb/ft^3$

Report

- · Results on forms approved by the agency
- Sample ID
- Corrected maximum dry density to the closest 1 kg/m³ (0.1 lb/ft³)
- · Corrected optimum moisture to the 0.1 percent

ANNEX B

Standardization of The Mold

Standardization is a critical step to ensure accurate test results when using this apparatus. Failure to perform the standardization procedure as described herein will produce inaccurate or unreliable test results.

Apparatus

- Mold and base plate
- Balance or scale Accurate to within 45 g (0.1 lb) or 0.3 percent of the test load, whichever is greater, at any point within the range of use.
- Cover plate A piece of plate glass, at least 6 mm (1/4 in) thick and at least 25 mm (1 in) larger than the diameter of the mold.
- Thermometers Standardized liquid-in-glass, or electronic digital total immersion type, accurate to 0.5°C (1°F)

Procedure

- 1. Create a watertight seal between the mold and base plate.
- 2. Determine and record the mass of the dry sealed mold, base plate, and cover plate.
- 3. Fill the mold with water at a temperature between 16°C and 29°C (60°F and 85°F) and cover with the cover plate in such a way as to eliminate bubbles and excess water.
- 4. Wipe the outside of the mold, base plate, and cover plate dry, being careful not to lose any water from the mold.
- 5. Determine and record the mass of the filled mold, base plate, cover plate, and water.
- 6. Determine and record the mass of the water in the mold by subtracting the mass in Step 2 from the mass in Step 5.
- 7. Measure the temperature of the water and determine its density from Table B1, interpolating as necessary.
- 8. Calculate the volume of the mold, V_m , by dividing the mass of the water in the mold by the density of the water at the measured temperature.

Calculations

$$V_m = \frac{M}{D}$$

Where:

V_m = volume of the mold M = mass of water in the mold

D = density of water at the measured temperature

Example

Mass of water in mold = 0.94061 kg (2.0737 lb)

Density of water at 23°C (73.4°F) = $997.54 \text{ kg/m}^3 (62.274 \text{ lb/ft}^3)$

$$V_m = \frac{0.94061 \, kg}{997.54 \, kg/m^3} = 0.000943 \, m^3$$
 $V_m = \frac{2.0737 \, lb}{62.274 \, lb/ft^3} = 0.0333 \, ft^3$

Table B1 Unit Mass of Water 15°C to 30°C

°C	(°F)	kg/m ³	(lb/ft ³)	°C	(°F)	kg/m ³	(lb/ft ³)
15	(59.0)	999.10	(62.372)	23	(73.4)	997.54	(62.274)
15.6	(60.0)	999.01	(62.366)	23.9	(75.0)	997.32	(62.261)
16	(60.8)	998.94	(62.361)	24	(75.2)	997.29	(62.259)
17	(62.6)	998.77	(62.350)	25	(77.0)	997.03	(62.243)
18	(64.4)	998.60	(62.340)	26	(78.8)	996.77	(62.227)
18.3	(65.0)	998.54	(62.336)	26.7	(80.0)	996.59	(62.216)
19	(66.2)	998.40	(62.328)	27	(80.6)	996.50	(62.209)
20	(68.0)	998.20	(62.315)	28	(82.4)	996.23	(62.192)
21	(69.8)	997.99	(62.302)	29	(84.2)	995.95	(62.175)
21.1	(70.0)	997.97	(62.301)	29.4	(85.0)	995.83	(62.166)
22	(71.6)	997.77	(62.288)	30	(86.0)	995.65	(62.156)

Report

- Mold ID
- · Date Standardized
- Temperature of the water
- Volume, V_m, of the mold

Performance Exam Checklist FOP for AASHTO T 180 Moisture-Density Relation of Soils

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	If damp, sample dried in air or drying apparatus, not exceeding 60°C (140°F)?		
2.	2. Sample broken up and an adequate amount sieved over the appropriate sieve (4.75 mm/No. 4 or 19.0 mm/3/4 in) to determine oversize (coarse particle) percentage?		
3.	Sample passing the sieve has appropriate mass?		
4.	If soil is plastic (clay types):		
	a. Multiple samples mixed with water varying moisture content by 1 to 2 percent, bracketing the optimum moisture content?		
	b. Samples placed in covered containers and allowed to stand for at least 12 hours?		
5.	Sample determined to be 4 to 8 percent below expected optimum moisture content?		
6.	Mold placed on rigid and stable foundation?		
7.	Layer of soil (approximately one fifth compacted depth) placed in mold with collar attached, loose material lightly tamped?		
8.	Soil compacted with appropriate number of blows (25 or 56)?		
9.	Material adhering to the inside of the mold trimmed?		
10.	Layer of soil (approximately two fifths compacted depth) placed in mold with collar attached, loose material lightly tamped?		
11.	Soil compacted with appropriate number of blows (25 or 56)?		
12.	Material adhering to the inside of the mold trimmed?		
13.	Layer of soil (approximately three fifths compacted depth) placed in mold with collar attached, loose material lightly tamped?		
14.	Soil compacted with appropriate number of blows (25 or 56)?		
15.	Material adhering to the inside of the mold trimmed?		
16.	Layer of soil (approximately four fifths compacted depth) placed in mold with collar attached, loose material lightly tamped?		
17.	Soil compacted with appropriate number of blows (25 or 56)?		
18.	Material adhering to the inside of the mold trimmed?		
19.	Mold filled with soil such that compacted soil will be above the mold, loose material lightly tamped?		
20.	Soil compacted with appropriate number of blows (25 or 56)?		

Proc	edure Eler	ment					Trial 1	Trial 2
21.	Collar rem	noved wit	hout shearing	off sample?				
22.	Approxim (without t	-		mpacted material above	the top (of the mold		
23.	Soil trimm	ned to top	of mold with	the beveled side of the s	straighte	dge?		
24.	Mass of m	nold and o	contents deter	mined to appropriate pre	ecision?			
25.	Wet dens	ity calcula	ated from the	wet mass?				
26.	Soil remov	ved from	mold using a s	ample extruder if neede	d?			
27.	Soil sliced	vertically	/ through cent	er (non-granular materia	l)?			
28.	Moisture	sample re	emoved ensuri	ng all layers are represen	ited?			
29.	Moist ma	ss determ	ined immediat	tely to 0.1 g?				
30.	Moisture	sample m	ass of correct	size?				
31.	Sample dr T 255/T 2		vater content o	determined according to	the FOF	P for		
32.				broken up until it will pa emainder of original test		-		
33.	33. Water added to increase moisture content of the remaining sample in 1 to 2 percent increments?							
34.	Steps 2 th	rough 20	repeated for	each increment of water	added?			
35.	If soil is pl	lastic (clay	/ types):					
	-		with water val optimum mois	rying moisture content b ture content?	y 1 to 2	percent,		
	b. Sampl	-	l in covered co	ntainers and allowed to	stand fo	r at least		
36.	If materia	l is degrad	dable:					
			nixed with wat mum moisture	er varying moisture cont e content?	ent by 1	to 2 percent,		
37.	Process co	ontinued	until wet dens	ity either decreases or st	tabilizes	?		
38.	Moisture	content a	nd dry density	calculated for each sam	ple?			
39.	-		l on vertical ax ed with a smo	is, moisture content plot oth curve?	tted on h	norizontal axis,		
40.			t peak of curv t 0.1 percent?	e recorded as optimum v	vater co	ntent and		
41.	Dry densi nearest 1	•		content reported as ma	ximum d	lensity to		
42.	Corrected	l for coars	se particles if a	applicable?				
First	Attempt:	Pass	Fail	Second Attempt:	Pass	Fail		
Signa	ture of Ex	aminer _			_ W	'AQTC #:		

Comments:

Performance Exam Checklist

Air Content of Concrete (Volumetric Method) for AASHTO T 196

Participant Name

Exam Date

Yes No

1. Bowl filled in two equal layers?

2. Each layer rodded 25 times?

Procedure Element

- 3. Bowl tapped (sharply) 10 to 15 times after rodding each layer?
- 4. Excess concrete removed with strike-off bar or plate?
- 5. Flange of bowl wiped clean?
- 6. Using funnel, water added, then alcohol added, then final water added until liquid level appears in neck?
- 7. Funnel removed & water adjusted to zero mark using rubber syringe?
- 8. Screw cap is attached and tightened?

Initial Reading

- 9. Unit inverted and agitated at 5 second intervals for a minimum of 45 seconds and until concrete is free from base?
- 10. Unit vigorously rolled ½ to ½ turn forward and back several times with base at a 45° angle. Then turn base about ½ turn and rolling process resumed.
- 11. Was meter checked for leaking?
 - a. If leak was found, was test started over with new sample?
- 12. Apparatus placed upright, top loosened and allowed to stand until air rises to the top?
 - a. < 0.25 percent change in 2 minutes (without excessive foam), initial reading recorded to the nearest 0.25%?
 - b. More than 6 minutes to stabilize or excessive foam, was test discarded and new test run?

Confirmation of Initial Meter Reading

- 13. 1 minute rolling repeated and liquid level checked?
- 14. Confirmation reading > 0.25 percent of initial, new reading recorded as new initial reading, repeat 1 minute rolling
- 15. Level of liquid read < 0.25 percent change, final meter reading recorded to nearest 0.25%?
- 16. Apparatus disassembled and checked for undisturbed concrete

Calculations

- 17. Correction factor from Table 1 subtracted for use of 2.5 pts or more of alcohol?
- 18. If required, number of calibration cups of water added to air content?
- 19. Air content reported to the nearest 0.25 percent air?

First Attempt: Pas	ss Fail	Second Attempt:	Pass	Fail
Signature of Exami	ner			
Comments:				

FOP for AASHTO T 209

Theoretical Maximum Specific Gravity (G_{mm}) and Density of Hot Mix Asphalt (Hma) Paving Mixtures

Scope

This procedure covers the determination of the maximum specific gravity (G_{mm}) of uncompacted hot mix asphalt (HMA) paving mixtures in accordance with AASHTO T 209-12. Two methods using different containers – bowl and pycnometer/volumetric flask– are covered.

Specimens prepared in the laboratory shall be cured according to agency standards.

Apparatus

- Balance or scale: 10,000 g capacity, readable to 0.1 g
- Container: A glass, metal, or plastic bowl, pycnometer or volumetric flask between 2000 and 10,000 mL as required by the minimum sample size requirements in Table 1 sample and capable of withstanding a partial vacuum
- Pycnometer/volumetric flask cover: A glass plate or a metal or plastic cover with a vented opening
- Vacuum lid: A transparent lid with a suitable vacuum connection, with a vacuum opening to be covered with a fine wire mesh
- Vacuum pump or water aspirator: Capable of evacuating air from the container to a residual pressure of 4.0 kPa (30 mm Hg)
- Residual pressure manometer or vacuum gauge: Traceable to NIST and capable of measuring residual pressure down to 4.0 kPa (30 mm Hg) or less
- Manometer or vacuum gauge: Capable of measuring the vacuum being applied at the source of the vacuum
- Water bath: A constant-temperature water bath (optional)
- Thermometers: Standardized liquid-in-glass, or electronic digital total immersion type, accurate to 0.5°C (1°F)
- Bleeder valve to adjust vacuum
- Timer

Standardization of Pycnometer or Volumetric Flask

Use a pycnometer/volumetric flask that is standardized to accurately determine the mass of water, at 25 ± 0.5 °C (77 \pm 1°F), in the pycnometer/volumetric flask. The pycnometer/volumetric flask shall be standardized periodically in conformance with procedures established by the agency.

Test Sample Preparation

Obtain samples in accordance with the FOP for AASHTO T 168 and reduce according to the FOP for AASHTO R 47.

Test sample size shall conform to the requirements of Table 1. Samples larger than the capacity of the container may be tested in two or more increments. Results will be combined and averaged. If the increments have a specific gravity difference greater than 0.014 the test must be re-run.

Table 1Test Sample Size for G_{mm}

Nominal Maximum* Aggregate Size mm (in)		Minimum Mass g
37.5 or greater	(1½)	4000
19 to 25	(¾ to 1)	2500
12.5 or smaller	(½)	1500

^{*}Nominal maximum size: One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained.

Procedure - General

Two procedures – bowl and pycnometer/volumetric flask – are covered. The first 11 steps are the same for both.

- 1. Separate the particles of the sample, taking care not to fracture the mineral particles, so that the particles of the fine aggregate portion are not larger than 6.3 mm (¼ in). If the mixture is not sufficiently soft to be separated manually, place it in a large flat pan and warm in an oven only until it is pliable enough for separation.
- 2. Cool the sample to room temperature.
- 3. Determine and record the mass of the dry container to the nearest 0.1 g.
- 4. Place the sample in the container.
- 5. Determine and record the mass of the dry container and sample to the nearest 0.1 g.
- 6. Determine and record the mass of the sample by subtracting the mass determined in Step 3 from the mass determined in Step 5. Designate this mass as "A".
- 7. Add sufficient water at approximately 25° C (77° F) to cover the sample by about 25 mm (1 in).
 - **Note 1:** The release of entrapped air may be facilitated by the addition of a wetting agent. Check with the agency to see if this is permitted and, if it is, for a recommended agent.
- 8. Place the lid on the container and attach the vacuum line. To ensure a proper seal between the container and the lid, wet the O-ring or use a petroleum gel.
- 9. Remove entrapped air by subjecting the contents to a partial vacuum of 3.7 ± 0.3 kPa (27.5 \pm 2.5 mm Hg) residual pressure for 15 \pm 2 minutes.

- 10. Agitate the container and contents, either continuously by mechanical device or manually by vigorous shaking, at 2 minute intervals. This agitation facilitates the removal of air.
- 11. Release the vacuum, increasing the pressure to atmospheric pressure in 10 to 15 seconds, turn off the vacuum pump, and remove the lid. When performing the pycnometer/volumetric flask method, complete steps 12B through 16B within 10 ± 1 minutes.

Procedure - Bowl

- 12A. Fill the water bath to overflow level with water at $25 \pm 1^{\circ}$ C (77 $\pm 2^{\circ}$ F) and allow the water to stabilize.
- 13A. Zero or tare the balance with the immersion apparatus attached, ensuring that the device is not touching the sides or the bottom of the water bath.
- 14A. Suspend and immerse the bowl and contents in water at $25 \pm 1^{\circ}$ C (77 $\pm 2^{\circ}$ F) for 10 ± 1 minutes. The holder shall be immersed sufficiently to cover both it and the bowl.
- 15A. Determine and record the submerged weight of the bowl and contents to the nearest 0.1 g.
- 16A. Refill the water bath to overflow level.
- 17A. Empty and re-submerge the bowl following Step 12A to determine the submerged weight of the bowl to the nearest 0.1 g.
- 18A. Determine and record the submerged weight of the sample to the nearest 0.1 g by subtracting the submerged weight of the bowl from the submerged weight determined in Step 15A. Designate this submerged weight as "C".

Procedure – Pycnometer or Volumetric Flask

- 12B. Immediately fill the pycnometer/volumetric flask with water without reintroducing air.
- 13B. Stabilize the temperature of the pycnometer/volumetric flask and contents so that the final temperature is within $25 \pm 1^{\circ}\text{C}$ (77 ± 2°F).
- 14B. Finish filling the pycnometer/volumetric flask with water that is $25 \pm 1^{\circ}$ C (77 $\pm 2^{\circ}$ F), place the cover or a glass plate on the pycnometer/volumetric flask, and eliminate all air.
 - **Note 2:** When using a metal pycnometer and cover, place the cover on the pycnometer and push down slowly, forcing excess water out of the hole in the center of the cover. Use care when filling the pycnometer to avoid reintroducing air into the water.
- 15B. Towel dry the outside of the pycnometer/volumetric flask and cover.
- 16B. Determine and record the mass of the pycnometer/volumetric flask, cover, de-aired water, and sample to the nearest 0.1 g. within 10 ± 1 minutes of completion of Step 11. Designate this mass as "E."

Procedure – Mixtures Containing Uncoated Porous Aggregate

If the pores of the aggregates are not thoroughly sealed by a bituminous film, they may become saturated with water during the vacuuming procedure, resulting in an error in maximum density. To determine if this has occurred, complete the general procedure and then:

- 1. Carefully drain water from sample through a towel held over the top of the container to prevent loss of material.
- 2. Spread sample in a flat shallow pan and place before an electric fan to remove surface moisture.
- 3. Determine the mass of the sample when the surface moisture appears to be gone.
- 4. Continue drying and determine the mass of the sample at 15-minute intervals until less than a 0.5 g loss is found between determinations.
- 5. Record the mass as the saturated surface dry mass to the nearest 0.1 g. Designate this mass as " A_{SSD} ".
- 6. Calculate, as indicated below, G_{mm}, using "A" and "A_{SSD}", and compare the two values.

Calculation

Calculate the G_{mm} to three decimal places as follows:

Bowl Procedure

$$G_{mm} = \frac{A}{A - C}$$
 or $G_{mm} = \frac{A}{A_{SSD} - C}$

(for mixes containing uncoated aggregate materials)

where:

A = mass of dry sample in air, g A_{SSD} = Mass of saturated surface dry sample in air, g C = submerged weight of sample in water, g

Example:

$$A = 1432.7 g$$

 $A_{SSD} = 1434.2 g$
 $C = 848.6 g$

$$G_{mm} = \frac{1432.7 \ g}{1432.7 \ g - 848.6 \ g} = 2.453$$
 or $G_{mm} = \frac{1432.7 \ g}{1434.2 \ g - 848.6 \ g} = 2.447$

Pycnometer/Volumetric Flask Procedure

$$G_{mm} = rac{A}{A+D-E}$$
 or $G_{mm} = rac{A}{A_{SSD}+D-E}$

(for mixtures containing uncoated materials)

where:

A = Mass of dry sample in air, g

A_{SSD} = Mass of saturated surface-dry sample in air, g

D = Mass of pycnometer/volumetric flask filled with water at 25°C (77°F), g, determined during the Standardization of Pycnometer/Volumetric Flask procedure

E = Mass of pycnometer/volumetric flask filled with water and the test sample at test temperature, g

Example (in which two increments of a large sample are averaged):

Increment 1	Increment 2
A = 2200.3 g	A = 1960.2 g
D = 7502.5 g	D = 7525.5 g
E = 8812.0 g	E = 8690.8 g
Temperature = 26.2°C	Temperature = 25.0°C
$G_{mm_1} =$	$\frac{2200.3 \ g}{2200.3 \ g + 7502.5 \ g - 8812.0 \ g} = 2.470$
$G_{mm_2} =$	$\frac{1960.2 \ g}{1960.2 \ a + 7525.5 \ a - 8690.8 \ a} = 2.466$

Allowable variation is: 0.014

2.470 - 2.466 = 0.004, which is < 0.014, so they can be averaged.

Average

$$4.936 \div 2 = 2.468$$

Theoretical Maximum Density

To calculate the theoretical maximum density at 25°C (77°F) use one of the following formulas. The density of water at 25°C (77°F) is 997.1 in Metric units or 62.245 in English units.

Theoretical maximum density kg/m³ = $G_{mm} \times 997.1 \text{ kg/m}^3$

$$2.468 \times 997.1 \text{ kg/m}^3 = 2461 \text{ kg/m}^3$$

or

Theoretical maximum density lb/ft³ = $G_{mm} \times 62.245$ lb/ft³

$$2.468 \times 62.245 \text{ lb/ft}^3 = 153.6 \text{ lb/ft}^3$$

Report

- Results on forms approved by the agency
- Sample ID
- \mathbf{G}_{mm} to three decimal places
- Theoretical maximum density to 1 kg/m³ (0.1 lb/ft³)

Performance Exam Checklist

FOP for AASHTO T 209

Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt (HMA) Paving Mixtures

Part	ticipant Name Exam Date		
Reco	cord the symbols "P" for passing or "F" for failing on each step of the chec	klist.	
Prod	ocedure Element	Trial 1	Trial 2
1.	Sample reduced to correct size?		
2.	Particles carefully separated insuring that aggregate is not fractured?		
3.	After separation, fine aggregate particles not larger than 6.3 mm ($\frac{1}{4}$ in)?		
4.	Sample at room temperature?		
5.	Mass of container determined to 0.1 g?		
6.	Mass of sample and container determined to 0.1 g?		
7.	Mass of sample calculated and conforms to required size?		
8.	Water at approximately 25°C (77°F) added to cover sample?		
9.	Entrapped air removed using partial vacuum for 15 ± 2 min?		
10.	Container and contents agitated continuously by mechanical device or r by vigorous shaking at intervals of about 2 minutes?	nanually	
11.	Bowl determination:		
	a. Water bath filled to the overflow level?		
	b. Bowl and contents suspended in water at $25 \pm 1^{\circ}$ C (77 ± 2° F) for 10 ± 1 minutes?		
	c. Submerged weight of bowl and contents determined to 0.1 g?		·
	d. Submerged weight of empty bowl determined to 0.1 g?		
	e. Net submerged weight of contents calculated?		

Prod	edu	re Element	Trial 1	Trial 2	
12.	Py	cnometer/Volumetric Flask determination:			
	a. Pycnometer/volumetric flask filled with water without reintroducing air into the sample?				
	b.	Contents stabilized at 25 ± 1 °C (77 ± 2°F)			
d. Mass of filled pycnometer/volumetric flask and cover determined to 0.1g , $10 \pm 1 \text{minutes}$ after removal of entrapped air completed?					
	e.	Mass of pycnometer/volumetric flask, cover, and water obtained from the Standardization of Pycnometer or Volumetric Flask procedure?			
13.	G_{m}	m calculated correctly and reported to 0.001?			
14.	14. Density calculated correctly and reported to 1 kg/m³ (0.1 lb/ft³)?				
First	: Att	empt: Pass Fail Second Attempt: Pass Fail			
Signa	ature	e of Examiner WAQTC #:			
Com	men	its:			

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.

¹ This Test Method is based on AASHTO T 231-05.

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 ¼ 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) think. 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). 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 in² (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 ½ in (13 mm). In no case shall the recess in the plate be deeper than ½ 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 ½ 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 ½ 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 buildup 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.

Cylinder Compressive Strength MPa (psi)	Minimum Strength of Capping Material	Maximum Average Thickness of Cap	Maximum Thickness Any Part of Cap
500 to 7,000 psi (3.5 to 50 MPa)	5,000 psi (35 MPa) or cylinder strength, whichever is greater	0.25 in (6 mm)	0.31 in (8 mm)
greater than 7,000 psi (50 MPa)	Compressive strength not less than cylinder strength, except as provided in Section 5.1.1.	0.125 in (3 mm)	0.20 in (5 mm)

Compressive Strength and Maximum Thickness of Capping Materials Table 1

- 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 7000 psi (50 MPa), 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 0.002 in (0.05 mm),
 - 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 2 in (50 mm) 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.

Note-Manufacture	er: Testing Supplies Co.
Capping Material	: Super Strong AAA-Sulfur mortar
Lot: 12a45 E	Date Tested: 11/3/98
Signed by:	and Responsible Official)

Item	Capping Material	Control Cylinders	Ratio	Criteria	Pass/Fail
Concrete Cylinder Test Data Type of Capping Material	Sulfur	Ground			
Average Concrete Strength, MPa (psi) Standard Deviation MPa (psi)	76.2 (11,061) 2.59 (376)	75.9 (11,008) 1.72 (250)	1.005 1.504	>0.98 Xc ≤1.57 C	Pass Pass
Number of cylinders tested	15	15			
Cap age when cylinders tested	7 days	n/a			
Capping Material Test Data Average cap thickness mm (in)	2.8 (0.11)	n/a			
Compressive strength of 50 mm (2 in) cubes, MPa (psi)	91 (12,195)				
Cube age when tested	7 days				
Maximum concrete strength qualified, MPa (psi) aNominally a specified strength of 75 MPa (11,000 psi) and perhaps somewhat higher.			1.2 Av.	Str = 91.5 (1	3,273)a

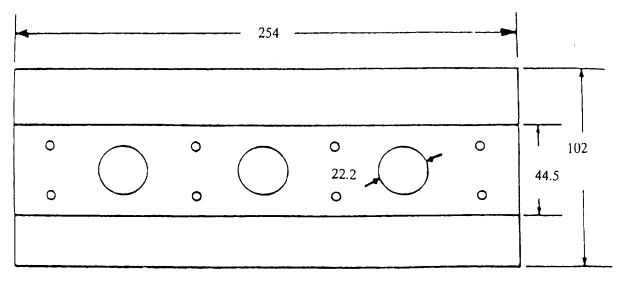
Sample Report of Qualifications of a Capping Material Table 2

- 5.1.3 The compressive strength of capping materials shall be determined by testing 2 in (50 mm) 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.

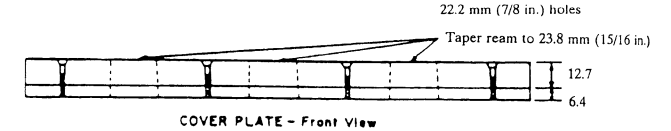
- 5.2 Neat Hydraulic Cement Paste:
 - 5.2.1 Make qualification test of the neat hydraulic cement paste prior to use for capping to establish the effects of water-cement ratio and age on compressive strength of 2 in (50 mm) cubes (Note 2).
 - **Note 2:** The cements used generally conform to Specification 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 2 in (50 mm) cubes. Retarders may be used to extend working time, but their effects on 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 (5,000 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.
- 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 30°C (68 to 86°F), 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-sulfur mortar in the pot within a range of 129 to 143°C (265 to 290°F), 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).
 - **Note 7:** If desired, a plane phenol formaldehyde (bakelite) plate of 0.125 in (3 mm) 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.



COVER PLATE - Plan View



Note: All dimensions are in millimeters unless otherwise indicated.

Dimensional Equivalents						
mm	6.4	12.7	22.2	44.5	102	254
in.	1/4	1/2	7/8	1 3/4	4	10

FIGURE 1 Sketch of Cover for 50-mm (2-in.) Cube Mold

Dimensional Equivalents							
mm	6.4	12.7	22.2	44.5	100	250	
in	1/4	1/2	7/8	1¾	4	10	
Notes: All dimensions shown in millimeters unless otherwise noted. Sketch of Cover for 2 in (50 mm) Cube Mold							

Figure 1

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 ½ in (3 mm) thick, and in no instance shall any part of a cap be more than ½ 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 0.125 in (3 mm) (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 ¼ 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 0.002 in (0.05 mm).

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.

A1.2Determination 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 Terreltype 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 = 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

Part	ticipant Name	Exam Date _			
Pro	cedure Element			Yes	No
1.	The tester has a copy of the current procedure on hand?				
2.	All equipment is functioning according to the test procedu current calibration/verification tags present?	re, and if requ	ired, has the		
Sul	fur Only				
1.	Is the temperature taken and the mortar stirred at the begin	ining of the op	eration?		
2.	Is the capping plate lightly oiled prior to use?				
Pro	cedure Element (cont.)				
3.	Are perpendicularity guides or leveling devices used effect	tively?			
4.	Are caps checked for planeness?				
	a. If yes, how often?				
5.	Are cylinders kept moist after capping?				
Firs	et Attempt: Pass Fail Second A	ttempt: Pass	Fail		
Sign	nature of Examiner				
Com	nments:				

WSDOT Errata to FOP for AASHTO T 255

Total Evaporable Moisture Content of Aggregate by Drying

WAQTC FOP for AASHTO T 255 has been adopted by WSDOT with the following changes:

Sample Preparation

TABLE 1 Sample Sizes for Moisture Content of Aggregate – *Shall include the note below.*

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 percent of the materials.

WSDOT Errata to FOP for AASHTO T 255

Total Evaporable Moisture Content of Aggregate by Drying

Scope

This procedure covers the determination of moisture content of aggregate in accordance with AASHTO T 255-00. It may also be used for other construction materials.

Overview

Moisture content is determined by comparing the wet mass of a sample and the mass of the sample after drying to constant mass. The term constant mass is used to define when a sample is dry.

Constant mass – the state at which a mass does not change more than a given percent, after additional drying for a defined time interval, at a required temperature.

Apparatus

- Balance or scale: Capacity sufficient for the principle sample mass, accurate to 0.1 percent of sample mass or readable to 0.1 g, meeting the requirements of AASHTO M 231.
- · Containers: clean, dry and capable of being sealed
- Suitable drying containers
- Microwave safe container with ventilated lids
- · Heat source, controlled
 - Forced draft oven
 - Ventilated oven
 - Convection oven
- · Heat source, uncontrolled
 - Infrared heater, hot plate, fry pan, or any other device/method that will dry the sample without altering the material being dried
 - Microwave oven (900 watts minimum)
- · Hot pads or gloves
- Utensils such as spoons

Sample Preparation

In accordance with the FOP for AASHTO T 2 obtain a representative sample in its existing condition. The representative sample size is based on Table 1 or other information that may be specified by the agency.

Sample Sizes for Moisture Content of Aggregate			
Nominal Maxim	minal Maximum Size* mm (in) Minimum Sample Mass g (lb		
150	(6)	50,000	(110)
100	(4)	25,000	(55)
90	(3½)	16,000	(35)
75	(3)	13,000	(29)
63	(2½)	10,000	(22)
50	(2)	8000	(18)
37.5	(1½)	6000	(13)
25.0	(1)	4000	(9)
19.0	(3/4)	3000	(7)
12.5	(1/2)	2000	(4)
9.5	(3/8)	1500	(3.3)
4.75	(No. 4)	500	(1.1)

Table 1 Sample Sizes for Moisture Content of Aggregate

Immediately seal or cover samples to prevent any change in moisture content or follow the steps in "Procedure."

Procedure

Determine all masses to the nearest 0.1 percent of the sample mass or to the nearest 0.1 g.

When determining the mass of hot samples or containers or both, place and tare a buffer between the sample container and the balance. This will eliminate damage to or interference with the operation of the balance or scale.

- 1. Determine and record the mass of the container (and lid for microwave drying).
- 2. Place the wet sample in the container.
 - a. For oven(s), hot plates, infrared heaters, etc.: Spread the sample in the container.
 - b. For microwave oven: Heap sample in the container; cover with ventilated lid.
- 3. Determine and record the total mass of the container and wet sample.
- 4. Determine and record the wet mass of the sample by subtracting the container mass determined in Step 1 from the mass of the container and sample determined in Step 3.

^{*}One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size.

- 5. Place the sample in one of the following drying apparatus:
 - a. Controlled heat source (oven): at 110 ± 5 °C (230 ± 9 °F).
 - b. Uncontrolled heat source (Hot plate, infrared heater, etc.): Stir frequently to avoid localized overheating.
- 6. Dry until sample appears moisture free.
- 7. Determine mass of sample and container.
- 8. Determine and record the mass of the sample by subtracting the container mass determined in Step 1 from the mass of the container and sample determined in Step 7.
- 9. Return sample and container to the heat source for additional drying.
 - a. Controlled (oven): 30 minutes
 - b. Uncontrolled (Hot plate, infrared heater, etc.): 10 minutes
 - c. Uncontrolled (Microwave oven): 2 minutes

Caution: Some minerals in the sample may cause the aggregate to overheat, altering the aggregate gradation.

- 10. Determine mass of sample and container.
- 11. Determine and record the mass of the sample by subtracting the container mass determined in Step 1 from the mass of the container and sample determined in Step 10.
- 12. Determine percent change by subtracting the new mass determination (M_n) from the previous mass determination (M_p) divide by the previous mass determination (M_p) multiply by 100.
- 13. Continue drying, performing steps 9 through 12, until there is less than a 0.10 percent change after additional drying time.
- 14. Constant mass has been achieved, sample is defined as dry.
- 15. Allow the sample to cool. Determine and record the total mass of the container and dry sample.
- 16. Determine and record the dry mass of the sample by subtracting the mass of the container determined in Step 1 from the mass of the container and sample determined in Step 15.
- 17. Determine and record percent moisture by subtracting the final dry mass determination (M_D) from the initial wet mass determination (M_W) divide by the final dry mass determination (M_D) multiply by 100.

Heat Source	Specific Instructions	Drying intervals to achieve constant mass (minutes)
Controlled:		
Forced Draft Oven (preferred), Ventilated Oven, or Convection Oven	110 ± 5°C (230 ± 9°F)	30
Uncontrolled:		
Hot plate, Infrared heater, etc.	Stir frequently	10
Microwave	Heap sample and cover with ventilated lid	2

Table 2 Methods of Drving

Calculation

Constant Mass:

Calculate constant mass using the following formula:

$$\frac{M_p - M_n}{M_n} \times 100 = \% \text{ Change}$$

Where:

 M_p = previous mass measurement M_n = new mass measurement

Example:

Mass of container: 1232.1 g

Mass of container after first drying cycle: 2637.2 g

Mass, M_p , of possibly dry sample: 2637.2 g - 1232.1 g = 1405.1 g

Mass of container and dry sample after second drying cycle: 2634.1 g

Mass, M_n , of dry sample: 2634.1 g - 1232.1 g = 1402.0 g

$$\frac{1405.1 \,\mathrm{g} - 1402.0 \,\mathrm{g}}{1405.1 \,\mathrm{g}} \times 100 = 0.22\%$$

0.22 percent is not less than 0.10 percent, so continue drying

Mass of container and dry sample after third drying cycle: 2633.0 g

Mass, M_n , of dry sample: 2633.0 g - 1232.1 g = 1400.9 g

$$\frac{1402.0 \text{ g} - 1400.9 \text{ g}}{1402.0 \text{ g}} \times 100 = 0.08\%$$

0.08 percent is less than 0.10 percent, so constant mass has been reached

Moisture Content:

Calculate the moisture content, w, as a percent, using the following formula:

$$\frac{M_W - M_D}{M_D} \times 100 = \% Moisture Content$$

Where:

$$M_W$$
 = wet mass M_D = dry mass

Example:

Mass of container: 1232.1 g

Mass of container and wet sample: 2764.7 g

Mass, M_W , of wet sample: 2764.7 g - 1232.1 g = 1532.6 g

Mass of container and dry sample (COOLED): 2633.0 g

Mass, M_D , of dry sample: 2633.0 g - 1232.1 g = 1400.9 g

$$w = \frac{1532.6g - 1400.9g}{1400.9g} \times 100 = \frac{131.7g}{1400.9g} = 9.40\%$$
 rounded to 9.4%

Report

- · Results on forms approved by the agency
- Sample ID
- M_W, wet mass
- M_D, dry mass
- w, moisture content to nearest 0.1 percent

Performance Exam Checklist FOP for AASHTO T 255 Total Moisture Content of Aggregate By Drying

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Representative sample of appropriate mass obtained?		
2.	Mass of container determined to 0.1 percent or 0.1 g?		
3.	Sample placed in container and wet mass determined to 0.1 percent or 0.1 g?		
4.	Test sample mass conforms to the required mass?		
5.	Wet mass of sample determined to 0.1 percent or 0.1 g?		
6.	Loss of moisture avoided prior to mass determination?		
7.	Sample dried by a suitable heat source?		
8.	If aggregate heated by means other than a controlled oven, is sample stirred to avoid localized overheating?		
9.	Is aggregate heated for the additional, specified time (forced draft, ventilated, convection – 30 minutes; microwave – 2 minutes; other – 10 minutes) and then mass determined and compared to previous mass – showing less than 0.10 percent loss?		
10.	Sample cooled prior to dry mass determination to 0.1 percent or 0.1 g?		
11.	Calculations performed properly and results reported to the nearest 0.1 percent?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
Com	ments:		

FOP for AASHTO T 255

Total Evaporable Moisture Content of Aggregate by Drying

FOP for AASHTO T 265

Laboratory Determination of Moisture Content of Soils

Scope

This procedure covers the determination of moisture content of aggregate and soil in accordance with AASHTO T 255-00 and AASHTO T 265-15. It may also be used for other construction materials.

Overview

Moisture content is determined by comparing the wet mass of a sample and the mass of the sample after drying to constant mass. The term constant mass is used to define when a sample is dry.

Constant mass – the state at which a mass does not change more than a given percent, after additional drying for a defined time interval, at a required temperature.

Apparatus

- Balance or scale: capacity sufficient for the principle sample mass, accurate to 0.1 percent of sample mass or readable to 0.1 g, and meeting the requirements of AASHTO M 231
- · Containers, clean, dry and capable of being sealed
- · Suitable drying containers
- · Microwave safe container with ventilated lid
- · Heat source, controlled:
 - Forced draft oven
 - Ventilated oven
 - Convection oven
- · Heat source, uncontrolled:
 - Infrared heater/heat lamp, hot plate, fry pan, or any other device/method that will dry the sample without altering the material being dried
 - Microwave oven (900 watts minimum)
- Utensils such as spoons
- · Hot pads or gloves

Sample Preparation

In accordance with the FOP for AASHTO T 2 obtain a representative sample in its existing condition.

For aggregates the representative sample size is based on Table 1 or other information that may be specified by the agency.

 Table 1
 Sample Sizes for Moisture Content of Aggregate

Nominal Maximum Size* mm (in)		Minimum Sam	ple Mass g (lb)
4.75	(No. 4)	500	(1.1)
9.5	(3/8)	1500	(3.3)
12.5	(½)	2000	(4)
19.0	(3/4)	3000	(7)
25.0	(1)	4000	(9)
37.5	(1½)	6000	(13)
50	(2)	8000	(18)
63	(2½)	10,000	(22)
75	(3)	13,000	(29)
90	(3½)	16,000	(35)
100	(4)	25,000	(55)
150	(6)	50,000	(110)

^{*}One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum.

For soils the representative sample size is based on Table 2 or other information that may be specified by the agency.

 Table 2
 Sample Sizes for Moisture Content of Soil

Maximum Particle Size mm (in)		Minimum Sample Mass g (lb)
0.425	(No. 40)	10
4.75	(No. 4)	100
12.5	(½)	300
25.0	(1)	500
50	(2)	1000

Immediately seal or cover samples to prevent any change in moisture content or follow the steps in "Procedure."

Procedure

Determine and record the sample mass as follows:

- For aggregate, determine and record all masses to the nearest 0.1 percent of the sample mass or to the nearest 0.1 g.
- For soil, determine and record all masses to the nearest 0.1 g.

When determining the mass of hot samples or containers or both, place and tare a buffer between the sample container and the balance. This will eliminate damage to or interference with the operation of the balance or scale.

- 1. Determine and record the mass of the container (and lid for microwave drying).
- 2. Place the wet sample in the container.
 - a. For oven(s), hot plates, infrared heaters, etc.: Spread the sample in the container.
 - b. For microwave oven: Heap sample in the container; cover with ventilated lid.
- 3. Determine and record the total mass of the container and wet sample.
- 4. Determine and record the wet mass of the sample by subtracting the container mass determined in Step 1 from the mass of the container and sample determined in Step 3.
- 5. Place the sample in one of the following drying apparatus:
 - a. For aggregate
 - i. Controlled heat source (oven): at 110 ± 5 °C (230 ± 9 °F).
 - ii Uncontrolled heat source (Hot plate, infrared heater, etc.): Stir frequently to avoid localized overheating.
 - b. For soil controlled heat source (oven): at 110 ± 5 °C (230 ± 9 °F).

Note 1: Soils containing gypsum or significant amounts of organic material require special drying. For reliable moisture contents dry these soils at 60°C (140°F). For more information see AASHTO T 265, Note 2.

- 6. Dry until sample appears moisture free.
- 7. Determine mass of sample and container.
- 8. Determine and record the mass of the sample by subtracting the container mass determined in Step 1 from the mass of the container and sample determined in Step 7.
- 9. Return sample and container to the heat source for additional drying.
 - a. For aggregate
 - i. Controlled heat source (oven): 30 minutes
 - ii. Uncontrolled heat source (Hot plate, infrared heater, etc.): 10 minutes
 - iii. Uncontrolled heat source (Microwave oven): 2 minutes

Caution: Some minerals in the sample may cause the aggregate to overheat, altering the aggregate gradation.

b. For soil - controlled heat source (oven): 1 hour

- 10. Determine mass of sample and container.
- 11. Determine and record the mass of the sample by subtracting the container mass determined in Step 1 from the mass of the container and sample determined in Step 10.
- 12. Determine percent change by subtracting the new mass determination (M_n) from the previous mass determination (M_p) divide by the previous mass determination (M_p) multiply by 100.
- 13. Continue drying, performing steps 9 through 12, until there is less than a 0.10 percent change after additional drying time.
- 14. Constant mass has been achieved, sample is defined as dry.
- 15. Allow the sample to cool. Immediately determine and record the total mass of the container and dry sample.
- 16. Determine and record the dry mass of the sample by subtracting the mass of the container determined in Step 1 from the mass of the container and sample determined in Step 15.
- 17. Determine and record percent moisture by subtracting the final dry mass determination (M_D) from the initial wet mass determination (M_W) divide by the final dry mass determination (M_D) multiply by 100.

Table 3 Methods of Drying

	Acquacata			
Heat Source	Aggregate Specific Instructions	Drying intervals to achieve constant mass (minutes)		
Controlled: Forced draft (preferred), ventilated, or convection oven	110 ± 5°C (230 ± 9°F)	30		
Uncontrolled: Hot plate, infrared heater, etc.	Stir frequently	10		
Microwave	Heap sample and cover with ventilated lid	2		
	Soil			
Heat Source	Specific Instructions	Drying increments (minutes)		
Controlled: Forced draft (preferred), ventilated, or convection oven	110 ± 5°C (230 ± 9°F)	1 hour		

Calculation

Constant Mass:

Calculate constant mass using the following formula:

$$\frac{M_p - M_n}{M_p} \times 100 = \% \text{ Change}$$

Where:

 M_p = previous mass measurement M_n = new mass measurement

Example:

Mass of container: 1232.1 g

Mass of container and sample after first drying cycle: 2637.2 g

Mass, M_p , of possibly dry sample: 2637.2 g - 1232.1 g = 1405.1 g

Mass of container and dry sample after second drying cycle: 2634.1 g

Mass, M_n , of dry sample: 2634.1 g - 1232.1 g = 1402.0 g

$$\frac{1405.1 \, g - 1402.0 \, g}{1405.1 \, g} \times 100 = 0.22\%$$

0.22 percent is not less than 0.10 percent, so continue drying

Mass of container and dry sample after third drying cycle: 2633.0 g

Mass, M_n , of dry sample: 2633.0 g - 1232.1 g = 1400.9 g

$$\frac{1402.0 \ g - 1400.9 \ g}{1402.0 \ g} \times 100 = 0.08\%$$

0.08 percent is less than 0.10 percent, so constant mass has been reached.

Moisture Content:

Calculate the moisture content, as a percent, using the following formula:

$$w = \frac{M_W - M_D}{M_D} \times 100$$

Where:

w = moisture content, percent M_W = wet mass M_D = dry mass

Example:

Mass of container: 1232.1 g

Mass of container and wet sample: 2764.7 g

Mass, M_W , of wet sample: 2764.7 g - 1232.1 g = 1532.6 g

Mass of container and dry sample (COOLED): 2633.5 g

Mass, M_D , of dry sample: 2633.5 g - 1232.1 g = 1401.4 g

$$w = \frac{1532.6 \ g - 1401.4 \ g}{1401.4 \ g} \times 100 = \frac{131.2 g}{1401.4 \ g} \times 100 = 9.36\% \ report \ 9.4\%$$

Report

- · Results on forms approved by the agency
- · Sample ID
- M_w, wet mass
- M_D, dry mass
- w, moisture content to nearest 0.1 percent

Performance Exam Checklist FOP for AASHTO T 265 Laboratory Determination of Moisture Content of Soils

Part	icipant Name Exam Date		
Reco	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Prod	cedure Element	Trial 1	Trial 2
1.	Representative sample of appropriate mass obtained?		
2.	Mass of container determined to 0.1 g?		
3.	Sample placed in container and mass determined to 0.1 g?		
4.	Test sample mass conforms to the required mass?		
5.	Wet sample mass determined to 0.1 g?		
6.	Loss of moisture avoided prior to mass determination?		
7.	Sample dried by a suitable heat source?		
8.	If aggregate heated by means other than a controlled oven, is sample stirred to avoid localized overheating?		
9.	For aggregate: Is aggregate heated for the additional, specified time (forced draft, ventilated, convection – 30 minutes; microwave – 2 minutes; other 10 minutes) and then mass determined and compared to previous mass showing less than 0.10 percent loss?		
10.	For soil: Is soil heated for at least 1hour additional drying time and then mass determined and compared to previous mass - showing less than 0.10 percent loss?		
11.	Sample cooled, dry mass determined and recorded to the nearest 0.1 percent?		
12.	Moisture content calculated correctly and recorded to the nearest 0.1 percent?		
First	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
Com	ments:		

FOP for AASHTO T 272

One-Point Method For Determining Maximum Dry Density And Optimum Moisture

Scope

This procedure provides for a rapid determination of the maximum dry density and optimum moisture content of a soil sample, using a one-point determination in accordance with AASHTO T 272-16. This procedure is related to the FOPs for AASHTO T 99/T 180 and R 75.

One-point determinations are made by compacting the soil in a mold of a given size with a specified rammer dropped from a specified height and then compared to an individual moisture/density curve (FOP for AASHTO T 99 or T 180) or a family of curves (FOP for AASHTO R 75). Four alternate methods – A, B, C, and D – are used and correspond to the methods described in the FOP for AASHTO T 99/T 180. The method used in AASHTO T 272 must match the method used for the reference curve or to establish the family of curves. For example, when moisture-density relationships as determined by T 99 - Method C are used to form the family of curves or an individual moisture density curve, then T 99 - Method C must be used to for the one-point determination.

Apparatus

See the FOP for AASHTO T 99/T 180. Use the method matching the individual curve or Family of Curves. Refer to Table 1 of the FOP for AASHTO T 99/T 180 for corresponding mold size, number of layers, number of blows, and rammer specification for the various test methods.

Sample

Sample size determined according to the FOP for AASHTO T 310. In cases where the existing individual curve or family cannot be used a completely new curve will need to be developed and the sample size will be determined by the FOP for AASHTO T 99/T 180.

If the sample is damp, dry it until it becomes friable under a trowel. Drying may be in air or by use of a drying apparatus maintained at a temperature not exceeding 60°C (140°F). Thoroughly break up aggregations in a manner that avoids reducing the natural size of individual particles.

Procedure

- 1. Determine the mass of the clean, dry mold. Include the base plate, but exclude the extension collar. Record the mass to the nearest 1 g (0.005 lb).
- 2. Thoroughly mix the sample with sufficient water to adjust moisture content to 80 to 100 percent of the anticipated optimum moisture.
- 3. Form a specimen by compacting the prepared soil in the mold (with collar attached) in approximately equal layers. For each layer:
 - a. Spread the loose material uniformly in the mold.
 - **Note 1**: It is recommended to cover the remaining material with a non-absorbent sheet or damp cloth to minimize loss of moisture.
 - b. Lightly tamp the loose material with the manual rammer or other similar device, this establishes a firm surface.
 - c. Compact each layer with uniformly distributed blows from the rammer.
 - d. Trim down material that has not been compacted and remains adjacent to the walls of the mold and extends above the compacted surface.
- 4. Remove the extension collar. Avoid shearing off the sample below the top of the mold. The material compacted in the mold should not be over 6 mm (¼ in) above the top of the mold once the collar has been removed.
- 5. Trim the compacted soil even with the top of the mold with the beveled side of the straightedge.
- 6. Determine the mass of the mold and wet soil to the nearest 1 g (0.005 lb) or better.
- 7. Determine the wet mass of the sample by subtracting the mass in Step 1 from the mass in Step 6.
- 8. Calculate the wet density as indicated below under "Calculations."
- 9. Extrude the material from the mold. For soils and soil-aggregate mixtures, slice vertically through the center and take a representative moisture content sample from one of the cut faces, ensuring that all layers are represented. For granular materials, a vertical face will not exist. Take a representative sample. This sample must meet the sample size requirements of the test method used to determine moisture content.
- 10. Determine the moisture content of the sample in accordance with the FOP for AASHTO T 255/T 265.

Calculations

1. Calculate the wet density, in kg/m³ (lb/ft³), by dividing the wet mass by the measured volume of the mold (T 19).

Example - Methods A or C mold:

Wet mass = 2.0055 kg (4.42 lb)

Measured volume of the mold = $0.0009469 \text{ m}^3 (0.03344 \text{ ft}^3)$

Wet Density =
$$\frac{2.0055 \, kg}{0.0009469 \, m^3}$$
 = 2118 kg/m³

Wet Density =
$$\frac{4.42 \text{ lb}}{0.03344 \text{ ft}^3}$$
 = 132.2 lb/ft³

2. Calculate the dry density as follows.

$$\rho_d = \left(\frac{\rho_w}{w + 100}\right) \times 100 \quad or \quad \rho_d = \frac{\rho_w}{\left(\frac{w}{100}\right) + 1}$$

Where:

 ρ_d = Dry density, kg/m³ (lb/ft³) ρ_w = Wet density, kg/m³ (lb/ft³) w = Moisture content, as a percentage

Example:

$$\rho_w = 2118 \text{ kg/m}^3 (132.2 \text{ lb/ft}^3)$$

 $w = 13.5\%$

$$\rho_d = \left(\frac{2118 \, kg/m^3}{13.5 + 100}\right) \times 100 = 1866 \, kg/m^3 \quad \rho_d = \left(\frac{132.2 \, lb/ft^3}{13.5 + 100}\right) \times 100 = 116.5 \, lb/ft^3$$

or

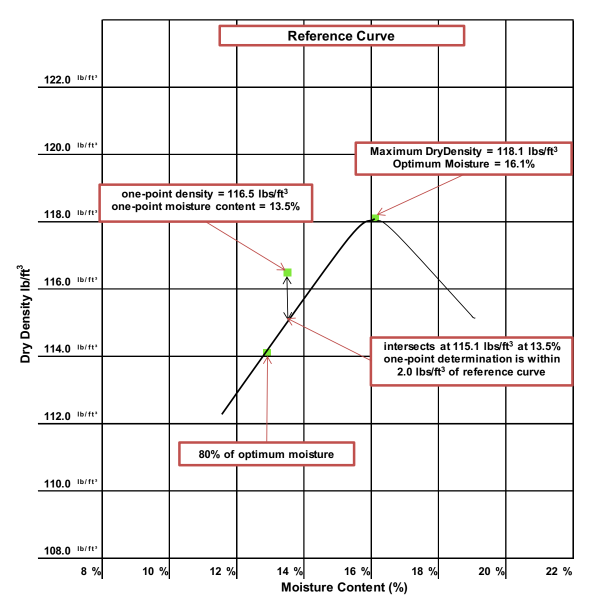
$$\rho_d = \left(\frac{2118 \, kg/m^3}{\frac{13.5}{100} + 1}\right) = 1866 \, kg/m^3 \, \rho_d = \left(\frac{132.2 \, lb/ft^3}{\frac{13.5}{100} + 1}\right) = 116.5 \, lb/ft^3$$

Maximum Dry Density and Optimum Moisture Content Determination Using an Individual Moisture / Density Curve

- 1. The moisture content must be within 80 to 100 percent of optimum moisture of the reference curve. Compact another specimen, using the same material, at an adjusted moisture content if the one-point does not fall in the 80 to 100 percent of optimum moisture range.
- 2. If the moisture content of the one-point determination is not within 80 to 100 percent of the optimum moisture content, compact another specimen, using the same material, at adjusted moisture content.
- 3. Plot the one-point, dry density on the vertical axis and moisture content on the horizontal axis, on the reference curve graph.

- 4. If the one-point falls on the reference curve or within ± 2.0 lbs/ft³, then the maximum dry density and optimum moisture content determined by the curve can be used.
- 5. Perform a full moisture-density relationship if the one-point does not fall on or within \pm 2.0 lbs/ft³ of the reference curve at 80 to 100 percent optimum moisture.

Example

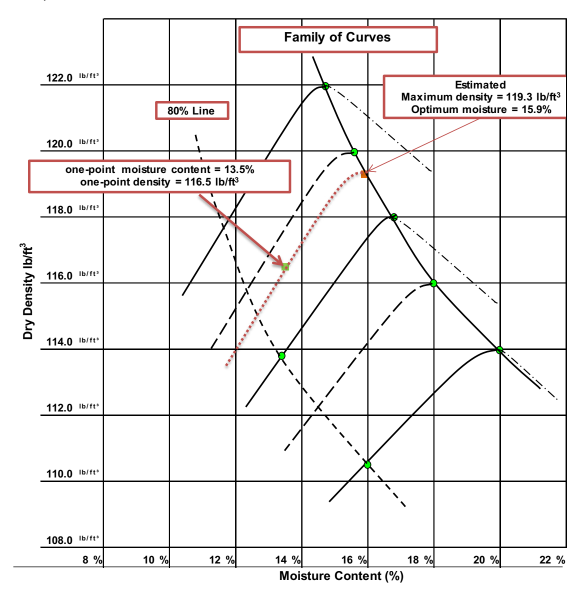


The results of a one-point determination were 116.5 lb/ft³ at 13.5 percent moisture. The point was plotted on the reference curve graph. The one-point determination is within 2.0 lb/ft³ of the point on the curve that corresponds with the moisture content.

Maximum Dry Density and Optimum Moisture Content Determination Using a Family of Curves

- 1. If the moisture-density 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 is used.
- 2. If the moisture-density one-point falls within the family of curves but not on an existing curve, draw a new curve through the plotted single point, parallel and in character with the nearest existing curve in the family of curves. Use the maximum dry density and optimum moisture content as defined by the new curve.
- 3. The one-point must fall either between or on the highest or lowest curves in the family. If it does not, then a full curve must be developed.
- 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 content, compact another specimen, using the same material, at an adjusted moisture content that will place the one point within this range.
- 5. If the new curve through a one-point is not well defined or is in any way questionable, perform a full moisture-density relationship to correctly define the new curve and verify the applicability of the family of curves.
 - **Note 2:** New curves drawn through plotted single point determinations shall not become a permanent part of the family of curves until verified by a full moisture-density procedure following the FOP for AASHTO T 99/T 180.





The results of a one-point determination were 116.5 lb/ft³ at 13.5 percent moisture. The point was plotted on the reference curve graph. The point was plotted on the appropriate family between two previously developed curves near and intermediate curve.

The "dotted" curve through the moisture-density one-point was sketched between the existing curves. A maximum dry density of 119.3 lb/ft³ and a corresponding optimum moisture content of 15.9 percent were estimated.

Report

- · Results on forms approved by the agency
- Sample ID
- Maximum dry density to the closest 1 kg/m³ (0.1 lb/ft³)
- Optimum moisture content to the closest 0.1 percent
- · Reference curve or Family of Curves used

Performance Exam Checklist FOP for AASHTO T 272 (T 99) One-Point Method

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	One-point determination of dry density and corresponding moisture content made in accordance with the FOP for AASHTO T 99?		
	a. Correct size (4.75 mm/No. 4 or 19.0 mm / ¾ in) material used?		
2.	If necessary, sample dried until friable in air or drying apparatus, not exceeding 60°C (140°F)?		
3.	Sample broken up and an adequate amount sieved over the appropriate sieve (4.75 mm / No. 4 or 19.0 mm / $\frac{3}{4}$ in) to determine oversize (coarse particle) percentage?		
4.	Sample passing the sieve has appropriate mass?		
5.	Layer of soil (approximately one third compacted depth) placed in mold with collar attached, loose material lightly tamped?		
6.	Soil compacted with appropriate number of blows (25 or 56)?		
7.	Material adhering to the inside of the mold trimmed?		
8.	Layer of soil (approximately two thirds compacted depth) placed in mold with collar attached, loose material lightly tamped?		
9.	Soil compacted with appropriate number of blows (25 or 56)?		
10.	Material adhering to the inside of the mold trimmed?		
11.	Mold filled with soil such that compacted soil will be above the mold, loose material lightly tamped?		
12.	Soil compacted with appropriate number of blows (25 or 56)?		
13.	Collar removed without shearing off sample?		
14.	Approximately 6 mm ($\frac{1}{4}$ in) of compacted material above the top of the mold (without the collar)?		
15.	Soil trimmed to top of mold with the beveled side of the straightedge?		
16.	Mass of mold and contents determined to appropriate precision?		
17.	Wet density calculated from the wet mass?		
18.	Soil removed from mold using a sample extruder if needed?		
19.	Soil sliced vertically through center (non-granular material)?		
20.	Moisture sample removed ensuring all layers are represented?		
21.	Moist mass determined immediately to 0.1 g?		
22.	Moisture sample mass of correct size?		

Pro	cedure Element	Trial 1	Trial 2
23.	Sample dried and water content determined according to the FOP for T 255/T 265?		
24.	One-point plotted on family of curves supplied?		
25.	One-point falls within 80 to 100 percent of optimum moisture content in order to be valid?		
26.	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?		
27.	Maximum dry density and corresponding optimum moisture content correctly estimated?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
Com	ments:		

Performance Exam Checklist ONE-POINT METHOD FOP FOR AASHTO T 272 (T 180)

Part	icipant Name Exam Date				
Rec	Record the symbols "P" for passing or "F" for failing on each step of the checklist.				
Pro	cedure Element	Trial 1	Trial 2		
1.	One-point determination of dry density and corresponding moisture content made in accordance with the FOP for AASHTO T 180?				
	a. Correct size (4.75 mm / No. 4 or 19.0 mm / ¾ in.) material used?				
2.	If necessary, sample dried until friable in air or drying apparatus, not exceedin 60° C (140°F)?	ng			
3.	Sample broken up and an adequate amount sieved over the appropriate sieve $(4.75 \text{ mm} / \text{No. 4 or } 19.0 \text{ mm} / \frac{3}{4} \text{ in.})$ to determine oversize (coarse particle) percentage?	e			
4.	Sample passing the sieve has appropriate mass?				
5.	Mold placed on rigid and stable foundation?				
6.	Layer of soil (approximately one fifth compacted depth) placed in mold with collar attached, loose material lightly tamped?				
7.	Soil compacted with appropriate number of blows (25 or 56)?				
8.	Material adhering to the inside of the mold trimmed?				
9.	Layer of soil (approximately two fifths compacted depth) placed in mold with collar attached, loose material lightly tamped?				
10.	Soil compacted with appropriate number of blows (25 or 56)?				
11.	Material adhering to the inside of the mold trimmed?				
12.	Layer of soil (approximately three fifths compacted depth) placed in mold wit collar attached, loose material lightly tamped?	h			
13.	Soil compacted with appropriate number of blows (25 or 56)?				
14.	Material adhering to the inside of the mold trimmed?				
15.	Layer of soil (approximately four fifths compacted depth) placed in mold with collar attached, loose material lightly tamped?				
16.	Soil compacted with appropriate number of blows (25 or 56)?				
17.	Material adhering to the inside of the mold trimmed?				
18.	Mold filled with soil such that compacted soil will be above the mold, loose material lightly tamped?				
19.	Soil compacted with appropriate number of blows (25 or 56)?				
20.	Collar removed without shearing off sample?				

Pro	cedure Element	Trial 1	Trial 2
21.	Approximately 6 mm ($\frac{1}{4}$ in) of compacted material above the top of the mold (without the collar)?		
22.	Soil trimmed to top of mold with the beveled side of the straightedge?		
23.	Mass of mold and contents determined to appropriate precision?		
24.	Wet density calculated from the wet mass?		
25.	Soil removed from mold using a sample extruder if needed?		
26.	Soil sliced vertically through center (non-granular material)?		
27.	Moisture sample removed ensuring all layers are represented?		
28.	Moist mass determined immediately to 0.1 g?		
29.	Moisture sample mass of correct size?		
30.	Sample dried and water content determined according to the FOP for T 255/T 265?		
31.	One-point plotted on family of curves supplied?		
32.	One-point falls within 80 to 100 percent of optimum moisture content in order to be valid?		
33.	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?		
34.	Maximum dry density and corresponding optimum moisture content correctly estimated?		
First	t Attempt: Pass Fail Second Attempt: Pass Fail		
Signa	ature of Examiner WAQTC #:		
_			

Comments:

Performance Exam Checklist

Participant Name

Determining Minimum Laboratory Soil Resistivity AASHTO T 288 Checklist

					<u></u>					
Procedure Element										
Laboratory method of Determining Minimum Resistivity							Yes	No		
1.	Sample dried at 140 F, and screened through # 10 sieve?									
2.	Quartered or split out 1500 grams of passing #10 material?									
3.	150 ml of distilled water added to the 1500 gram and thoroughly mixed?									
4.	Sample covered with a wet cloth and allow to stabilize or cure for 12 hours?									
5.	Sample placed & compacted in soil box in layers and the excess trimmed off with a straightedge?									
6.	Resistivity measured with the instrument?									
7.	Soil removed and retained from box and 100 ml of distilled water added and thoroughly mixed?									
8.	Soil box cleaned with distilled water?									
9.	Repeat procedure by increasing moisture content by 100 ml until minimum resistivity can be established?									
10.	Record the lowest value during the repeated measurements?									
11.	Report the resistivity reading.									
Firs	t Attempt:	Pass	Fail		Second Attempt:	Pass	Fail			
Signature of Examiner										
Com	ments:									

Exam Date

WSDOT FOP for AASHTO T 3041

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 Section 9).

Note: WSDOT Specifications require Method A.

- 1.2.2 See the Section 5 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

2.1 AASHTO Standards

T 84 Specific Gravity and Absorption of Fine Aggregate

2.1 WSDOT Standards

T 2 FOP for AASHTO for the Sampling of Aggregates

R 76 FOP for AASHTO for Reducing Field Samples of Aggregates to Testing Size

T 27/T 11 FOP for WAQTC for the Sieve Analysis of Fine and Coarse Aggregates

¹This FOP is based on AASHTO T 304-11 and has been modified per WSDOT standards. To view the redline modifications, contact the WSDOT Quality Systems Manager at 360-709-5412.

2.1 ASTM Standards

В	88	Specification for Seamless Copper water Tube
В	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 Minera 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 Terminology¹

3. Terminology

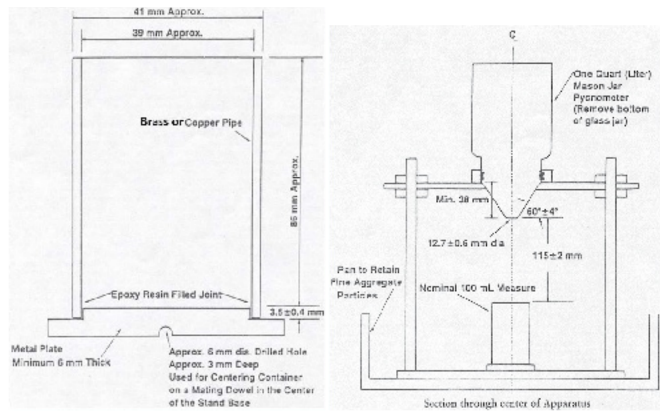
3.1 Terms used in this standard are defined in ASTM C 125 or ACI 116R.

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, the percent void content is determined directly and the average value from two runs is reported.

5. Significance and Use

- 5.1 Methods A 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 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.



Nominal 100-ml Cylindrical Measure Figure 1

Suitable Funnel Stand Apparatus With Cylindrical Measure in Place Figure 2

- 5.3 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 Method A will be useful as an indicator of properties such as 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 brass or 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 $60 \pm 4^{\circ}$ 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).
 - *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).
- 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. The pan shall not be warped so as to prevent rocking of the apparatus during testing.
- 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 edge of the spatula blade is used to strike off the fine aggregate.
- 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 and FOP for AASHTO R 76, or from sieve analysis samples used for FOP for WAQTC/AASHTO T 27/11, or from aggregate extracted from a bituminous concrete specimen. For Method A, 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 and then dried and sieved into separate size fractions according to FOP for WAQTC/AASHTO T 27/11 procedures. Maintain the necessary size fractions obtained from one (or more) sieve analysis in a dry condition in separate containers for each size.

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:

$$V = 1000 \, \frac{M}{D}$$

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.

Individual Size Fraction				
Passing	Retained On	Mass, g		
No. 8 (2.36 mm)	No. 16 (1.18 mm)	44		
No. 16 (1.18 mm)	No. 30 (600 um)	57		
No. 30 (600 um)	No. 50 (300 um)	72		
No. 50 (300 um)	No. 100 (150 um)	17		
	Total	190		

The tolerance on each of these amounts is ± 0.2 g.

- 9.2 Method B Individual Size Fractions 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. 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 1 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 Section 11).
- 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 = volume of cylindrical measure, mL;

F = net mass, g, of fine aggregate in measure (gross mass minus the mass of the empty measure);

G = Bulk dry specific gravity of fine aggregate; and

U = uncompacted voids, percent, in the material

For the standard Graded Sample (Method A), calculate the average uncompacted voids for the two determinations and report the result as U_s.

12. Report

- 12.1 For the Standard Graded Sample (Method A) report:
 - 12.1.1 The Uncompacted Voids (U_s) in percent to the nearest 1 percent.
 - 12.1.2 The specific gravity value used in the calculations.
- 12.2 Report the results using one or more of the following:
 - Materials Testing System (MATS)
 - DOT Form 350-161
 - Form approved in writing by the State Materials Engineer

13. Precision and Bias

See AASHTO T 304 for precision and bias.

Performance Exam Checklist

Uncompacted Void Content of Fine Aggregate FOP AASHTO T 304

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?

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.

- Field sample obtained per FOP for AASHTO T 2?
- 2. Sample reduced to testing size per FOP for AASHTO R 76?
- Sample washed over No. 100 or No. 200 sieve in accordance with FOP for WAQTC/AASHTO T 27/11?
- 4. Sample dried to constant weight?
- 5. Standard Graded sample achieved per FOP for WAQTC/AASHTO T 27/11?
- 6. Necessary size fractions obtained, maintained in a dry condition in separate containers for each size?
- 7. Standard Graded sample-weighed out and combined per Section 9.1, FOP for AASHTO T 304?

Procedure Element

Procedure (Method A)

N*ote:* If Bulk Dry Specific Gravity is unknown, determine it on the minus No. 4 (4.75 mm) material according to AASHTO T 84.

Yes No

- 1. Test sample mixed until it appears to be homogeneous?
- 2. Jar and funnel section positioned in stand and cylindrical measure centered on stand?
- 3. Finger used to block the opening of the funnel?
- 4. Test sample poured into the funnel and leveled?
- 5. Finger removed and sample allowed to fall freely into cylindrical measure?
- 6. After funnel empties, is excess material struck off w/single pass of upright spatula?
- 7. Was care taken to avoid any vibration or disturbance that could cause compaction of material?
- 8. All adhering grains brushed off before weighing the cylindrical measure?
- 9. Mass of the cylindrical measure and contents weighed to nearest 0.1 gram?
- 10. All fine aggregate particles retained and re-homogenized for a second test run?
- 11. Percent (%) of Uncompacted Voids calculated for each run, as per FOP for AASHTO T 304, Method A?
- 12. Were the results for each run averaged for a final result?
- 13. Was the (%) percent of Uncompacted voids reported to the nearest one percent (1%)?
- 14. All calculations performed correctly?

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 1,000 pound-force, and an accuracy of ± 2.5 1,000 pound-force.

Section 6.3.3:

The LVDTs will be inside the test chamber.

Section 6.3.3.2 through Section 6.3.3:

Note 2 Following Section 6.3.1, Section 6.3.3.2, 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 road 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 1,000 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 Errata to FOP for AASHTO T 308

Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method

WAQTC FOP for AASHTO T 308 has been adopted by WSDOT with the following changes:

Procedure - Method B (External Balance) - Method not recognized by WSDOT.

Annex - Correction Factors

Asphalt Binder and Aggregate

Asphalt binder correction factor – Shall read as below:

A correction factor must be established by testing a set of correction specimens for each Job Mix Formula (JMF).

Aggregate correction factor - Method not recognized by WSDOT.

Procedure

Steps 9 - 13 not recognized by WSDOT.

FOP for AASHTO T 308

Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method

Scope

This procedure covers the determination of asphalt binder content of hot mix asphalt (HMA) by ignition of the binder in accordance with AASHTO T 308-16.

Overview

The sample is heated in a furnace at 538°C (1000°F) or less; samples may be heated by convection or direct infrared irradiation (IR). The aggregate remaining after burning can be used for sieve analysis using the FOP for AASHTO T 30.

Some agencies allow the use of recycled HMA. When using recycled HMA, check with the agency for specific correction procedures.

Asphalt binder in the HMA is ignited in a furnace. Asphalt binder content is calculated as the percentage difference between the initial mass of the HMA and the mass of the residual aggregate, with the asphalt binder correction factor, and moisture content subtracted. The asphalt binder content is expressed as percent of moisture-free mix mass.

Two methods, A and B, are presented.

Apparatus

Note 1: The apparatus must be calibrated for the specific mix design. See "Correction Factors" at the end of this FOP.

There are two methods – A and B. The apparatus for the two methods are the same except that the furnace for Method A has an internal balance.

• Ignition Furnace: A forced-air ignition furnace that heats the specimens by either the convection or direct IR irradiation method. The convection-type furnace must be capable of maintaining the temperature at 538 ± 5 °C (1000 ± 9 °F).

For Method A, the furnace will be equipped with an internal scale thermally isolated from the furnace chamber and accurate to 0.1 g. The scale shall be capable of determining the mass of a 3500 g sample in addition to the sample baskets. A data collection system will be included so that mass can be automatically determined and displayed during the test. The furnace shall have a built-in computer program to calculate the 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 binder content, test time, and test temperature. The furnace shall provide an audible alarm and indicator light when the sample mass loss does not exceed 0.01 percent of the total sample mass for three consecutive minutes. Perform lift test according to manufacturer's instructions weekly during use.

Note 2: The furnace shall be designed to permit the operator to change the ending mass loss percentage from 0.01 percent to 0.02 percent.

For both Method A and Method B, the furnace chamber dimensions shall be adequate to accommodate a 3500 g sample. The furnace door shall be equipped so that it cannot be opened during the ignition test. A method for reducing furnace emissions shall be provided and the furnace shall be vented so that no emissions escape into the laboratory. The furnace shall have a fan to pull air through the furnace to expedite the test and to eliminate the escape of smoke into the laboratory.

- Sample Basket Assembly: consisting of sample basket(s), catch pan, and basket guards. Sample basket(s) will be of appropriate size allowing samples to be thinly spread and allowing air to flow through and around the sample particles. Sets of two or more baskets shall be nested. A 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. Basket guards will completely enclose the basket and be made of screen mesh, perforated stainless steel plate, or other suitable material.
- Thermometer, or other temperature measuring device, with a temperature range of 10 -260°C (50-500°F).
- Oven capable of maintaining 110 ± 5°C (230 ± 9°F).
- Balance or scale: Capacity sufficient for the sample mass and conforming to the requirements of M 231, Class G2.
- Safety equipment: Safety glasses or face shield, high temperature gloves, long sleeved jacket, a heat resistant surface capable of withstanding 650°C (1202°F), a protective cage capable of surrounding the sample baskets during the cooling period, and a particle mask for use during removal of the sample from the basket assembly.
- Miscellaneous equipment: A pan larger than the sample basket(s) for transferring sample after ignition, spatulas, bowls, and wire brushes.

Sampling

- 1. Obtain samples of HMA in accordance with the FOP for AASHTO T 168.
- 2. Reduce HMA samples in accordance with the FOP for AASHTO R 47.
- 3. 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 $110 \pm 5^{\circ}$ C ($230 \pm 9^{\circ}$ F) until soft enough.
- 4. Test sample size shall conform to the mass requirement shown in Table 1.
 - **Note 3:** When the mass of the test specimen exceeds the capacity of the equipment used or for large samples of fine mixes, the test specimen may be divided into suitable increments, tested, and the results appropriately combined through a weighted average for calculation of the asphalt binder content.

Table 1

	num Aggregate nm (in)	Minimum Mass Specimen g	Maximum Mass Specimen g
37.5	(1½)	4000	4500
25.0	(1)	3000	3500
19.0	(3/4)	2000	2500
12.5	(1/2)	1500	2000
9.5	(3/8)	1200	1700
4.75	(No. 4)	1200	1700

^{*} One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size.

Procedure - Method A (Internal Balance)

- For the convection-type furnace, preheat the ignition furnace to 538 ± 5°C (1000 ± 9°F) or to the temperature determined in the "Correction Factor" section, Step 9 of this method. Manually record the furnace temperature (set point) prior to the initiation of the test if the furnace does not record automatically. For the direct IR irradiation-type furnace, use the same burn profile as used during the correction factor determination.
- 2. Dry the sample to constant mass, according to the FOP for AASHTO T 329; or determine the moisture content of a companion sample in accordance with the FOP for AASHTO T 329.
- 3. Determine and record the mass to the nearest 0.1 g of the sample basket assembly.
- 4. Evenly distribute the sample in the sample basket assembly, taking care to keep the material away from the edges of the basket. Use a spatula or trowel to level the sample.
- 5. Determine and record the total mass of the sample and sample basket assembly to the nearest 0.1 g. Calculate and record the initial mass of the sample (total mass minus the mass of the sample basket assembly) to the nearest 0.1 g. Designate this mass as (Mi).
- 6. Record the correction factor or input into the furnace controller for the specific HMA.
- 7. Input the initial mass of the sample (Mi) into the ignition furnace controller. Verify that the correct mass has been entered.
 - **CAUTION:** Operator should wear safety equipment high temperature gloves, face shield, fire-retardant shop coat when opening the door to load or unload the sample.
- 8. Open the chamber door and gently set the sample basket assembly in the furnace. Carefully position the sample basket assembly so it is not in contact with the furnace wall. Close the chamber door and verify that the sample mass displayed on the furnace scale equals the total mass of the sample and sample basket assembly recorded in Step 5 within ± 5 g.

Note 4: Furnace temperature will drop below the set point when the door is opened, but will recover when the door is closed and ignition begins. Sample ignition typically increases the temperature well above the set point – relative to sample size and asphalt binder content.

9. Initiate the test by pressing the start button. This will lock the sample chamber and start the combustion blower.

Safety note: Do not attempt to open the furnace door until the asphalt binder has been completely burned off.

- 10. Allow the test to continue until the stable light and audible stable indicator indicate that the change in mass does not exceed 0.01 percent for three consecutive minutes. Press the stop button. This will unlock the sample chamber and cause the printer to print out the test results.
 - **Note 5:** An ending mass loss percentage of 0.02 may be used, if allowed by the agency, when aggregate that exhibits an excessive amount of loss during ignition testing is used.
- 11. Open the chamber door, remove the sample basket assembly, and place on the cooling plate or block. Place the protective cage over the sample basket assembly and allow it to cool to room temperature (approximately 30 minutes).
- 12. Determine and record the total after ignition mass to the nearest 0.1 g. Calculate and record the mass of the sample, after ignition (total after ignition mass minus the mass of the sample basket assembly) to the nearest 0.1 g. Designate this mass as M_f.
- 13. Use the asphalt binder content percentage from the printed ticket. Subtract the moisture content from the printed ticket asphalt binder content and report the difference as the corrected asphalt binder content.
- 14. Asphalt binder content percentage can also be calculated using the formula from "Method B" Step 16.

Calculation

Corrected asphalt binder content:

$$P_b = BC - MC - C_f$$

(if not input in the furnace controller)

where:

 $P_b =$ the corrected asphalt binder content as a percent by mass of the HMA

BC = asphalt binder content shown on printed ticket

MC = moisture content of the companion HMA sample, percent, as determined by the FOP for AASHTO T 329 (if the specimen was oven-dried prior to initiating the procedure, MC=0)

C_f = correction factor as a percent by mass of the HMA sample

Procedure - Method B (External Balance)

- 1. Preheat the ignition furnace to $538 \pm 5^{\circ}\text{C}$ ($1000 \pm 9^{\circ}\text{F}$) or to the temperature determined in the "Correction Factor" section, Step 9 of this method. Manually record the furnace temperature (set point) prior to the initiation of the test if the furnace does not record automatically.
- 2. Dry the sample to constant mass, according to the FOP for AASHTO T 329; or determine the moisture content of a companion sample in accordance with the FOP for AASHTO T 329.
- 3. Determine and record the mass of the sample basket assembly to the nearest 0.1 g.
- 4. Place the sample basket(s) in the catch pan. Evenly distribute the sample in the sample basket(s), taking care to keep the material away from the edges of the basket. Use a spatula or trowel to level the sample.
- 5. Determine and record the total mass of the sample and sample basket assembly to the nearest 0.1 g. Calculate and record the initial mass of the sample (total mass minus the mass of the sample basket assembly) to the nearest 0.1 g. Designate this mass as (M_i).
- 6. Record the correction factor for the specific HMA.
- 7. Open the chamber door and gently set the sample basket assembly in the furnace. Carefully position the sample basket assembly so it is not in contact with the furnace wall. Burn the HMA sample in the furnace for 45 minutes or the length of time determined in the "Correction Factors" section.
- 8. Open the chamber door, remove the sample basket assembly, and place on the cooling plate or block. Place the protective cage over the sample and allow it to cool to room temperature (approximately 30 min).
- 9. Determine and record the total after ignition mass to the nearest 0.1 g. Calculate and record the mass of the sample, after ignition (total after ignition mass minus the mass of the sample basket assembly) to the nearest 0.1 g.
- 10. Place the sample basket assembly back into the furnace.
- 11. Burn the sample for at least 15 minutes after the furnace reaches the set temperature.
- 12. Open the chamber door, remove the sample basket assembly, and place on the cooling plate or block. Place the protective cage over the sample basket assembly and allow it to cool to room temperature (approximately 30 min).
- 13. Determine and record the total after ignition mass to the nearest 0.1 g. Calculate and record the mass of the sample, after ignition (total after ignition mass minus the mass of the sample basket assembly) to the nearest 0.1 g.
- 14. Repeat Steps 10 through 13 until the change in measured mass of the sample after ignition does not exceed 0.01 percent of the previous sample mass after ignition.
 - **Note 6:** An ending mass loss percentage of 0.02 may be used, if allowed by the agency, when aggregate that exhibits an excessive amount of loss during ignition testing is used.

- 15. Determine and record the total after ignition mass to the nearest 0.1 g. Calculate and record the mass of the sample, after ignition (total after ignition mass minus the mass of the sample basket assembly) to the nearest 0.1 g. Designate this mass as M_f.
- 16. Calculate the asphalt binder content of the sample.

Calculations

Calculate the asphalt binder content of the sample as follows:

$$P_b = \frac{M_i - M_f}{M_i} \times 100 - MC - C_f$$

Where:

P_b = the corrected asphalt binder content as a percent by mass of the HMA sample

 M_f = the final mass of aggregate remaining after ignition M_i = the initial mass of the HMA sample prior to ignition MC = moisture content of the companion HMA sample, percent, as determined by the FOP for AASHTO T 329 (if the specimen was oven-dried prior to initiating the procedure, MC=0). C_f = correction factor as a percent by mass of the HMA sample

Example

Correction Factor = 0.42 percent **Moisture Content** = 0.04 percent Initial Mass of Sample and Basket = 5292.7 g= 2931.5 gMass of Basket Assembly M_{i} = 2361.2 g= 5154.4 gTotal Mass after First ignition + basket = 2222.9 gSample Mass after First ignition = 2222.7 gSample Mass after additional 15 min ignition

$$\frac{2222.9 \ g - 2222.7 \ g}{2222.9 \ g} \times 100 = 0.009\%$$

Not greater than 0.01 percent, so M_f = 2222.7 g

$$P_b = \frac{2361.2 \ g - 2222.7 \ g}{2361.2 \ g} \times 100 - 0.42\% - 0.04\% = 5.41\%$$

$$P_h = 5.41$$
 percent

Gradation

- 1. Empty contents of the basket(s) into a flat pan, being careful to capture all material. Use a small wire brush to ensure all residual fines are removed from the baskets.
 - **Note 7:** Particle masks are a recommended safety precaution.
- 2. Perform the gradation analysis in accordance with the FOP for AASHTO T 30.

Report

- · Results on forms approved by the agency
- Sample ID
- Method of test (A or B)
- Corrected asphalt binder content, Pb, per agency standard
- Correction factor, Cf, to 0.01 percent
- Temperature compensation factor (if applicable)
- Total percent loss
- Sample mass
- Moisture content to 0.01 percent
- · Test temperature

Attach the original printed ticket with all intermediate values (continuous tape) to the report for furnaces with internal balances.

Annex – Correction Factors

(Mandatory Information)

Asphalt Binder and Aggregate

Asphalt binder content results may be affected by the type of aggregate in the mixture and by the ignition furnace. Asphalt binder and aggregate correction factors must, therefore, be established by testing a set of correction specimens for each Job Mix Formula (JMF) mix design. Each ignition furnace will have its own unique correction factor determined in the location where testing will be performed.

This procedure must be performed before any acceptance testing is completed, and repeated each time there is a change in the mix ingredients or design. Any changes greater than 5 percent in stockpiled aggregate proportions should require a new correction factor.

Historical data or scientific studies may be used to determine the correction factor(s) in lieu of using this testing procedure if the testing agency provides reference to the studies/data.

All correction samples will be prepared by a central/regional laboratory unless otherwise directed.

Asphalt binder correction factor: A correction factor must be established by testing a set of correction specimens for each Job Mix Formula (JMF). Certain aggregate types may result in unusually high correction factors (> 1.00 percent). Such mixes should be corrected and tested at a lower temperature as described below.

Aggregate correction factor: Due to potential aggregate breakdown during the ignition process, a correction factor will need to be determined for the following conditions:

- a. Aggregates that have a proven history of excessive breakdown
- b. Aggregate from an unknown source.

This correction factor will be used to adjust the acceptance gradation test results obtained according to the FOP for AASHTO T 30.

Procedure

- 1. Obtain samples of aggregate in accordance with the FOP for AASHTO T 2.
- 2. Obtain samples of asphalt binder in accordance with the FOP for AASHTO R 66.
 - **Note 8:** Include other additives that may be required by the JMF.
- 3. Prepare an initial, or "butter," mix at the design asphalt binder content. Mix and discard the butter mix prior to mixing any of the correction specimens to ensure accurate asphalt content.
- 4. Prepare two correction specimens at the JMF design asphalt binder content. Aggregate used for correction specimens shall be sampled from material designated for use on the project. An agency approved method will be used to combine aggregate. An additional "blank" specimen shall be batched and tested for aggregate gradation in accordance with the FOP for AASHTO T 30. The gradation from the "blank" shall fall within the agency specified mix design tolerances.

- 5. Place the freshly mixed specimens directly into the sample basket assembly. If mixed specimens are allowed to cool prior to placement in the sample basket assembly, the specimens must be dried to constant mass according to the FOP for AASHTO T 329. Do not preheat the sample basket assembly.
- 6. Test the specimens in accordance with Method A or Method B of the procedure.
- 7. Once both of the correction specimens have been burned, determine the asphalt binder content for each specimen by calculation or from the printed oven tickets, if available.
- 8. If the difference between the asphalt binder contents of the two specimens exceeds 0.15 percent, repeat with two more specimens and, from the four results, discard the high and low result. Determine the correction factor from the two original or remaining results, as appropriate. Calculate the difference between the actual and measured asphalt binder contents for each specimen to 0.01 percent. The asphalt binder correction factor, C_f, is the average of the differences expressed as a percent by mass of HMA.
- 9. If the asphalt binder correction factor exceeds 1.00 prcent, the test temperature must be lowered to 482 ± 5°C (900 ± 9°F) and new samples must be burned. The temperature for determining the asphalt binder content of HMA samples by this procedure shall be the same temperature determined for the correction samples.
- 10. For the direct IR irradiation-type burn furnaces, the **default** burn profile should be used for most materials. The operator may select burn-profile Option 1 or Option 2 to optimize the burn cycle. The burn profile for testing HMA samples shall be the same burn profile selected for correction samples.
 - **Option 1** is designed for aggregate that requires a large asphalt binder correction factor (greater than 1.00 percent) typically very soft aggregate (such as dolomite).
 - **Option 2** is designed for samples that may not burn completely using the default burn profile.
- 11. Perform a gradation analysis on the residual aggregate in accordance with the FOP for AASHTO T 30, if required. The results will be utilized in developing an "Aggregate Correction Factor" and should be calculated and reported to 0.1 percent.
- 12. From the gradation results subtract the percent passing for each sieve, for each sample, from the percent passing each sieve of the "Blank" specimen gradation results from Step 4.
- 13. Determine the average difference of the two values. If the difference for any single sieve exceeds the allowable difference of that sieve as listed in Table 2, then aggregate gradation correction factors (equal to the resultant average differences) for all sieves shall be applied to all acceptance gradation test results determined by the FOP for AASHTO T 30. If the 75 μm (No. 200) is the only sieve outside the limits in Table 2, apply the aggregate correction factor to only the 75 μm (No. 200) sieve.

Table 2Permitted Sieving Difference

Sieve	Allowable Difference
Sizes larger than or equal to 2.36 mm (No.8)	± 5.0%
Sizes larger than to 75 μm (No.200) and smaller than 2.36 mm (No.8)	± 3.0%
Sizes 75 μm (No.200) and smaller	± 0.5%

Examples

Sieve Siz	ze mm (in)	Correction Factor Blank Sample % Passing	Correction Factor Sample #1 % Passing	Correction Factor Sample #2 % Passing	Difference ½	Avg. Diff.	Sieves to adjust
19.0	(3/4)	100	100	100	0/0	0.0	
12.5	(1/2)	86.3	87.4	86.4	-1.1/-0.1	-0.6	
9.5	(3/8)	77.4	76.5	78.8	+0.9/-1.4	-0.3	
4.75	(No. 4)	51.5	53.6	55.9	-2.1/-4.4	-3.3	
2.36	(No. 8)	34.7	36.1	37.2	-1.4/-2.5	-2.0	
01.18	(No. 16)	23.3	25.0	23.9	-1.7/-0.6	-1.2	
0.600	(No. 30)	16.4	19.2	18.1	-2.8/-1.7	-2.3	
0.300	(No. 50)	12.0	11.1	12.7	+0.9/-0.7	+0.1	
0.150	(No. 100)	8.1	9.9	6.3	-1.8/+1.8	0.0	
75 μm	(No. 200)	5.5	5.9	6.2	-0.4/-0.7	-0.6	- 0.6

In this example, all gradation test results performed on the residual aggregate (FOP for AASHTO T 30) would have an aggregate correction factor applied to the percent passing the 75 μm (No. 200) sieve. The correction factor must be applied because the average difference on the 75 μm (No. 200) sieve is outside the tolerance from Table 2.

In the following example, aggregate correction factors would be applied to each sieve because the average difference on the 4.75 mm (No. 4) is outside the tolerance from Table 2.

Sieve Siz	ze mm (in)	Correction Factor Blank Sample % Passing	Correction Factor Sample #1 % Passing	Correction Factor Sample #2 % Passing	Difference ½	Avg. Diff.	Sieves to adjust
19.0	(3/4)	100	100	100	0/0	0.0	0.0
12.5	(1/2)	86.3	87.4	86.4	-1.1/-0.1	-0.6	-0.6
9.5	(3/8)	77.4	76.5	78.8	+0.9/-1.4	-0.3	-0.3
4.75	(No. 4)	51.5	55.6	57.9	-4.1/-6.4	-5.3	-5.3
2.36	(No. 8)	34.7	36.1	37.2	-1.4/-2.5	-2.0	-2.0
01.18	(No. 16)	23.3	25.0	23.9	-1.7/-0.6	-1.2	-1.2
0.600	(No. 30)	16.4	19.2	18.1	-2.8/-1.7	-2.3	-2.3
0.300	(No. 50)	12.0	11.1	12.7	+0.9/-0.7	+0.1	+0.1
0.150	(No. 100)	8.1	9.9	6.3	-1.8/+1.8	0.0	0.0
75 μm	(No. 200)	5.5	5.9	6.2	-0.4/-0.7	-0.6	-0.6

Performance Exam Checklist

FOP for AASHTO T 308 Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method

Part	icipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Oven at correct temperature $538 \pm 5^{\circ}$ C ($1000 \pm 9^{\circ}$ F) or correction factor temperature?		
	Or: for IR ovens, correct burn profile applied?		
2.	Sample reduced to correct size?		
3.	HMA sample or companion moisture sample taken and dried per FOP for AASHTO T 329?		
4.	Mass of sample basket assembly recorded to 0.1 g?		
5.	With pan below basket(s) sample evenly distributed in basket(s)?		
6.	Sample conforms to the required mass and mass recorded to 0.1 g?		
7.	Method A		
	a. Initial mass entered into furnace controller?		
	b. Sample correctly placed into furnace?		
	c. Test continued until stable indicator signals?		
	d. Uncorrected asphalt binder content obtained on printed ticket?		
	e. Sample mass determined to nearest 0.1 g.?		
8.	Method B		
	a. Sample correctly placed into furnace?		
	b. Sample burned for 45 min or time determined by correction process?		
	c. Sample cooled to room temperature?		
	d. Sample burned to constant mass?		
	e. Sample mass determined to nearest 0.1 g.?		
	f. Uncorrected asphalt binder content calculated correctly and recorded?		
9.	Asphalt binder content corrected for Correction Factor if needed?		
10.	Asphalt binder content corrected for moisture per the FOP for AASHTO T 329 if needed?		

Procedure Element					Irial 1	Irial 2
11. Corrected asphalt	binder content recor	ded?				
12. Contents of the ba	asket(s) carefully emp	tied into a pan?				
First Attempt: Pass	Fail	Second Attempt:	Pass	Fail		
Signature of Examiner			_ WA	AQTC #:		
Comments:						

FOP for AASHTO T 309

Temperature of Freshly Mixed Portland Cement Concrete

Scope

This procedure covers the determination of the temperature of freshly mixed Portland Cement Concrete in accordance with AASHTO T 309-15.

Warning—Fresh Hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.

Apparatus

- Container The container shall be made of non-absorptive material and large enough to provide at least 75 mm (3 in) of concrete in all directions around the sensor; concrete cover must also be a least three times the nominal maximum size of the coarse aggregate.
- Temperature measuring device The temperature measuring device shall be
 calibrated and capable of measuring the temperature of the freshly mixed concrete
 to ± 0.5°C (± 1°F) throughout the temperature range likely to be encountered.
 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.
- Reference temperature measuring device The reference temperature measuring device shall be a thermometric device readable to 0.2°C (0.5°F) 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.

Calibration of Temperature Measuring Device

Each temperature measuring device shall be verified for accuracy annually and whenever there is a question of accuracy. Calibration shall be performed by comparing readings on the temperature measuring device with another calibrated instrument at two temperatures at least 15°C or 27°F apart.

Sample Locations and Times

The temperature of freshly mixed concrete may be measured in the transporting equipment, in forms, or in sample containers, provided the sensor of the temperature measuring device has at least 75 mm (3 in) of concrete cover in all direction around it.

Complete the temperature measurement of the freshly mixed concrete within 5 minutes of obtaining the sample.

Concrete containing aggregate of a nominal maximum size greater than 75 mm (3 in) may require up to 20 minutes for the transfer of heat from the aggregate to the mortar after batching.

Procedure

- 1. Dampen the sample container.
- 2. Obtain the sample in accordance with the FOP for WAQTC TM 2.
- 3. Place sensor of the temperature measuring device in the freshly mixed concrete so that it has at least 75 mm (3 in) of concrete cover in all directions around it.
- 4. Gently press the concrete in around the sensor of the temperature measuring device at the surface of the concrete so that air cannot reach the sensor.
- 5. Leave the sensor of the temperature measuring device in the freshly mixed concrete for a minimum of two minutes, or until the temperature reading stabilizes.
- 6. Complete the temperature measurement of the freshly mixed concrete within 5 minutes of obtaining the sample.
- 7. Read and record the temperature to the nearest 0.5°C (1°F).

Report

- · Results on forms approved by the agency
- Sample ID
- Measured temperature of the freshly mixed concrete to the nearest 0.5°C (1°F).

Performance Exam Checklist FOP for AASHTO T 309 Temperature of Freshly Mixed Concrete

Part	icipant Name	Exam Da	te		
Rec	ord the symbols "P" for passing or "F" fo	or failing on each step of the	checklist.		
Pro	cedure Element			Trial 1	Trial 2
1.	Obtain sample of concrete large enoug of concrete cover around sensor in all		75 mm (3 in)		
2.	Place temperature measuring device in cover around sensor?	n sample with a minimum of	75 mm (3 in)		
3.	Gently press concrete around thermor	neter?			
4.	Read temperature after a minimum of stabilizes?	2 minutes or when tempera	ture reading		
5.	Complete temperature measurement v	within 5 minutes of obtaining	g sample?		
6.	Record temperature to nearest 0.5°C (1°F)?			
Firs	t Attempt: Pass Fail	Second Attempt: Pass	Fail		
Sign	ature of Examiner	WA	AQTC #:		
	checklist is derived, in part, from copyri rican Concrete Institute.	ghted material printed in AC	I CP-1, publis	hed by th	ne
Com	ments:				

WSDOT Errata to FOP for AASHTO T 310

In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)

WAQTC FOP for AASHTO T 310 has been adopted by WSDOT with the following changes:

Procedure

Replace step 1 with below:

- 1. WSDOT requires test location selected per WSDOT SOP 615.
- 6. Place the gauge on the prepared surface so the source rod can enter the hole without disturbing loose material.

Include note below:

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

- 10. Perform one of the following methods, per agency requirements:
 - a. Method A Single Direction: Method not recognized by WSDOT.
- 11. Step not required by WSDOT.

Percent Compaction

Determined using WSDOT SOP 615.

In Diana Danaite and Maintena (to by Nuclear Methods (Shallow Denth)

FOP for AASHTO T 310

In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)

Scope

This procedure covers the determination of density, moisture content, and relative compaction of soil, aggregate, and soil-aggregate mixes in accordance with AASHTO T 310-13. This field operating procedure is derived from AASHTO T 310. The nuclear moisture-density gauge is used in the direct transmission mode.

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 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.
- Sealable containers and utensils for moisture content determinations.

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.

Calibration

Calibrate the nuclear gauge as required by the agency. This calibration may be performed by the agency using manufacturer's recommended procedures or by other facilities approved by the agency. Verify or re-establish calibration curves, tables, or equivalent coefficients every 12 months.

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 Operator's Manual.

Note 1: New standard counts may be necessary more than once a day. See agency requirements.

Overview

There are two methods for determining in-place density of soil/soil aggregate mixtures. See agency requirements for method selection.

- · Method A Single Direction
- Method B Two Direction

Procedure

- 1. Select a test location(s) randomly and in accordance with agency requirements. Test sites should be relatively smooth and flat and meet the following conditions:
 - a. At least 10 m (30 ft) away from other sources of radioactivity
 - b. At least 3 m (10 ft) away from large objects
 - c. The test site should be at least 150 mm (6 in) away from any vertical projection, unless the gauge is corrected for trench wall effect.
- 2. Remove all loose and disturbed material, and remove additional material as necessary to expose the top of the material to be tested.
- 3. Prepare a flat area sufficient in size to accommodate the gauge. Plane the area to a smooth condition so as to obtain maximum contact between the gauge and the material being tested. For Method B, the flat area must be sufficient to permit rotating the gauge 90 or 180 degrees about the source rod.
- 4. Fill in surface voids beneath the gauge with fines of the material being tested passing the 4.75 mm (No. 4) sieve or finer. Smooth the surface with the guide plate or other suitable tool. The depth of the filler should not exceed approximately 3 mm (1/4 in).
- 5. Make a hole perpendicular to the prepared surface using the guide plate and drive pin. The hole shall be at least 50 mm (2 in) deeper than the desired probe depth, and shall be aligned such that insertion of the probe will not cause the gauge to tilt from the plane of the prepared area. Remove the drive pin by pulling straight up and twisting the extraction tool.
- 6. Place the gauge on the prepared surface so the source rod can enter the hole without disturbing loose material.

- 7. Insert the probe in the hole and lower the source rod to the desired test depth using the handle and trigger mechanism.
- 8. Seat the gauge firmly by partially rotating it back and forth about the source rod. Ensure the gauge is seated flush against the surface by pressing down on the gauge corners, and making sure that the gauge does not rock.
- 9. Pull gently on the gauge to bring the side of the source rod nearest to the scaler/detector firmly against the side of the hole.
- 10. Perform one of the following methods, per agency requirements:
 - a. Method A Single Direction: Take a test consisting of the average of two, one-minute readings, and record both density and moisture data. The two wet density readings should be within 32 kg/m³ (2.0 lb/ft³) of each other. The average of the two wet densities and moisture contents will be used to compute dry density.
 - b. Method B Two Direction: Take a one-minute reading and record both density and moisture data. Rotate the gauge 90 or 180 degrees, pivoting it around the source rod. Reseat the gauge by pulling gently on the gauge to bring the side of the source rod nearest to the scaler/detector firmly against the side of the hole and take a one-minute reading. (In trench locations, rotate the gauge 180 degrees for the second test.) Some agencies require multiple one-minute readings in both directions. Analyze the density and moisture data. A valid test consists of wet density readings in both gauge positions that are within 50 kg/m³ (3.0 lb/ft³). If the tests do not agree within this limit, move to a new location. The average of the wet density and moisture contents will be used to compute dry density.
- 11. If required by the agency, obtain a representative sample of the material, 4 kg (9.0 lb) minimum, from directly beneath the gauge full depth of material tested. This sample will be used to verify moisture content and/or identify the correct density standard. Immediately seal the material to prevent loss of moisture.
 - The material tested by direct transmission can be approximated by a cylinder of soil approximately 300 mm (12 in) in diameter directly beneath the centerline of the radioactive source and detector. The height of the cylinder will be approximately the depth of measurement. When organic material or large aggregate is removed during this operation, disregard the test information and move to a new test site.
- 12. To verify the moisture content from the nuclear gauge, determine the moisture content with a representative portion of the material using the FOP for AASHTO T 255/T 265 or other agency approved methods. If the moisture content from the nuclear gauge is within ± 1 percent, the nuclear gauge readings can be accepted. Moisture content verification is gauge and material specific. Retain the remainder of the sample at its original moisture content for a one-point compaction test under the FOP for AASHTO T 272, or for gradation, if required.
 - **Note 2:** Example: A gauge reading of 16.8 percent moisture and an oven dry of 17.7 percent are within the ± 1 percent requirements. Moisture correlation curves will be developed according to agency guidelines. These curves should be reviewed and possibly redeveloped every 90 days.

13. Determine the dry density by one of the following.

- a. From nuclear gauge readings, compute by subtracting the mass (weight) of the water (kg/m³ or lb/ft³) from the wet density (kg/m³ or lb/ft³) or compute using the percent moisture by dividing wet density from the nuclear gauge by 1 + moisture content expressed as a decimal.
- b. When verification is required and the nuclear gauge readings cannot be accepted, the moisture content is determined by the FOP for AASHTO T 255/T 265 or other agency approved methods. Compute dry density by dividing wet density from the nuclear gauge by 1 + moisture content expressed as a decimal.

Percent Compaction

Percent compaction is determined by comparing the in-place dry density as
determined by this procedure to the appropriate agency density standard. For soil
or soil-aggregate mixes, these are moisture-density curves developed using the
FOP for AASHTO T 99/T 180. When using maximum dry densities from the FOP
for AASHTO T 99/T 180 or FOP for AASHTO T 272, it may be necessary to use the
Annex in the FOP for T 99/T 180 to determine corrected maximum dry density and
optimum moisture content.

For coarse granular materials, the density standard may be density-gradation curves developed using a vibratory method such as AKDOT&PF's ATM 212, ITD's T 74, WSDOT's TM 606, or WFLHD's Humphres.

See appropriate agency policies for use of density standards.

Calculation

Wet density readings from gauge: 1963 kg/m³ (121.6 lb/ft³)

1993 kg/m³ (123.4 lb/ft³)

Avg: 1978 kg/m³ (122.5 lb/ft³)

Moisture readings from gauge: 14.2% and 15.4% = Avg 14.8%

Moisture content from the FOP's for AASHTO T 255/ T 265: 15.9%

Moisture content is greater than 1 percent different so the gauge moisture cannot be used.

Calculate the dry density as follows:

$$\rho_d = \left(\frac{\rho_w}{w + 100}\right) \times 100 \quad or \quad \rho_d = \left(\frac{\rho_w}{\frac{w}{100} + 1}\right)$$

Where:

 ρ_d = Dry density, kg/m³ (lb/ft³) ρ_w = Wet density, kg/m³ (lb/ft³)

w = Moisture content from the FOP's for AASHTO T 255/T 265, as a

percentage

$$\rho_d = \left(\frac{1978 \, kg/m^3 \, or \, 122.5 \, lb/ft^3}{15.9 + 100}\right) \times 100 \quad \rho_d = \left(\frac{1978 \, kg/m^3 \, or \, 122.5 \, lb/ft^3}{\frac{15.9}{100} + 1}\right)$$

Corrected for moisture Dry Density: 1707 kg/m³ (105.7 lb/ft³)

Calculate percent compaction as follows:

% Compaction =
$$\frac{\rho_d}{Agency\ density\ standard} \times 100$$

Example:

% Compaction =
$$\frac{105.7 \ lb/ft^3}{111.3 \ lb/ft^3} \times 100 = 95\%$$

Where:

$$ho_d$$
 = Dry density, kg/m³ (lb/ft³)
Agency density standard = Corrected maximum dry density from the FOP from T 99/T 180 Annex

Report

- Results on forms approved by the agency
- Sample ID
- Location of test, elevation of surface, and thickness of layer tested.
- Visual description of material tested.
- Make, model and serial number of the nuclear moisture-density gauge.
- Wet density to 0.1 lb/ft³.
- Moisture content as a percent, by mass, of dry soil mass to 0.1 percent.
- Dry density to 0.1 lb/ft³.
- Density standard to 0.1 lb/ft³.
- · Percent compaction.
- Name and signature of operator.

In-Place Density and Meisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Denth)				
	In Diago Density and Maintens (Contant of Cail and Cail As	unun maka bu Almalanu M	lathada (Challan Danth)

Performance Exam Checklist

FOP for AASHTO T 310

In-Place Density and Moisture Content of Soil and Soil-Aggregate by Nuclear Methods (Shallow Depth)

Part	icip	ant Name Exam Date			
Rec	ord 1	the symbols "P" for passing or "F" for failing on each step of the checklist.			
Pro	cedu	ıre Element	Trial 1	Trial 2	
1.	Ga	uge turned on 10 to 20 minutes before use?			
2.	Ca	libration verified?			
3.		ndard count taken and recorded in accordance with nufacturer's instructions?			
4.		st location selected appropriately 10 m (30 ft) from other radioactive sources, n (10 ft) from large objects, 150 mm (6 in) away from vertical projections?			
5.	Loc	ose, disturbed material removed?			
6.	Fla	t, smooth area prepared?			
7.		rface voids filled with native fines (-No. 4) to 3 mm (½ in) maximum ckness?			
8.	Но	le driven 50 mm (2 in) deeper than probe depth?			
9.					
10.	Мє	ethod A:			
	a.	Gauge firmly seated, and gently pulled back so that the source rod is against the side of the hole toward the scaler/detectors?			
	b.	Two, one-minute reading taken; wet density within 32 kg/m³ (2.0 lb/ft³)?			
	c.	Density and moisture data averaged?			
11.	Мє	ethod B:			
	а.	Gauge firmly seated, and gently pulled back so that the source rod is against the side of the hole toward the scaler/detectors?			
	b.	A minimum of a one-minute reading taken; density and moisture data recorded?			
	c.	Gauge turned 90° or 180° (180° in trench)?			
	d.	Gauge firmly seated, and gently pulled back so that the source rod is against the side of the hole toward the scaler/detectors?			
	e.	A minimum of a one-minute reading taken; density and moisture data recorded?			
	f.	Wet densities within 50 kg/m³ (3.0 lb/ft³)?			
	g.	Density and moisture data averaged?			

Pro	cedure Element			Trial 1	Trial 2
12.	Representative sa	mple (4 kg or 9 lb) obtained from test location?		
13.	Sample sealed imr	mediately to preve	ent moisture loss?		
14.	Moisture content density gauge read	•	ned using other means than the nuclear		
15.	Dry Density calcu	lated using prope	r moisture content?		
16.	Percent compaction	on calculated corr	rectly?		
Firs	t Attempt: Pass	Fail	Second Attempt: Pass Fail		
Sign	ature of Examiner		WAQTC #:		
Com	ments:				

WSDOT Errata to FOP for AASHTO T 312

Asphalt Mixture Specimens by Means of the Superpave Gyratory Compactor

WAQTC FOP for AASHTO T 312 has been adopted by WSDOT with the following changes:

Sample Preparation

Plant Produced Asphalt Mixtures

Replace step 3 with below:

3. Place in the oven set no more than 25° F above the compaction temperature range until the material is within the compaction temperature range.

Compaction Procedure

Replace step 3 with below:

3. 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.

FOP for AASHTO T 312

Asphalt Mixture Specimens by Means of the Superpave Gyratory Compactor

Scope

This procedure covers preparing specimens, using samples of plant produced asphalt mixtures, for determining the mechanical and volumetric properties of asphalt mixtures in accordance with AASHTO T 312-15.

Apparatus

- Superpave Gyratory Compactor (SGC) meeting the requirements of AASHTO T 312
- Molds meeting the requirements of AASHTO T 312
- Chute, mold funnel or both (Optional)
- Scale meeting the requirements of AASHTO M 231 Class G 5
- Oven, thermostatically controlled, capable of maintaining set temperature within ± 3°C (± 5°F)
- Thermometers accurate to ± 1°C (± 2°F) between 10 and 232°C (50 450°F)
 - **Note 1:** Non-Contact thermometers are not acceptable.
- Miscellaneous pans, spoons, spatulas, hot pads, gloves, paper discs, markers, etc.

Equipment Requirements

The calibration shall be performed on the SGC per the Manufacturer's instructions. See agency requirements for the calibration frequency.

The mold and base plate dimensions shall be checked every twelve months or 80 hours of operation to determine that they are within the tolerances listed in AASHTO T 312.

Equipment Preparation

Prepare the equipment in accordance with manufacturer's recommendations. At a minimum preparation includes:

- Warm-up gyratory compactor
- · Verify machine settings
- Internal Angle: 1.16 ± 0.02°
- Ram Pressure: 600 kPa ± 18 kPa
- Number of gyrations

Note 2: The number of gyrations (N_{des}) is obtained from the Job Mix Formula (JMF).

- · Lubricate bearing surfaces
- Prepare recording device as required
- Pre-heat molds and plates at the compaction temperature range (minimum of 30 min) or before reuse reheat (minimum of 5 min)
 - **Note 3:** The use of multiple molds will speed up the compaction process.
- Pre-heat chute, mold funnel, spatulas, and other apparatus (not to exceed the maximum compaction temperature)

Sample Preparation

Laboratory Prepared Asphalt Mixtures

This is a sample produced during the Mix Design process using aggregate and binder that is combined in the laboratory. When designing asphalt mixtures using the gyratory compactor refer to AASHTO T 312.

Plant Produced Asphalt Mixtures

- Determine initial sample size, number of gyrations (N_{des}), and compaction temperature range from the Job Mix Formula (JMF).
- Obtain the sample in accordance with the FOP for AASHTO T 168.
- Reduce the sample in accordance with the FOP for AASHTO R 47.
- The sample size should be such that it results in a compacted specimen that is 115 ± 5mm at the desired number of gyrations.

Note 4: Replicate specimens are generally prepared. Refer to agency requirements.

If the material is not in the compaction temperature range:

- 1. Place the appropriate sample mass into a container.
- 2. Spread to a depth of 1 to 2 in for even heating of mixture.
- 3. Place in the oven until the material is within the compaction temperature range.

Note 5: The material properties may be altered when the times of delivery of the test sample and the placement of the material on the roadway are different.

Compaction Procedure

Follow the manufacturer's recommended loading procedure. This may require the steps below to be performed in a different order. Steps 1 through 8 must be performed before the sample and mold cools below minimum compaction temperature.

- 1. Remove pre-heated mold and plate(s) from the oven (verify mold and plate(s) has been cleaned if previously used).
- 2. Place the base plate and paper disc in bottom of mold.
- 3. Place the mix into the mold in a single lift (care should be taken to avoid segregation or loss of material).
- 4. Level the mix in the mold.
- 5. Place a paper disc and the heated upper plate (if required) on top of the leveled sample.
- 6. Load the mold into the compactor; check settings.
- 7. Start the compaction process.
 - a. Check the pressure ($600 \pm 18 \text{ kPa}$).
 - b. Check the angle $(1.16 \pm 0.02^{\circ})$.
- 8. Upon completion of the compaction process, record the number of gyrations and specimen height.

Note 6: If the specimen is not 115 ± 5 mm follow agency requirements.

- 9. Extrude the specimen from the mold; a brief cooling period may be necessary before fully extruding some specimens to ensure the specimens are not damaged.
 - Note 7: Clean molds after each use.
- 10. Carefully remove the paper discs.
- 11. Cool the compacted specimen to room temperature.
- 12. Identify the specimen with chalk or other marker.

Report

- On forms approved by the agency
- Sample ID
- Number of gyrations
- · Specimen height

Performance Exam Checklist FOP for AASHTO T 312 Gyratory Compaction of Asphalt Mixtures

Part	ticipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Angle, pressure and number of gyrations set?		
2.	Bearing surfaces, rotating base surface, and rollers lubricated?		
3.	Representative sample obtained according to the FOP for AASHTO T 168?		
4.	Sample reduced according to FOP AASHTO R 47?		
5.	Asphalt mixture heated to compaction temperature range?		
6.	Mold, base plate, and upper plate heated to compaction temperature range?		
7.	Mold, base plate, and upper plate (if required) removed from oven and paper disk placed on bottom of mold?		
8.	Mix placed into mold in one lift without segregation?		
9.	Paper disk placed on top of the asphalt mixture?		
10.	Mold placed into compactor and upper plate clamped into place?		
11.	Pressure applied at 600 kPa ± 18 kPa?		
12.	Specified number of gyrations applied?		
13.	Proper angle confirmed from display?		
14.	Compacted specimen removed from mold, paper disc(s) removed, and allowed to cool to room temperature?		
15.	Asphalt mixture sample measured to a height of 115 ± 5 mm at required gyrations?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	ature of Examiner WAQTC #:		
Com	ments:		

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 315/32 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 $\frac{1}{16}$ in (1.6 mm) nor more than $\frac{1}{8}$ 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.
- f. Balance Must conform to ASTM C 1005.
- g. Tapping Stick Made of hardwood, a diameter of ½ in (3.2 mm), and a length of 6 in (152 mm).
- h. Tamper Made of hardwood, a diameter of 7/16 in (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 3½ 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 follwing table:

		Batch Weights (grams)		
U.S.	No. Screen Size	Individual Wt.	Accumulative Wt.	
3⁄₅ in - #4	(9.5 mm - 4.75 mm)	14	14	
#4 - #8	(4.75 mm - 2.36 mm)	205	219	
#8 - #16	(2.36 mm - 1.18 mm)	186	405	
#16 - #30	(1.18 mm - 600 um)	230	635	
#30 - #50	(600 μm - 300 μm)	561	1196	
#50 - minus	(600 µm - minus)	278	1474	

- 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/reflectometer 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. Calibrate and zero the reflectometer, Photovolt Model 577 or equivalent per ASTM E 1347, according to the manufacturer's User Manual.
 - a. Select standard plaque for calibration that is slightly higher than the expected percent reflectance of the samples.

b. Reflectance Determination

1. For testing lane markers, move the unit to determine an average reading. Recheck the standard plaque reading and the zero reading between each sample measurement.

2. Report

a. Report the average result of all readings for a sample to the nearest percent.

WSDOT SOP 318

Standard Operating Procedure for Melting of Flexible Bituminous Pavement Marker Adhesive for Evaluation

1. Scope

- 1.1. This standard covers the handling, cutting, and melting of Flexible Bituminous Adhesive.
- 1.2. 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 establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

- 2.1. ASTM Standards:
 - D 5167 Standard Practice for Melting of Hot-Applied Joint and Crack Sealant and Filler for Evaluation.

3. Significance and Use

3.1. This standard establishes the procedure for handling, cutting and melting of Flexible Bituminous Pavement Marker Adhesives in preparation for the making of test specimens used in the laboratory evaluations of the Flexible Bituminous Pavement Marker Adhesives.

4. Apparatus

- 4.1. Laboratory Melter:
 - An appropriate laboratory melter as described in section 6.1 of ASTM D5167.
- 4.2. Cutting Device:
 - A heated knife or saw capable of cutting a vertical cross section from a solid adhesive sample.

5. Hazards

- 5.1. Use standard safety precautions and Personal Protective Equipment (PPE) when handling hot materials and preparing test specimens.
- 5.2 Prior to handling, cutting, or melting flexible bituminous pavement marker adhesive, a Material Safety Data Sheet (MSDS) must be obtained from the manufacturer so that proper safe handling techniques will be used.

6. Procedure

- 6.1 Cut a complete vertical section from the block of adhesive in order to obtain a uniform representative sample (see figure 1) and to supply enough product to pour all specimens for testing.
- 6.2 To avoid compatibility problems clean the melting container so that it is free of all cleaning solvents and previously melted material.
- 6.3 All segments from one vertical section must be melted in the same melting container. If necessary cut the sample to fit into container.
- 6.4 Heat and maintain the oil bath to $405 \pm 5^{\circ}$ F. Place the sample container into the heating apparatus. Insert paddle for stirring as soon as sample begins to melt. Begin continuous stirring immediately after inserting the paddle. Check the sample temperature frequently. Keep the sample container covered except when checking the sample temperature.
- 6.5 Heat the sample to 398 ± 2°F. Once the sample has become fluid and reached temperature, stop the mechanical stirring device and immediately remove sample container from the melter. Clean oil residue or wrap container with towel to ensure bath oil does not contaminate sample. Pour all required specimens immediately following removal of the sample container from the melter to minimize temperature loss.
- 6.6 Do not reuse sample once melted and split for testing. If retesting is necessary, a new specimen must be cut from the original sample.

Cut vertically through sample block to remove entire cross section for testing.

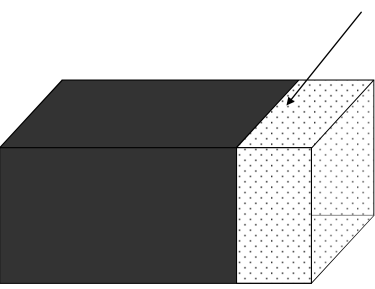


Figure 1

WSDOT Errata to AASHTO T 324

Hamburg Wheel-Track Testing of Compacted Hot Mix Asphalt (HMA)

AASHTO T 324 has been adopted by WSDOT with the following changes:

7. Determining Air Void Content

7.3. Determine the air void content of the specimens in accordance with T 269. The recommended target air void content is 7.0 ± 1.0 percent for laboratory-compacted SGC cylindrical specimens and 7.0 ± 1.0 percent for laboratory-compacted slab specimens. Field specimens may be tested at the air void content at which they are obtained.

8. Procedure

8.6.1. Select a test temperature of 50° C.

Tester Qualification Practical Exam Checklist

Hamburg Wheel-Track Testing of Compacted Hot Mix Asphalt (HMA) FOP for AASHTO T 324

Part	icipant Name Exam Date		
Pro	cedure 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.	Specimen height is 62 ± 1.0 mm (2.44 ± 0.04 in.) or 38.1 mm (1.5 inch) minimum for cores?		
4.	Specimen meets air void tolerance of $7.0 + 1.0 \%$?		
5.	Specimens placed in molds and loaded into trays with a maximum gap of 7.5 mm between molds?		
6.	Tray mounted in machine and securely fastened?		
7.	Sample data and testing parameters entered into computer? (e.g., sample name, agg source, wheel speed, maximum rut depth, number of passes, and water temperature)		
8.	Wheels gently lowered and samples allowed to soak at testing temperature for 30 minutes?		
9.	Wheel tracking device shut off when test parameters are reached?		
10.	Test data obtained for charting and analysis?		
First	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	nature of Examiner		
Com	ments:		

FOP for AASHTO T 329

Moisture Content of Asphalt Mixtures by Oven Method

Scope

This procedure covers the determination of moisture content of asphalt mixtures in accordance with AASHTO T 329-15.

Overview

Moisture content is determined by comparing the wet mass of a sample and the mass of the sample after drying to constant mass. The term constant mass is used to define when a sample is dry.

Constant mass – the state at which a mass does not change more than a given percent, after additional drying for a defined time interval, at a required temperature.

Apparatus

- Balance or scale: 2 kg capacity, readable to 0.1 g and conforming to AASHTO M 231.
- Forced draft, ventilated, or convection oven: Capable of maintaining the temperature surrounding the sample at 163 ± 14 °C (325 ± 25 °F).
- Sample Container: Clean, dry, not affected by heat and of sufficient size to contain a test sample without danger of spilling.
- Thermometer or other suitable device with a temperature range of 10-260°C (50-500°F).

Sample

The test sample shall be obtained in accordance with the FOP for AASHTO T 168, and reduced in accordance with the FOP for AASHTO R 47. The size of the test sample shall be a minimum of 1000 g.

Procedure

- 1. Preheat the oven to the Job Mix Formula (JMF) mixing temperature range. If the mixing temperature is not supplied, a temperature of 163 ± 14 °C (325 ± 25 °F) is to be used.
 - **Note 1:** For repeatability between laboratories, the preferred practice is to dry the sample at no less than 9° C (15° F) below the JMF mixing temperature.
- 2. Determine and record the mass of the sample container, including release media, to the nearest 0.1 g.
 - **Note 2:** When using paper or other absorptive material to line the sample container ensure it is dry before determining initial mass of sample container.
- 3. Place the test sample in the sample container.
- 4. Determine and record the temperature of the test sample.
- 5. Determine and record the total mass of the sample container and test sample to the nearest 0.1 g.

- 6. Calculate the initial, moist mass (M_i) of the test sample by subtracting the mass of the sample container as determined in Step 2 from the total mass of the sample container and the test sample as determined in Step 5.
- 7. The test sample shall be initially dried for 90 ± 5 minutes, and its mass determined. Then it shall be dried at 30 ± 5 min intervals until further drying does not alter the mass by more than 0.05 percent.
- 8. Cool the sample container and test sample to \pm 9°C (\pm 15°F) of the temperature determined in Step 4.
- 9. Determine and record the total mass of the sample container and 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 mass.
- 10. Calculate the final, dry mass (M_f) of the test sample by subtracting the mass of the sample container as determined in Step 2 from the total mass of the sample container and the test sample as determined in Step 9.

Note 4: Moisture content and the number of samples in the oven will affect the rate of drying at any given time. Placing wet samples in the oven with nearly dry samples could affect the drying process.

Calculations

Constant Mass:

Calculate constant mass using the following formula:

$$\%Change = \frac{M_p - M_n}{M_n} \times 100$$

Where:

 M_p = previous mass measurement M_n = new mass measurement

Example:

Mass of container: 232.6 g

Mass of container and sample after first drying cycle: 1361.8 g

Mass, M_p , of possibly dry sample: 1361.8 g - 232.6 g = 1129.2 g

Mass of container and possibly dry sample after second drying cycle: 1360.4 g

Mass, M_p , of possibly dry sample: 1360.4 g - 232.6 g = 1127.8 g

$$\frac{1129.2 \ g - 1127.8 \ g}{1129.2 \ g} \times 100 = 0.12\%$$

0.12 percent is not less than 0.05 percent, so continue drying the sample.

Mass of container and possibly dry sample after third drying cycle: 1359.9 g Mass, M_n , of dry sample: 1359.9 g - 232.6 g = 1127.3 g

$$\frac{1127.8 \ g - 1127.3 \ g}{1127.8 \ g} \times 100 = 0.04\%$$

0.04 percent is less than 0.05 percent, so constant mass has been reached.

Moisture Content:

Calculate the moisture content, as a percent, using the following formula.

$$Moisture\ Content = \frac{M_i - M_f}{M_f} \times 100$$

Where:

 M_i = initial, moist mass M_f = final, dry mass

Example:

$$Moisture\ Content = \frac{1134.9\ g - 1127.3\ g}{1127.3\ g} \times 100 = 0.674, \text{say}\ 0.67\%$$

Where:

 $M_i = 1134.9 g$ $M_f = 1127.3 g$

Report

- · Results on forms approved by the agency
- Sample ID
- Moisture content to 0.01 percent

Performance Exam Checklist FOP for AASHTO T 329 Moisture Content of Asphalt Mixtures by Oven Method

Part	ticipant Name Exam Date		_	
Rec	ord the symbols "P" for passing or "F" for failing on each step of the check	list.		
Pro	cedure Element	Trial 1 Tri	al 2	
1.	Mass of clean dry container including release media determined to 0.1 g	?		
2.	Representative sample obtained; 1000 g minimum?			
3.	Initial temperature taken and recorded?			
4.	Mass of sample determined to 0.1 g?			
5.	Sample placed in drying oven for 90 ± 5 minutes?			
6.	Sample dried at a temperature not to exceed the JMF mixing temp?			
7.	Constant mass checked at 30 ± 5 minute intervals and reached?			
8.	Sample and container cooled to \pm 9°C (15°F) of the initial temperature b final mass determined to 0.1 g?	efore		
9.	Calculation of moisture content performed correctly to 0.01 percent?			
	$Moisture\ Content = rac{M_i - M_f}{M_f} imes 100$			
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail			
Signa	Signature of Examiner WAQTC #:			
Com	ments:			

WSDOT FOP for AASHTO T 331

Bulk Specific Gravity (G_{mb}) and Density of Compacted Hot Mix Asphalt (HMA) Using Automatic Vacuum Sealing Method

AASHTO T 331 has been adopted by WSDOT with the following changes:

6. Procedure

AASHTO R 79 shall be used throughout section 6 in lieu of ASTM D7227/D7227M

Note 3: Laboratory specimens 3000 grams or greater shall be cooled to room temperature for a minimum of 15 hours and a maximum of 24 hours at $77 \pm 9^{\circ}F$ ($25 \pm 5^{\circ}C$).

8. Verification

- 8.1 WSDOT VP 103 shall be used for vacuum system verification.
- 8.2 This section is deleted.

Performance Exam Checklist

Bulk Specific Gravity of Compacted HMA Using Automatic Vacuum Sealing Method FOP for AASHTO T 331

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. Water bath of suitable size to entirely submerge and suspend the specimen with an adequate holder?
- 4. Water bath equipped with an overflow outlet?
- 5. Water bath controlled to 77 ± 1.8 °F (25 ± 1 °C)?
- 6. Plastic bag meets procedure specifications?

Sample Preparation

- 1. Specimen dried to constant mass per AASHTO R 79 or according to T 331 Section 6.1?
- 2. Specimen at room temperature, $77 \pm 9^{\circ}F$ (25 ± 5°C)? Laboratory compacted specimens cooled for 15 24 hours at $77 \pm 9^{\circ}F$ (25 ± 5°C)?
- 3. Sharp edges removed from specimen (recommended)?

Procedure

- 1. Specimen mass, A, determined at room temperature, $77 \pm 9^{\circ}F$ (25 ± 5°C)?
- 2. Appropriate size bag selected, inspected for holes and it's mass determined?
- 3. Sealed dry mass of specimen determine by adding specimen and bag masses together then recorded as B?
- 4. If needed, filler plates added or removed before placing bag inside vacuum chamber and inserting specimen into bag?
- 5. Specimen placed in bag with the smoothest side down?
- 6. End of bag pulled over sample and centered over sealing bar with minimum of 1" overlap?
- 7. Bag wrinkles smoothed out over seal bar just prior to closing lid?
- 8. CorLok operation initiated by closing and latching lid?
- 9. CorLok test cycle allowed to continue until chamber door opens?
- 10. Sealed specimen carefully removed from vacuum chamber without puncturing bag?
- 11. Bag inspected for loose areas which indicate poor seal or bag puncture?
- 12. If needed, test started over because seal ruptured or bag punctured?
- 13. Sealed specimen fully submerged in water bath within 1 minute of vacuum chamber door releasing?
- 14. Bag is not touching the sides of the water bath and no trapped air bubbles exist under specimen?

Procedure (continued)

Yes No

- 15. Mass of sealed specimen underwater, E, at 77 ± 1.8 °F (25 ± 1 °C) recorded as soon as scale stabilizes?
- 16. Specimen removed from bag and mass recorded as C then checked to be no more than 5 grams of the mass recorded as A?
- 17. Process restarted at section 6.1 if test fails section 6.5 check? Section 6.5 check: If difference between C and A are greater than 5 grams the specimen is acceptable if less than 0.08 percent is lost (material loss) or 0.04 percent is gained (from water) as compared to A.
- 18. All calculations performed correctly?

First Attempt:	Pass	Fail	Second Attempt:	Pass	Fail
Signature of Ex	aminer				
Comments:					

WSDOT Errata to FOP for AASHTO T 335

Determining the Percent Fracture in Coarse Aggregate

WAQTC FOP for AASHTO T 335 has been adopted by WSDOT with the following changes:

Sampling and Sample Preparation

4. Method 2 – Individual Sieve Fracture Determination – *Method not recognized by WSDOT*.

FOP for AASHTO T 335

Determining the Percent Fracture in Coarse Aggregate

Scope

This procedure covers the determination of the percentage, by mass, of a coarse aggregate (CA) sample that consists of fractured particles meeting specified requirements in accordance with AASHTO T 335-09.

In this FOP, a sample of aggregate is screened on the sieve separating CA and fine aggregate (FA). This sieve will be identified in the agency's specifications, but might be the 4.75 mm (No. 4) sieve. CA particles are visually evaluated to determine conformance to the specified fracture. The percentage of conforming particles, by mass, is calculated for comparison to the specifications.

Apparatus

- Balance or scale: Capacity sufficient for the principle sample mass, accurate to 0.1 percent of the sample mass or readable to 0.1 g, and meeting the requirements of AASHTO M 231.
- Sieves: Meeting requirements of the FOP for AASHTO T 27/T 11.
- Splitter: Meeting the requirements of FOP for AASHTO R 76.

Terminology

- 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.
- 2. Fractured particle: A particle of aggregate having at least the minimum number of fractured faces specified. (This is usually one or two.)

Sampling and Sample Preparation

- 1. Sample and reduce the aggregate in accordance with the FOPs for AASHTO T 2 and R 76.
- 2. When 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.
- 3. Method 1 Combined Fracture Determination
 - a. Dry the sample sufficiently to obtain a clean separation of FA and CA material in the sieving operation.
 - b. Sieve the sample in accordance with the FOP for AASHTO T 27/T 11 over the 4.75 mm (No. 4) sieve, or the appropriate sieve listed in the agency's specifications for this material.
 - **Note 1:** Where necessary, wash the sample over the sieve designated for the determination of fractured particles to remove any remaining fine material, and dry to a constant mass in accordance with the FOP for AASHTO T 255.

c. Reduce the sample using Method A – Mechanical Splitter, in accordance with the FOP for AASHTO R 76, to the appropriate test size. This test size should be slightly larger than shown in Table 1, to account for loss of fines through washing if necessary.

Table 1 Sample Size Method 1 (Combined Sieve Fracture)

Nominal Maxim	um Size* mm (in)		Cumulative Retained on 4) Sieve g (lb)
37.5	(1½)	2500	(6)
25.0	(1)	1500	(3.5
19.0	(3/4)	1000	(2.5)
12.5	(1/2)	700	(1.5)
9.5	(3/8)	400	(0.9)
4.75	(No. 4)	200	(0.4)

^{*}One sieve larger than the first sieve to retain more than 10 percent of the material using an agency specified set of sieves based on cumulative percent retained. Where large gaps in specification sieves exist, intermediate sieve(s) may be inserted to determine nominal maximum size.

4. Method 2 - Individual Sieve Fracture Determination

- a. Dry the sample sufficiently to obtain a clean separation of FA and CA material in the sieving operation. A washed sample from the gradation determination (the FOP for T 27/T 11) may be used.
- b. If not, sieve the sample in accordance with the FOP for AASHTO T 27 over the sieves listed in the specifications for this material.
 - **Note 2:** If overload (buffer) sieves are used the material from that sieve must be added to the next specification sieve.
- c. The size of test sample for each sieve shall meet the minimum size shown in Table 2. Utilize the total retained sieve mass or select a representative portion from each sieve mass by splitting or quartering in accordance with the FOP for AASHTO R 76.
 - **Note 3:** Where necessary, wash the sample over the sieves designated for the determination of fractured particles to remove any remaining fine material, and dry to a constant mass in accordance with the FOP for AASHTO T 255.

Sieve Size mm (in) Minimum Sample Mass g (lb) 31.5 $(1\frac{1}{4})$ 1500 (3.5)25.0 (1)1000 (2.2)19.0 $(\frac{3}{4})$ 700 (1.5)16.0 (5/8) 500 (1.0)12.5 300 (0.7) $(\frac{1}{2})$ 9.5 $(\frac{3}{8})$ 200 (0.5)6.3 $(\frac{1}{4})$ 100 (0.2)4.75 (No. 4) 100 (0.2)2.36 25 (No. 8) (0.1)2.00 (No. 10) 25 (0.1)

Table 2 Sample Size Method 2 (Individual Sieve Fracture)

Note 4: If fracture is determined on a sample obtained for gradation, use the mass retained on the individual sieves, even if it is less than the minimum listed in Table 2. If less than 5 percent of the total mass is retained on a single specification sieve, include that material on the next smaller specification sieve. If a smaller specification sieve does not exist, this material shall not be included in the fracture determination.

Procedure

- 1. After cooling, spread the dried sample on a clean, flat surface.
- 2. Examine each particle face and determine if the particle meets the fracture criteria.
- 3. Separate the sample into three categories:
 - Fractured particles meeting the criteria
 - Particles not meeting the criteria
 - Questionable or borderline particles
- 4. Determine the dry mass of particles in each category to the nearest 0.1 g.
- 5. Calculate the percent questionable particles.
- 6. Resort the questionable particles when more than 15 percent is present. Continue sorting until there is no more than 15 percent in the questionable category.
- 7. Calculate the percent fractured particles meeting criteria to nearest 0.1 percent. Report to 1 percent.

Calculation

Calculate the mass percentage of questionable particles to the nearest 1 percent using the following formula:

$$\%Q = \frac{Q}{F + O + N} \times 100$$

Where:

%Q = Percent of questionable fractured particles

F = Mass of fractured particles

Q = Mass of questionable or borderline particles

N = Mass of unfractured particles

Example:

Mass of unfractured particles = 632.6 g

Mass of questionable particles = 97.6 g

Mass of unfractured particles = 352.6 g

$$\%Q = \frac{97.6 \ g}{632.6 \ g + 97.6 \ g + 352.6 \ g} \times 100 = 9.0\%$$

Calculate the mass percentage of fractured faces to the nearest 0.1 percent using the following formula:

$$P = \frac{\frac{Q}{2} + F}{F + Q + N} \times 100$$

where:

P = Percent of fracture

F = Mass of fractured particles

Q = Mass of questionable particles

N = Mass of unfractured particles

Example:

Mass of fractured particles = 632.6 g

Mass of questionable particles = 97.6 g

Mass of unfractured particles = 352.6 g

$$P = \frac{\frac{97.6 g}{2} + 632.6 g}{632.6 g + 97.6 g + 352.6 g} \times 100 = 62.9\%$$
 Report 63%

Report

- Results on forms approved by the agency
- · Sample ID
- Fractured particles to the nearest 1 percent.

Performance Exam Checklist FOP for AASHTO T 335 Determining The Percentage of Fracture in Coarse Aggregate

Par	ticipant Name Exam Date		
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Sample properly sieved through specified sieve(s)?		
2.	Sample reduced to correct size?		
3.	Sample dried and cooled, if necessary?		
4.	Particles separated into fractured, unfractured, and questionable categories?		
5.	Dry mass of each category determined to nearest 0.1 g?		
6.	Questionable category resorted if more than 15 percent of total mass falls in that category?		
7.	Fracture calculation performed correctly?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	ature of Examiner WAQTC #:		
Com	ments:		

WSDOT Errata to FOP for AASHTO T 355

In-Place Density of Asphalt Mixtures by Nuclear Method

WAQTC FOP for AASHTO T 355 has been adopted by WSDOT with the following changes:

Material

• Filler material: Not used by WSDOT.

Test Site Location

Replace step 1 with below:

1. WSDOT requires test location selected per WSDOT Test Method 716.

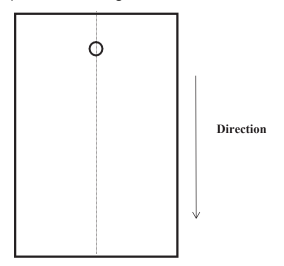
Procedure

Steps 1 - 7 not recognized by WSDOT use Method 2 Below:

Method 2 - One four minute test

- 1. For Thin Layer Mode (Use with Troxler 3450 only) ensure the depth entered into the gauge matches the pavement depth and the depth at which the gauge was correlated or set the gauge depth to Backscatter.
 - Thin Layer or Backscatter are different modes of measurement, ensure that the correct form of measurement is utilized.
- 2. Draw an outline around the entire gauge base for correlation coring.
- 3. Take one 4-minute test and record the wet density reading parallel to the direction of travel.

Method 2 Footprint of The Gauge Test Site Parallel to Direction of Travel



APPENDIX - CORRELATION WITH CORES

Correlation with Cores

Replace step 2 with below:

1. Obtain a pavement core from each of the test sites according to WSDOT SOP 734. The core should be taken from the center of the nuclear gauge footprint.

FOP for AASHTO T 355 IN-PLACE DENSITY OF ASPHALT MIXTURES by Nuclear Method

Scope

This test method describes a procedure for determining the density of asphalt mixtures by means of a nuclear gauge using the backscatter method in accordance with AASHTO T 355-16. Correlation with densities determined under the FOP for AASHTO T 166 is required by some agencies.

Apparatus

- Nuclear density gauge with the factory-matched standard reference block.
- 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 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.

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.

Calibration

Calibrate the nuclear gauge as required by the agency. This calibration may be performed by the agency using the manufacturer's recommended procedures or by other facilities approved by the agency. Verify or re-establish calibration curves, tables, or equivalent coefficients every 12 months.

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, recalibrated, or both.
- 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 Operator's Manual.

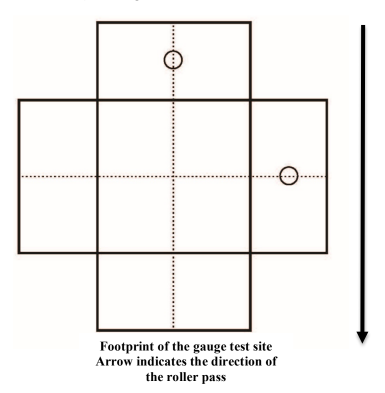
Note 1: New standard counts may be necessary more than once a day. See agency requirements.

Test Site Location

- 1. Select a test location(s) randomly and in accordance with agency requirements. Test sites should be relatively smooth and flat and meet the following conditions:
 - a. At least 10 m (30 ft) away from other sources of radioactivity.
 - b. At least 3 m (10 ft) away from large objects.
 - c. If the gauge will be closer than 600 mm (24 in) to any vertical mass, or less than 300 mm (12 in) from a vertical pavement edge, use the gauge manufacturer's correction procedure.

Procedure

- 1. Maintain maximum contact between the base of the gauge and the surface of the material under test. Use filler material to fill surface voids. Spread a small amount of filler material over the test site surface and distribute it evenly. Strike off the surface with a straightedge (such as a lathe or flat-bar steel) to remove excess material.
- 2. Place the gauge on the test site, perpendicular to the roller passes. Using a crayon (not spray paint), mark the outline or footprint of the gauge. Extend the probe to the backscatter position.
- 3. Take a one-minute test and record the wet density reading.
- 4. Rotate the gauge 90 degrees centered over the original footprint. Mark the outline or footprint of the gauge.
- 5. Take another one-minute test and record the wet density reading.
- 6. If the difference between the two one-minute tests is greater than 40 kg/m³ (2.5 lb/ft³), retest in both directions. If the difference of the retests is still greater than 40 kg/m³ (2.5 lb/ft³) test at 180 and 270 degrees.
- 7. The density reported for each test site shall be the average of the two individual one-minute wet density readings.



Calculation of Results

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.

Example:

Reading #1: 141.5 lb/ft³

Reading #2: 140.1 lb/ft³ Are the two readings within the tolerance? (YES)

Reading average: 140.8 lb/ft^3 Core correction: $+2.1 \text{ lb/ft}^3$ Corrected reading: 142.9 lb/ft^3

From the FOP for

AASHTO T 209: $G_{mm} = 2.466$

Maximum Laboratory Dry Density = $2.466 \times 62.245 lb/ft^3 = 153.5 lb/ft^3$

Percent compaction =
$$\frac{142.9 \, lb/ft^3}{153.5 \, lb/ft^3} \times 100 = 93.1\%$$

Report

- · Results on forms approved by the agency
- Test ID
- · Location of test and thickness of layer tested
- Mixture type
- Make, model and serial number of the nuclear moisture-density gauge
- Calculated wet density of each measurement and any adjustment data
- · Density standard
- Compaction 0.1 percent
- Name and signature of operator

APPENDIX - CORRELATION WITH CORES

(Nonmandatory Information)

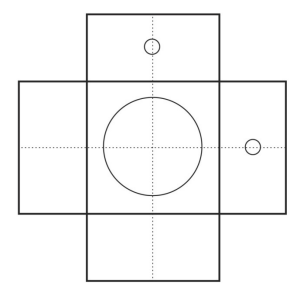
The Bulk Specific Gravity (Gmb) of the core is a physical measurement of the in-place HMA and can be compared with the nuclear density gauge readings. Comparing the core value to the corresponding gauge values, a correlation can be established.

The correlation can then be used to adjust the gauge readings to the in-place density of the cores. The core correlation is gauge specific and must be determined without traffic allowed on the pavement between nuclear density gauge readings and obtaining the core. When using multiple nuclear density gauges each gauge should be correlated to the core locations prior to removal of the core.

When density correlation with the FOP for AASHTO T 166 is required, correlation of the nuclear gauge with pavement cores shall be made on the first day's paving (within 24 hours) or from a test strip constructed prior to the start of paving. Cores must be taken before traffic is allowed on the pavement.

Correlation with Cores

- Determine the number of cores required for correlation from the agency's specifications. Cores shall be located on the first day's paving or on the test strip. Locate the test sites in accordance with the agency's specifications. Follow the "Procedure" section above to establish test sites and obtain densities using the nuclear gauge.
- 2. Obtain a pavement core from each of the test sites according to AASHTO R 67. The core should be taken from the center of the nuclear gauge footprint.



Footprint of the gauge test site. Core location in the center of the footprint.

3. Determine the density of the cores by the FOP for AASHTO T 166, Bulk Specific Gravity of Compacted Asphalt Mixtures Using Saturated Surface Dry Specimens.

- 4. Calculate a correlation factor for the nuclear gauge reading as follows:
 - a. Calculate the difference between the core density and the average nuclear gauge density at each test site to the nearest 1 kg/m^3 (0.1 lb/ft³). Calculate the average difference and standard deviation of the differences for the entire data set to the nearest 1 kg/m^3 (0.1 lb/ft³).
 - b. If the standard deviation of the differences is equal to or less than 40 kg/m³ (2.5 lb/ft³), the correlation factor applied to the average nuclear gauge density shall be the average difference calculated above in 4.a.
 - c. If the standard deviation of the differences is greater than 40 kg/m³ (2.5 lb/ft³), the test site with the greatest variation from the average difference shall be eliminated from the data set and the data set properties and correlation factor recalculated following 4.a and 4.b.
 - d. If the standard deviation of the modified data set still exceeds the maximum specified in 4.b, additional test sites will be eliminated from the data set and the data set properties and correlation factor recalculated following 4.a and 4.b. If the data set consists of less than five test sites, additional test sites shall be established.

Note A1: The exact method used in calculating the nuclear gauge correlation factor shall be defined by agency policy.

Note A2: The above correlation procedure must be repeated if there is a new job mix formula. Adjustments to the job mix formula beyond tolerances established in the contract documents will constitute a new job mix formula. A correlation factor established using this procedure is only valid for the particular gauge and at the probe depth used in the correlation procedure. If another gauge is brought onto the project, it shall be correlated using the same procedure. Multiple gauges may be correlated from the same series of cores if done at the same time.

Note A3: For the purpose of this procedure, a job mix formula is defined as the percent and grade of paving asphalt used with a specified gradation of aggregate from a designated aggregate source. A new job mix formula may be required whenever compaction of the wearing surface exceeds the agency's specified maximum density or minimum air voids.

Core Correlation Example:

	Core results from T 166:	Average Gauge reading:	Difference:	X	X2
1	144.9 lb/ft ³	142.1 lb/ft ³	2.8 lb/ft ³	-0.7	0.49
2	142.8 lb/ft ³	140.9 lb/ft ³	1.9 lb/ft ³	0.2	0.04
3	143.1 lb/ft ³	140.7 lb/ft ³	2.4 lb/ft ³	-0.3	0.09
4	140.7 lb/ft ³	138.9 lb/ft ³	1.8 lb/ft ³	0.3	0.09
5	145.1 lb/ft ³	143.6 lb/ft ³	1.5 lb/ft ³	0.6	0.36
6	144.2 lb/ft ³	142.4 lb/ft ³	1.8 lb/ft ³	0.3	0.09
7	143.8 lb/ft ³	141.3 lb/ft ³	2.5 lb/ft ³	-0.4	0.16
8	142.8 lb/ft ³	139.8lb/ft ³	3.0 lb/ft ³	0.9	0.81
9	144.8 lb/ft ³	143.3 lb/ft ³	1.5 lb/ft ³	-0.6	0.36
10	143.0 lb/ft ³	141.0 lb/ft ³	2.0 lb/ft ³	-0.1	0.01
	Average Difference:		+2.1 lb/ft ³		

$$\sqrt{\frac{\sum x^2}{n-1}}$$

Where:

∑ = Sum

x = Difference from the average Difference

n-1 = number of data sets minus 1

Example: 10 - 1 = 9

$$\sqrt{\frac{2.5}{9}} = 0.53$$

X1.1.1.

The Sum of X2 = 2.5 and the number of data sets = 9 for a computed standard deviation of 0.53. This is within the allowable 2.5 therefore no cores are eliminated, use the average difference from all ten cores.

Performance Exam Checklist FOP for AASHTO T 355 In-Place Density of Asphalt Mixtures by Nuclear Method

Participant Name Exam Date			
Rec	ord the symbols "P" for passing or "F" for failing on each step of the checklist.		
Pro	cedure Element	Trial 1	Trial 2
1.	Gauge turned on approximately 10 to 20 minutes before use?		
2.	Gauge calibrated and standard count recorded?		
3.	3. Test location selected appropriately [600 mm (24 in) from vertical projections or		
4.	Procedure:		
	a. Filler spread evenly over test site?		
	b. Excess filler material removed by striking off the surface?		
	c. Gauge placed on pavement surface and footprint of gauge marked?		
	d. Probe extended to backscatter position?		
	e. One-minute count taken; gauge rotated 90°, reseated, and another one-minute count taken?		
	f. Densities averaged?		
	g. If difference of the wet densities is greater than 40 kg/m3 (2.5 lb/ft3), retest conducted in both directions?		
5.	Core correlation applied if required?		
6.	Percent compaction calculated correctly?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	ature of Examiner WAQTC #:		
Com	nments:		

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 \times 10K" should be 1.56 \pm 0.05 V.; the range of "R \times 100K" should be 1.43 \pm 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 $-12 \times 12 \times \frac{1}{8}$ in. (304.8 mm \times 304.8 mm \times 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 $12 \times 12 \times 1$ in. (304.8 mm \times 304.8 mm \times 25.4 mm), medium density (see Figure 1, below).

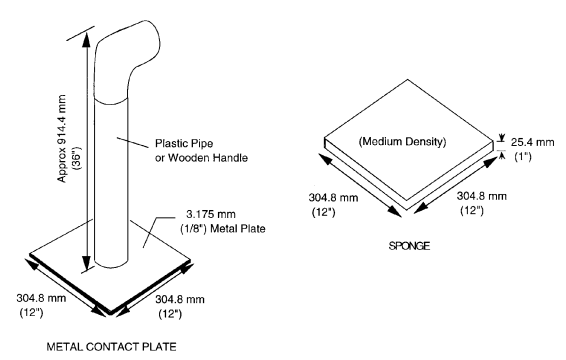


Figure 1

- 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. ³/₄-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 ½ 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

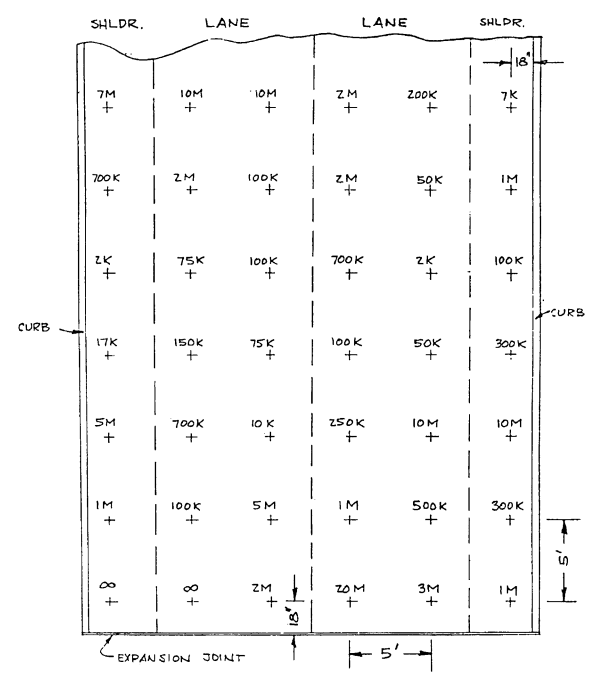
- (1) New pavement Procedure steps 3.a.(1) through 3.a.(5) may be used in measuring the resistance of the membrane-pavement system, except that in step 3.a.(5) the test locations should be prewetted with the wetting solution. Only enough wetting solution should be used to saturate the test zone. Surface 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 tin 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 its 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)."



7K= 7,000 OHMS

ZM = 7,000,000 OHMS

∞ = INFINITE RESISTANCE

Example of Test Grid and Resistance Tabulation Figure 2

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 THCKft.	
GROUND CHECK RDGohms (avg.) OHMMETER RANGE US	ED R x
GRID SPACINGft. CREW_	
BRIDGE & GRID LAYOUT AND MEASUREMENT	TS

SCALE: VERT.___in. =___ft.; HORIZ.___in. =___ft. NOTE: Use additional sheets (blank) if necessary.

Example of Electrical Resistivity Measurements Figure 3

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.
 - (3) Method of determining pH of soil.
 - (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.
- (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 10 lb (4.5 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. Laboratory Method of Determining pH OF SOIL

a. Scope

This method covers the laboratory procedure for determining pH of soil samples selected as indicated in Section 2.

b. Apparatus and Materials

- (1) pH meter suitable for laboratory testing.
- (2) Suitable containers constructed of glass or wax coated paper, with moisture proof covers.
- (3) pH buffer solutions of pH 4.0, 7.0 & 10.0 (or those recommended by the pH meter manufacturer for meter standardization.)
- (4) Distilled water and wash bottle.
- (5) Thermometer (if required) readable to $0.2^{\circ}F$ ($0.1^{\circ}C$).
- (6) U.S. No. 8 (2.36 mm) sieve.
- (7) Balance, with sufficient capacity and readable to 0.1% of the sample mass, or better, conforming to the requirements of AASHTO M 231.
- (8) Oven capable of maintaining a temperature of 140°F (60°C) around sample.
- (9) Glass stirring rod.

c. Initial Preparation of Test Samples

- (1) As received samples are to be tested for pH in a "moist" condition. If the soil as received is too wet to facilitate proper screening and reduction to test size it shall be air dried or dried to a "moist" condition in an oven at a temperature not to exceed 140°F (60°C).
- (2) Split or quarter a sufficient amount of the moist sample to yield approximately 100g of material after the material has been pulverized or mulled, taking care not to crush rock particles or naturally occurring grains, 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. Procedure for pH Determination

- (1) Place a 30.0 ± 0.1 gram sample of prepared soil into the test container.
- (2) Add 30.0 ± 0.1 grams of distilled water to the soil sample. Stir the sample to obtain a slurry and cover.
- (3) Allow the sample to stand for a minimum of 1 hour, stirring every 10 to 15 minutes.
- (4) Standardize the pH meter in accordance with the manufacturer's instructions.
- (5) Stir the sample with a glass rod immediately prior to placing the pH meter electrode into the sample. Place the electrode in the sample taking precaution to ensure good contact between the electrode and the soil slurry. DO NOT place the electrode into any soil that may have accumulated in the bottom of the container, only into the soil slurry.
- (6) Allow the electrode to remain immersed in the soil slurry for a sufficient time for the meter to stabilize. Refer to the manufacturer's instructions for recommended pH determination procedure and stabilization time.
- (7) Read and record the pH of the sample to the nearest tenth of a whole number. If the meter reads to the hundredth place it shall be rounded to the nearest tenth place.
- (8) Clean pH meter electrode and store in accordance with the manufacturer's instructions

e. Precautions

(1) Follow all manufacturer's recommendations regarding proper use of the pH meter.

f. Report

(1) Report the pH value to the nearest tenth of a whole number.

5. Laboratory Method of Determining Minimum Soil Resistivity

a. Scope

This method covers the procedure for determining the minimum resistivity of soil 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) Non-absorbent pans, bowls or other containers of sufficient size to eliminate spilling during mixing, moisture conditioning, and sample handling.
 - (5) Oven capable of maintaining a temperature of 140°F (60°C) around sample.
 - (6) Balance, with sufficient capacity and readable to 0.1% of the sample mass, or better, and conform to the requirements of AASHTO M 231.
 - (7) Distilled or deionized water.
 - (8) Spoon or spatula.
 - (9) Graduated cylinder or other suitable device of sufficient size to accurately add quantities of moisture to sample.
 - (10) Straightedge
- c. Preparation of Soil Samples
 - (1) 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 or mulled, taking care not to crush rock particles or naturally occurring grains, 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) 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) 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) 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*.

- (5) Remove the soil from the soil box and recombine it with the remainder of the original sample then add an additional 5% by *original dry soil* weight of distilled water and thoroughly mix.
- (6) Rinse the soil box with distilled water then immediately place the soil in the soil box and compact as described in step 3.
- (7) Measure the resistivity of the soil in accordance with the instructions furnished with the meter and record the value.
- (8) Repeat steps 5 through 7 until a minimum value can be determined.
- (9) 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.
- (10) Report the minimum resistivity of the soil in ohms-cm.

6. Laboratory Method of Determining Water Resistivity

- a. Measuring the Resistivity of a Water Sample
 - (l) 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.

b. Recording Data

Record data in a field record book and report the results to the Project Engineer and in the Regional Soils Report.

7. 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

Participant Name	Exam Date			
Procedure Element		Yes No		

Determining pH of H₂O.

- 1. pH meter standardized in accordance with manufacturer's instructions?
- 2. H₂O sample placed in suitable non-metallic container for testing?
- 3. pH of H₂O determined in accordance with pH meter manufacturer's instructions?
- 4. pH recorded and reported to the nearest one tenth of a whole number?

Determining pH of soil.

- 1. Sample dried (if required) to a moist condition at a temperature not to exceed 140°F (60°C)?
- 2. Sample cooled, pulverized or mulled, and screened over a U.S. #8 sieve?
- 3. Only natural material passing U.S. #8 sieve used for test?
- 4. Approximately 100 grams of passing #8 material selected for testing?
- 5. 30 ± 0.1 grams of soil and 30 ± 0.1 grams of distilled H₂O added to suitable non-metallic testing container?
- 6. Sample immediately stirred to produce slurry and covered?
- 7. Sample allowed to stand for 1 hour, stirring every 10 to 15 minutes?
- 8. pH meter standardized in accordance with manufacturer's instructions?
- 9. Soil stirred immediately prior to pH determination?
- 10. pH of soil slurry correctly determined?
- 11. pH of soil read, rounded (if necessary) and reported to the nearest one tenth of a whole number?

Determining minimum resistivity of soil.

Yes No

- 1. As received sample dried to a constant mass at a temperature not to exceed 140°F (60°C)?
- 2. Sample cooled, pulverized or mulled, and screened over a U.S. #8 sieve?
- 3. Only natural material passing U.S. #8 sieve used for test?
- 4. Approximately 4 times the volume of the soil box of material passing the U.S. #8 sieve split or quartered for testing?
- 5. 10% by weight of distilled H₂O added to sample?
- 6. Sample mixed, covered and allowed to stand for a minimum of 12 hours in a moisture proof container?
- 7. Sample re-mixed, moderately compacted in soil box and resitivity determined?
- 8. Resistivity value recorded?
- 9. Sample from soil box removed, placed with remainder of sample and additional 5% by original dry soil weight of distilled H₂O added?
- 10. Sample remixed and resistivity determined and recorded?
- 11. Steps 8 and 9 repeated until minimum resistivity can be determined?
- 12. Minimum resistivity of soil reported in ohms/cm?

Determination of H₂O resistivity.

- 1. Soil box thoroughly cleaned and rinsed at least three times with distilled H₂O?
- 2. Soil box filled level full with distilled H₂O and resistivity determined?
- 3. Resistivity from step #2 measures as infinite?
- 4. If yes, soil box emptied and resistivity of test sample determined and recorded?
- 5. If no, soil box further cleaned until condition described in step 3 satisfied?
- 6. Resistivity of H₂O sample reported in ohms/cm?

First Attempt:	Pass	Fail	Second Attempt:	Pass	Fail
----------------	------	------	-----------------	------	------

Signature of Examiner

Comments:

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 ½ 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 in not in an optimal state, then the following should be performed:
 - 7.9.1 If the sample is hot, it should be covered and allowed to cool to room temperature before testing.
 - 7.9.2 If the sample is too wet, it should be dried until it passes the squeeze test.
 - 7.9.3 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 gelpack 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 $68-77^{\circ}$ F (20-25° C) out of direct sunlight for 4 hours \pm 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

10.1 Report both the readings for the "A" paddle and the "C" paddle in the Inspector's Daily Report.

Performance Exam Checklist

Determining the Maturity of Compost (Solvita Test) WSDOT Test Method T 420

Participant Name		Exam Date				
Pro	ocedure Element		Yes No			
1.	The tester has a copy of the current procedure on hand?					
San	mple 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?					
San	mple Preprocedure					
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 \pm 10 minutes?					
9.	Maturity determined per Manufacturers instructions?					
Firs	st Attempt: Pass Fail Second	l Attempt: Pass	Fail			

Signature of Examiner

Comments:

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

9)

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:

As a minimum, the following items shall be inspected.					
1)	Mylar and four cabinet prints	13)	Phone jack for modem		
2)	Manuals	14)	Radio interference suppresser		
3)	Labeling	15)	Door light switch(s)		
4)	Air filter	16)	Pedestrian switches		
5)	All wire laced and clamped	17)	Cabinet light		
6)	Field wire terminal blocks	18)	120 VAC receptacle outlet		
7)	Door locks	19)	Ground fault interrupter		
8)	Police keys	20)	Equipment/Cabinet clearance		

- Police panel switches 21) Load switches
- 10) Circuit breakers
 11) Transient voltage suppresser
 22) Intersection display panel
 23) Isolated 120VAC bus bar (neutral)
- 12) Modem(s) 24) Phase Selectors

50) Loop Amplifiers

- 25) Flash transfer relays 38) Absence of red assembly (170) 26) Supplemental resistor loads 39) Controller 27) Two position door stop 40) Software 28) Emergency indicator lights 41) AC isolator (170) 29) Railroad preemption 42) DC isolator (170) 30) Cabinet Construction 43) Aux. file (170) 31) Detector Panel 44) External logic (NEMA) 32) Detector Panel shorting plug (NEMA) 45) CMU door interlock switch 33) Plastic document envelope (NEMA) 46) Stop Time switch 34) DB9 socket and C20 plug (170) 47) Cabinet Ground Bus bar 35) C-2 plug and cable (170) 48) Conflict Monitor 49) Inside auto/flash switch 36) Rack mounted document drawer (170)
- a. The traffic controller cabinet shall be connected to the resistor load bank.

Verify circuit breakers capacities

b. WSDOT Test Method T 425, Environmental Chamber Test shall be performed. Any deficiencies shall be documented on the test report.

6. Report

37)

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.

Test Method for Traffic Controller Inspection and Test Procedure Method T 421 Checklist

Part	articipant Name			Exam D	ate			
Prep	cedure Ele varation Cabinet in		or damage d	uring shippin	g.			Yes No
2.	Letter to p	roject off	ice sent	- 11				
3.	Traffic con	ntroller as	sessed for co	ompliance wi	th contract provisions			
4.	Simulated	load com	nected to the	Controller.				
5.	Perform E	nvironme	ntal test WS	DOT Method	d T 425			
6.	Document	test resul	ts on report.					
Firs	t Attempt:	Pass	Fail		Second Attempt:	Pass	Fail	
Sign	nature of Ex	kaminer						

Comments:

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 $300 \text{ V} \pm 5\%$ 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.

Test Method for Traffic Controller Transient Voltage Test (Spike Test) Procedure Method T 422 Checklist

Participant	Participant Name				Exam D	Date				
Procedure	Elemer	ıt							Yes	No
1. Progr	m contr	oller f	or minimu	ım recall						
2. Conn	ct line r	ioise g	enerator to	o controller j	per manufactur	es recom	mendatio	ons.		
3. With	he noise	gener	rator set to	minimum v	erify controller	is opera	ting norr	nally.		
4. Adjus	noise g	generat	or for 300	V spike.						
5. Reven	se polari	ity and	l repeat 4							
6. Redu	e spike	to min	imum and	l restore norr	nal power.					
7. Docu	nent test	t result	ts on repor	rt.						
First Atten	pt: Pas	SS	Fail		Second A	ttempt:	Pass	Fail		
Signature	f Exami	iner ₋								
Comments:										

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. binet.

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.

Participant Name

Test Method for Traffic Controller Conflict Monitor Testing WSDOT Test Method T 423

	-						
Pro	cedure Element					Yes	No
1.	Remove Conflict M	Ionitor from controller	cabinet.				
2.	Configure CMU tes	ster for CMU to be test	ed.				
3.	Connect CMU to te	ester.					
4.	Perform CMU test	as defined in appropria	te manufactures test prod	cedures.			
5.	File raw data as req	uired.					
6.	Document test resu	lts on report.					
Firs	t Attempt: Pass	Fail	Second Attempt:	Pass	Fail		
Sign	nature of Examiner						
Com	ments:						

Exam Date

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.

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.

Test Method for Traffic Controller Power Interruption Test Procedure WSDOT Test Method T 424

Par	ticipant Name		Exa	ım Date			
Pro	ocedure Element					Yes	No
1.	Program traffic contro	oller for minimum re	ecall.				
2.	Connect traffic contro recommendations.	ller to power interru	upter per the manufa	ectures			
3.	Verify traffic controlle	er operates normally	for the prescribed i	nterruptio	n.		
4.	Verify traffic controlle	er reverted to the sta	art-up sequence.				
5.	Restore normal power	r to the cabinet.					
6.	Document test results	on test report.					
Firs	st attempt: Pass 🗖 1	Fail 🗖	Second attempt:	Pass	Fail 🗖		
Sig	nature of Examiner _						
Con	nments:						

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 30°F (-34°C) to 165°F (74°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 (-30°F and 165°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 \pm 3°F. The air inside the chamber shall be circulated so that no more than a 3°F difference will occur. The chamber control shall maintain constant absolute humidity from 109°F to 165°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 30°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 & $2070 \pm \text{modified } 2010 \text{ECL}$; $102 \text{VAC } 2070 \pm \text{standard } 2010 \text{ECL}$ 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 -30°F at a rate not to exceed 30°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 30°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 -30°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 30°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

- 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.

- 1. Verify function of intersection display panel switches
- 2. Verify function of police panel switches (on/off, auto/flash)
- 3. Verify stop time switch function (inside)
- 4. Verify auto/flash switch function (inside)
- 5. Reserve for future use
- 6. Verify function of external logic (NEMA)
- 7. Verify operation of loop detection panel
- 8. Verify function of preemption push button on door (NEMA)
- 9. Verify function of preemption switches on phase selectors
- 10. Verify operation of emergency indicator light
- 11. Verify CMU functioning properly

Switches Table 1

Dry Bulb		Relative	Wet Bulb			
°F	°C	Humidity, Percent*	°F	°C		
40	4.4	75	37	2.8		
50	10.0	80	47	8.3		
60	15.6	83	57	13.9		
70	21.1	86	67	19.4		
80	26.7	87	77	25.0		
90	32.2	89	87	30.6		
100	37.8	89	97	36.1		
110	43.3	90	107	41.7		
120	48.9	70	109	42.8		
130	54.4	50	109	42.8		
140	60.0	38	109	42.8		
150	65.6	28	109	42.8		
160	71.1	21	109	42.8		
165	73.9	18	109	42.8		
*For dyr	*For dynamic testing.					

Wet-Bulb Dry-Bulb Relative Humidity at Barometric Pressure of 29.92 in hg

Table 2

Test Method for Traffic Controller NEMA and 170 Type Environmental Chamber Test WSDOT Test Method T 425

Participant Name	Exam	Date		
Procedure Element				Yes No
1. Place Traffic Controller	into the Environmental chamber.			
2. Perform Low-Temperatu	ure Low Voltage Test.			
3. Perform Low- Temperat	ture High Voltage Test.			
4. Perform High- Temperat	ture Low Voltage Test.			
5. Perform High-Temperat	ure High Voltage Test.			
6. Document test results or	n report.			
First Attempt: Pass Fa	ail Second Attempt	t: Pass	Fail	
Signature of Examiner		_		
Comments:				

Pull-Off Test for Hot Melt Traffic Button Adhesive

1. Scope

This method describes the procedure for determining the force (psi) required to pull a Type 1 raised pavement marker, from an asphalt or concrete surface that has been adhered with hot melt button adhesive.

2. Apparatus and Materials

- a. Asphalt or concrete surface, conditioned for 24 hours at standard laboratory conditions prior to testing.
- b. Raised pavement marker WSDOT Type 1 plastic or thermoplastic, drilled in the center to accept a threaded steel rod.
- c. Laboratory melter as described in ASTM D5167.
- d. Threaded steel eye bolt for attaching to the raised pavement marker.
- e. Tensile testing apparatus as described in AASHTO T 237 Section 15, 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 a 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 surface and a pre-drilled 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 pre-drilled hole in the button.
- e. The puck/block 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 the button pulls off from the surface 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 (in²)

5. Report

The pull-off strength reported shall be the average of the three determinations.

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.

Participant Name

Test Method for Loop Amplifier Testing Procedure WSDOT Test Method T 427

	-						
Pro	cedure Element					Yes	No
1.	Maximum Sensitivi	ty Check					
2.	Sustained Presence	Check					
3.	Sustained Presence	Recovery Check					
4.	Pulse Check						
5.	Second Vehicle Che	eck					
6.	Delay Check						
7.	Extension Check						
Firs	t Attempt: Pass	Fail	Second Attempt:	Pass	Fail		
Sign	nature of Examiner						
Com	iments:						

Exam Date

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 Amplilifier 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.

Test Method for Traffic Controller Inspection and Test Procedure Method T 428 Checklist

Par	Participant Name Exam Date					
Pro	ocedure Element				Yes	No
1.	Cabinet inspected for damage during shipping.					
2.	2. Letter to project office sent					
3.	3. Traffic controller assessed for compliance with contract provisions.					
4. Simulated load connected to the Controller.						
5.	Perform Transient Voltage Test WSDOT Method T	422 (optio	onal)			
6.	Perform Conflict Monitor Test WSDOT Method T	423				
7.	Perform Power Interruption Test WSDOT Method	Т 424				
8.	-					
9.	Document test results on report.					
Firs	st attempt: Pass Fail Second	attempt:	Pass 🗖	Fail 🗖		
Sig	nature of Examiner					
Con	nments:					

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

- T 421 Receiving Inspection and Test Procedure
- T 422 Transient Voltage (Spike Test) Procedure
- T 423 Conflict Monitor Testing
- T 424 Power interruption Test
- T 425 Environmental Chamber Test
- T 427 Loop Amplifier Test
- T 428 Compliance Inspection and Test Procedure

3. Process

WSDOT Test Method T 421 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 T 425 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 T 428.

WSDOT Test Method T 428 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 T 422 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 T 423 Conflict Monitor Testing

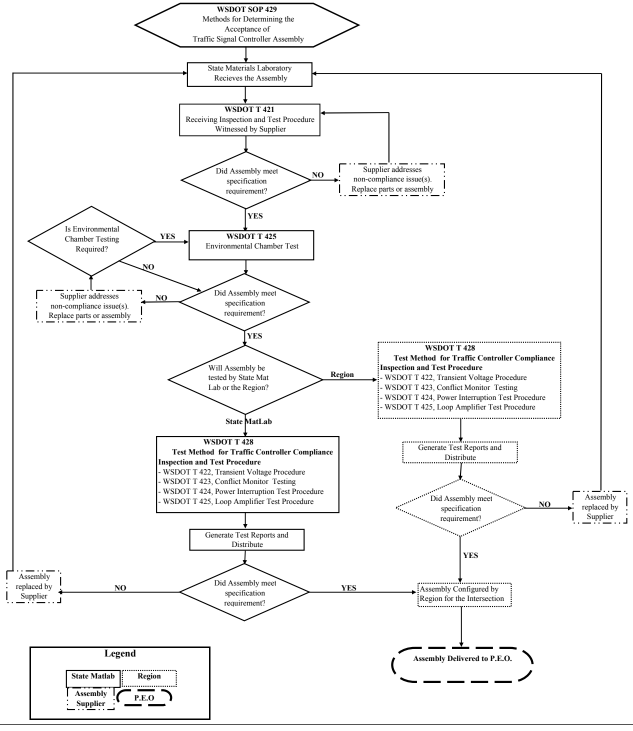
WSDOT Test Method T 424 Power interruption Test

Test Method T424 is not to be performed on type 2070 controllers

WSDOT Test Method T 427 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.



Flexibility Test for Hot-Melt Adhesives

1. Scope

This method describes the determination of flexibility of hot-melt adhesives under specific conditions.

2. Referenced Documents

- a. WSDOT SOP 318 Standard Operating Procedure for Melting of Bituminous Pavement Marker Adhesive.
- b. ASTM D3111 Standard Test Method for Flexibility of Hot-Melt Adhesives by Mandrel Bend Test Method modified to meet WSDOT specification.

3. Apparatus and Materials

- a. 1 diameter Mandrel and holder.
- b. Three-specimen stainless steel flexibility mold, 1/8 x 1 x 6 dimensions.

4. Procedure

- a. Adhesive material is melted and prepared by Liquid Asphalt lab per WSDOT SOP 318.
- b. Test specimens poured into the flexibility mold.
- c. Test specimens allowed to cure at room temperature for at least one hour.
- d. The test specimens removed from the mold and conditioned at 20°F for minimum of four hours.
- e. The 1 diameter Mandrel and its holder are also conditioned at 20°F for minimum of four hours
- f. Flexibility test is done in the same environment used to condition the specimens, by bending each specimen over the 1 Mandrel in an arc of 90° at a uniform rate for ten seconds.

5. Report

Flexibility shall be reported as Pass/Fail. Failure is a visible fracture, crazing, or cracking of the hot-melt adhesive that can occur at any time during the bending of two out of the three specimens.

Test Method to Determine Durability of Very Weak Rock

1. Scope

- 1.1 This test method covers the determination of the Jar Slake Index, Ij, of weak rock.
- 1.2 The values stated in SI units are 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
 - D 653 Terminology Relating to Soil, Rock, and Contained Fluids
 - D 2113 Practice for Rock Core Drilling and Sampling of Rock for Site Investigation
 - D 5079 Practices for Preserving and Transporting Rock Core Samples

3. Terminology

- 3.1 For terminology used in this test method, refer to Terminology D 653.
- 3.2 Definitions of terms specific to this test method:
 - 3.2.1 Jar Slake Index, Ij the visual determination of degradation of weak rock at some elapsed time after immersion in water or polymer slurry. This value is an integer ranging from 1 to 6.

4. Significance and Use

- 4.1 The Jar Slake test is a simple test developed to determine the reaction of weak rock material to water and/or polymer slurry during a certain period of time which can be tested on irregular bulk samples. Results of this test have implications on the porosity, grain interactions and density of the material.
- 4.2 This test method is used to qualitatively estimate and assign durability values to weak rocks.

5. Apparatus

- 5.1 300 ml to 600 ml clear glass laboratory jar, no taper.
- 5.2 Drying Apparatus Any suitable device capable of drying samples at a temperature not exceeding 60°C [140°F].

6. Test Sample and Specimen

- 6.1 Collect, transport, and store test samples in such manner to retain the natural water content using the guidelines in ASTM D 2113 and D 5079.
- 6.2 Test specimen shall be an air dried intact rock fragment with minimum dimensions of 25 mm × 25 mm and maximum dimensions of 65 mm × 50 mm. Specimen may be dried in an oven not exceeding 60°C [140°F].

7. Procedure

- 7.1 Place the specimen into an empty jar taking care not to break or fracture any part of it.
- 7.2 Photograph the specimen in the empty jar.
- 7.3 Add enough distilled water to cover the specimen by at least 15mm taking care not to disturb the specimen. A pre-mixed polymer slurry conforming to construction industry standards may be used in place of distilled water to investigate the retardation effects the polymer may have on the slaking process.
- 7.4 After two minutes of immersion, visually inspect the specimen to determine the Jar Slake Index, Ij, using the criteria contained in Table 1 and record the Ij for the reading.
- 7.5 Repeat Step 7.4 after 4, 6, 8, 10, 15, 20, 60, and 1440 minutes.
- 7.6 Take a final photograph of the specimen.

Jar Slake Index, Ij General behavior during test		
1	Degrades rapidly into a pile of flakes or mud	
2	Breaks readily and/or forms many chips	
3	Breaks slowly and/or forms few chips	
4	Breaks rapidly and/or develops several fractures	
5	Breaks slowly and/or develops few fractures	
6	Very little or no change	

Jar Slake Index Descriptions *Table 1*

8. Report

- 8.1 The report shall include the following:
 - 8.1.1 Specimen identification and description, test date, and test fluid used.
 - 8.1.2 Jar Slake Index value for all required readings.
 - 8.1.3 The Jar Slake Index Table (Table 1).
 - 8.1.4 Beginning and final photographs.

WSDOT Test Method T 606

Method of Test for Compaction Control of Granular Materials

1. Scope

This test method is used to establish the theoretical maximum density of granular materials and non-granular materials with more than 30 percent by weight of the original specimen is retained on the No. 4 sieve or more than 30 percent by weight of the original specimen is retained on the ³/₄ in sieve

2. Reference Documents

2.1 AASHTO Standards

- T 99 Moisture-Density Relations of Soils Using a 5.5 lb (2.5 kg) Rammer and a 12 in (305 mm) Drop (Method A only)
- M 92 Standard Specification for Wire-Cloth Sieves for Testing Purposes
- M 231 Standard Specification for Weighing Devices Used in the Testing of Materials

2.2 WSDOT Standards

- T 2 FOP for AASHTO Standard Practice for Sampling Aggregates
- R 76 FOP for AASHTO Reducing Samples of Aggregate to Testing Size
- T 255 FOP for AASHTO Total Moisture Content of Aggregate by Drying

3. Definitions

- 3.1 Fine Aggregate Portion Material passing the No. 4 Sieve.
- 3.2 Coarse Aggregate Portion Material retained on the No. 4 Sieve.

4. Significance and Use

This test method consists of three separate tests which present a method for establishing the proper theoretical maximum density values to be used for controlling the compaction of granular materials. In general, this test method is applicable to granular materials having 30 to 70 percent of the material passing the No. 4 (4.75 mm) sieve. These methods account for variations of maximum obtainable density of a given material for a given compactive effort, due to fluctuations in gradation.

5. Apparatus

- 5.1 A vibratory spring-loaded compactor. Information on where to obtain this equipment will be provided by the State Materials Laboratory.
- 5.2 Small Mold height = $8 \text{ in} \pm 0.1$ internal diameter = $6 \text{ in} \pm 0.15$, a piston to fit inside the mold with a maximum $\frac{1}{16}$ in clearance between piston and mold.
- 5.3 Large Mold-Approximately ½ ft³ (internal height 85-150 percent of diameter) with a piston to fit inside mold having a maximum ¼6 in clearance between piston and mold.
 - 5.3.1 The molds and pistons will be constructed of metal of such dimensions as to remain rigid and inflexible under test conditions.
- 5.4 Spacer blocks of varying heights compatible with the compactor and pistons.
- 5.5 Measuring device, accurate and readable to 0.01 in with a minimum 6 in length.
- 5.6 Pycnometer calibrated at the test temperature having a capacity of at least 1 quart (100 ml). Glass pycnometers shall be used to determine the specific gravity of the fine particles. The glass pycnometer shall have a companion glass plate large enough to cover the jar's opening when calibrating or weighing the pycnometer.
- 5.7 Absolute pressure gauge or vacuum gauge, used for annual standardization 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.
 - *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.
- 5.8 One vacuum pump or aspirator (pressure not to exceed 100 mm mercury).
- 5.9 One balance accurate to 0.1 g.
- 5.10 3 in (75 mm), ³/₄ in (19 mm), and a No. 4 (4.75 mm) sieve conforming to ASTM E11 requirements.
- 5.11 Balance or Scale Capacity sufficient for the principle sample mass, readable to 0.1 percent or 0.1 g, and meeting the requirements of AASHTO M 231.
- 5.12 Manually Operated Metal Rammer As specified in AASHTO T 99, Apparatus.
- 5.13 Tamping rod of straight steel, % in (16 mm) in diameter and approximately 24 in (400 mm) long having at least one end rounded to a hemispherical tip.
- 5.14 Graduated cylinder.
- 5.15 A stopwatch or timer readable to 1 second.

6. Selection of T 606 Test and Procedure

To select the proper method for determining the maximum density of the fine aggregate portion of the sample, refer to the Fine Aggregate Split of Original Sample section of Table 1.

To select the proper procedure in Test 2 for determining the maximum density of the coarse aggregate portion of the sample, refer to the Coarse Aggregate Split of Original Sample section of Table 1.

Fine Aggregate Split of Original Sample					
Soil Type	Test Method				
Sandy, non-plastic, permeable soils or non-cohesive soils.	T 606, Test 1				
Silt, some plasticity, low permeability.	T 99, Method A				
Sandy/silt, some plasticity, permeable.	T 606, Test 1/T 99, Method A (use highest results)				
Coarse Aggregate Split of Original Sample					
No more than 15 percent by weight of the original aggregate specimen exceeds 3/4 in	T 606, Test 2, Procedure 1				
15 percent or more by weight of the original aggregate specimen is greater than 3/4 in (19 mm), but does not exceed 3 in (76 mm).	T 606, Test 2, Procedure 2				

Test Selection Table 1

7. Sampling Material

- 7.1 Sample the material in accordance with WSDOT FOP for AASHTO T 2.
- 7.2 Native soils within the contract limits to be used for embankment construction and/or backfill material do not require sampling by a qualified tester.
- 7.3 For material that requires gradation testing such as but not limited to manufactured aggregates and gravel borrow, sampling shall be performed by a qualified testers.

8. Sample Preparation

- 8.1 Prepare the field sample by splitting out a representative portion in accordance with WSDOT FOP for AASHTO R 76.
- 8.2 Dry the compaction sample in accordance with WSDOT FOP for AASHTO T 255.
- 8.3 Scalp the plus 75 mm (3 in) material from the compaction sample and discard, if not required for other tests.
- 8.4 Separate the remainder of the compaction sample into coarse and fine aggregate fractions as follows:
 - 8.4.1 Fine Aggregate (No. 4 minus) Minimum of three portions approximately 13 lb (6 kg) each.

8.4.2 Coarse Aggregate

- 8.4.2.1 Procedure 1 (Aggregate Size: No. 4 to ³/₄ in (19 mm) Separate a representative specimen of 10 to 11 lbs (4.5 to 5 kg) and weigh to 0.01 lbs (5 g) or less if using a balance that is more accurate than 0.1 lbs.
- 8.4.2.2 Procedure 2 (Aggregate Size: No. 4 to 3 in (76 mm) Separate a representative specimen of 45 lbs (20 kg) and weigh to 0.1 lbs (50 g) or less if using a balance that is more accurate than 0.1 lbs.

9. Procedure

- 9.1 Test No. 1 Compaction Test of the Fine Fraction (No. 4 Minus Material)
 - 9.1.1 Assemble the small mold and determine its mass, along with the piston, to the nearest 0.01 lb (5 g). Record this as the Mass of Mold Assembly.
 - 9.1.2 Using one of the fine aggregate portions, add an amount of water estimated to produce a saturated sample (see Note 1). Mix the water and aggregate until the sample is homogenous.
 - **Note 1:** The sample is considered saturated when one to two drops of free water are visible at the base of the mold at the end of the first 2-minute cycle. Do not over saturate the material.
 - 9.1.3 Set the piston aside and place the sample in the mold in three approximately equal layers. Consolidate each lift by 25 strokes of the tamping rod followed by 25 blows of the manually operated metal rammer. The surface of the top lift should be finished as level as possible.
 - 9.1.4 Place the piston on top of the sample and mount the mold on the jack platform in the compactor. Spacers between the load spring and piston must be used to adjust the elevation of the mold so the hammers strike the mold in the center of the lift area.
 - 9.1.5 Elevate the mold until the loading head seats on top of the piston. Apply an initial seating load of approximately 100 lbs on the sample.
 - 9.1.6 Start the compactor hammers and, by elevating the jack, begin the loading procedure. The load is gradually applied over the time stated in the table below.

Load Application Rate				
Load Time				
0 to 500 lb	1 minute			
500 lb to 1,000 lb	30 sec			
1000 lb to 2,000 lb	30 sec			

- 9.1.7 Upon reaching the 2,000 lb load at the end of the 2-minute cycle, stop the hammers, release the load on the jack, return to zero pressure, and check for free water.
 - **Note 2:** If dirty water is flooding off the base of the mold or excessive material is pumping around the sides of the top piston, the sample is beyond the saturation point. Stop the test, remove the material from the mold, prepare a new sample at lower moisture content, and begin the test again.
- 9.1.8 Repeat Steps 9.1.5 through 9.1.7 four additional times (excluding check for free water). After the last run, remove the mold from the compactor.
- 9.1.9 Measure the height of the compacted sample to the nearest 0.01 in (0.1 mm) and record as the "Depth."

- 9.1.10 Determine the mass of the specimen in the mold to the nearest 0.01 lb (5 g). Record this as: Mass of Mold + Sample.
- 9.1.11 Remove the specimen from the mold and determine the moisture content in accordance with WSDOT FOP for AASHTO T 255.
- 9.1.12 Vertically slice through the center of the specimen, take a representative specimen (at least 1.1 lbs (500 g)) of the materials from one of the cut faces (using the entire specimen is acceptable), weigh immediately, dry in accordance with AASHTO T 255 to determine the moisture content, and record the results.
- 9.1.13 Calculate and record the dry density of fine fraction.
- 9.2 Test No. 2 Compaction Test of the Coarse Fraction
 - 9.2.1 Procedure $1 \frac{3}{4}$ in (19 mm) to No. 4 (4.75 mm) Aggregates
 - 9.2.1.1 Determine the mass of the coarse aggregate to the nearest 0.01 lb (5 g).
 - 9.2.1.2 Add 2.5 percent moisture to the sample, mix thoroughly.
 - 9.2.1.3 Place in 0.1 ft³ (0.0028 m³) mold in approximately three equal lifts.

 Tamp each lift lightly to consolidate material and achieve a level surface.

 Avoid the loss of any material during placement.
 - 9.2.1.4 Follow steps 9.1.5 through 9.1.8.
 - 9.2.1.5 Measure the height of the compacted sample to the nearest 0.01 in (0.1 mm) and record as the "Depth."
 - 9.2.1.6 Calculate and record the dry density of coarse fraction.
 - 9.2.2 Procedure 2-3 in (76 mm) to No. 4 Aggregates
 - 9.2.2.1 Determine the mass of the coarse aggregate to the nearest 0.01 lb (5 g) or better.
 - 9.2.2.2 Divide the sample into five representative, approximately equal portions.
 - 9.2.2.3 Place one of the portions into the ½ ft³ (0.014 m³) mold and level the surface.
 - 9.2.2.4 Position the piston on the material, mount the mold in the compactor, and compact as described in steps 9.1.5 through 9.1.7.
 - *Note 3:* Spacers may be needed between the load spring and piston to adjust the elevation of the mold to the height of the lift being compacted.
 - 9.2.2.5 Repeat 9.2.2.3 and 9.2.2.4 for the remaining four portions of material.
 - 9.2.2.6 After the final portion is compacted, determine the height of the compacted sample to the nearest 0.01 in (0.1 mm) and record as the "Depth."
 - 9.2.2.7 Calculate and record the dry density of coarse fraction (see Calculations section).

- 9.3 Test No. 3 Specific Gravity Determination for Maximum Density Test
 - 9 3 1 Material
 - 9.3.1.1 Fine fraction No. 4 (4.75 mm) minus 1.1 lbs (500 g) minimum.
 - 9.3.1.2 Coarse fraction No. 4 (4.75 mm) plus 2.2 lbs (1,000 g) minimum.
 - 9.3.2 Procedure
 - 9.3.2.1 Place dry materials, either fine or coarse fraction, in pycnometer.
 - 9.3.2.2 Fill the pycnometer approximately ³/₄ full with 68°F (20°C) water.
 - 9.3.2.3 Connect the pycnometer to the vacuum system. Apply a partial vacuum of 30 mm Hg or less absolute pressure for a period of 20 minutes.
 - 9.3.2.4 Agitate container either continuously by mechanical device or manually by vigorous shaking at 2-minute intervals.
 - 9.3.2.5 Release vacuum and disconnect the hoses.
 - 9.3.2.6 Fill pycnometer with water. Water temperature during test should be maintained as close to $68^{\circ} \pm 1^{\circ}F$ ($20^{\circ} \pm 0.5^{\circ}C$) as possible.
 - **Note 4:** It may be necessary to place the pycnometer in a water bath for 10 minutes, after release of vacuum, to bring the water temperature back to $68^{\circ} \pm 1^{\circ}$ F ($20^{\circ} \pm 0.5^{\circ}$ C).
 - 9.3.2.6.1 Metal Pycnometer (Coarse Specific Gravity Only) Fill the vessel, according to the manufacturer's instructions, with $68^{\circ} \pm 1^{\circ}F$ ($20^{\circ} \pm 0.5^{\circ}C$) water. Dry the outside of the vessel and weigh to the nearest 0.1g. Record the weight.
 - 9.3.2.6.2 Glass Pynometer (Fine or Coarse Specific Gravity) Completely fill the pycnometer with $68^{\circ} \pm 1^{\circ}F$ ($20^{\circ} \pm 0.5^{\circ}C$) water, then slide the calibrated glass plate over the mouth of the jar making sure air bubbles are not trapped under the glass plate. Dry the outside of the pycnometer and glass plate and weigh to the nearest 0.1g. Record the weight.

Calculations

10. Determine the dry density of each of the fine aggregate points as follows:

10.1 Calculate Specific Gravity as follows:

Sp. Gr. =
$$=\frac{a}{(a+b-c)}$$

Where:

= Weight of dry material, grams

= Weight of pycnometer + water, grams

= Weight of pycnometer + material + water, grams

10.2 Calculate the wet sample weight:

$$e = c - d$$

Where:

= Wet sample weight, g

= mold and wet sample weight

= Tare of mold assembly

10.3 Calculate the wet density by:

$$g = \frac{e}{b \times f}$$

Where:

g = wet density, lb/ft³

= wet sample weight, lbs

= mold constant, ft³/in

= height of sample, in (height constant-depth)

10.4 Calculate the dry density of each of the fine fraction specimens as follows:

$$h = \frac{g}{1+n}$$
Where:

h = dry density, lb/ft³

= wet density, lb/ft³

= moisture content, expressed as a decimal

11. Reports

11.1 Enter information into the WSDOT Materials Testing System (MATS) or other form approved in writing by the State Materials Engineer to obtain the theoretical maximum density curve.

Performance Exam Checklist

WSDOT Test Method T 606

Method of Test for Compaction Control of Granular Materials

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?

Fine Fraction – 100% Passing the No. 4 (4.75 mm) Sieve

Specimen Preparation

Participant Name

- 1. Has the specimen been oven-dried?
- 2. Has the specimen been separated on the No. 4 (4.75 mm) sieve?
- 3. Is the specimen weight approximately 13 lbs?

Procedure

- 1. Is specimen saturated when compacted?
- 2. Has specimen been placed in three layers, rodded 25, and tamped 25 times, each layer?
- 3. Is the hammer blow approximately a 12 in free fall to prevent severe displacement of the specimen?
- 4. The specimen is as level as possible?
- 5. Has piston been placed on top of the specimen?
- 6. Has the mold been mounted on the jack in the compactor?
- 7. Has the mold been elevated until the load-spring retainer sits on top of the piston?
- 8. Has the initial load been set at 100 lbs?
- 9. Is the loading rate applied as specified in the test procedure?
- 10. Has the hammer been stopped, jack released, and pressure returned to zero when 2,000 lbs pressure was reached?
- 11. Are one to two drops of free water visible at the base of the mold at the end of the first 2-minute cycle?
- 12. Steps 7 through 10 repeated four additional times?
- 13. The mold removed from the compactor?
- 14. Has the height of the specimen been determined?
- 15. Has specimen been weighed?
- 16. Has specimen 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. Theoretical maximum density determined by testing fresh specimens, as necessary, at different moisture contents and entered on the testing sheets?

Procedure Element Yes No

Aggregate Size: No. 4 to ¾ in (19 mm)

Specimen Preparation

- 1. Has the specimen been oven-dried?
- 2. Has the specimen been separated on the No. 4 (4.75 mm) sieve?
- 3. Does more than 85 percent of the material pass the ³/₄ in (19 mm) sieve?

Procedure

- 1. Weight and record specimen weight?
- 2. Has the specimen been dampened to $2\frac{1}{2}$ percent and placed in three lifts in a 0.1 ft³ mold?
- 3. Specimen lightly tamped to archive a level surface?
- 4. Piston placed on top of specimen 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 zero when 2,000 lb load has been reached?
- 9. Steps 5 through 8 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

Specimen Preparation

- 1. Has the specimen been oven-dried?
- 2. Has the specimen been separated on the No. 4 (4.75 mm) sieve?
- 3. Is the specimen weight approximately 45 lbs?
- 4. Does the specimen contain 15 percent or more $\frac{3}{4}$ + material?
- 5. Has material greater than 3 in (76 mm) been removed?
- 6. Specimen separated into five approximately equal parts?

Procedure

- 1. Specimen placed in the mold in five separate lifts?
- 2. The specimen is as level as possible?
- 3. After each lift, mold placed in compactor and compacted according to test procedure?
- 4. After compacting final lift, specimen removed from compactor and volume determined?
- 5. Dry density determined calculated and entered onto testing sheet?

Procedure Element Yes No

Specific Gravity Determination for Theoretical Maximum Density Test

Specimen Preparation

- 1. Has the specimen been oven-dried?
- 2. Has the specimen been separated on the No. 4 (4.75 mm) sieve?
- 3. Weight of fine fraction approximately 500 g?
- 4. Weight of coarse fraction approximately 1000 g?

Procedure

- 1. Material placed in pycnometer and 68°F water added?
- 2. Vacuum applied for at least 20 minutes?
- 3. Container and contents agitated manually by shaking at intervals of 2 minutes?
- 4. Pycnometer filled with water at 68°F?
- 5. Pycnometer dried, weighted, and recorded on testing sheet?
- 6. Specific Gravity calculated and entered onto testing sheet?

First Attempt:	Pass	Fail	Second Attempt:	Pass	Fail
Signature of Exa Comments:	miner				

WSDOT Test Method T 610

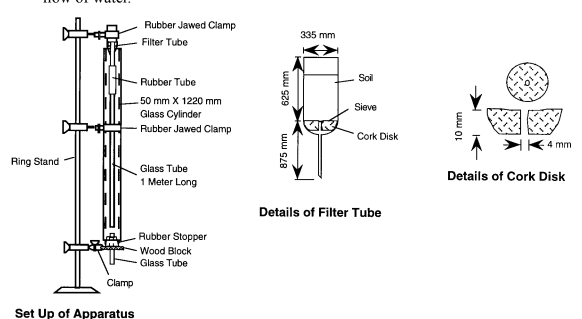
Method of Test for the Capillary Rise of Soil

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 (1,000 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.



Capillary Apparatus Figure 1

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 Standard Operating Procedure SOP 615

Determination of the % Compaction for Embankment & Untreated Surfacing Materials Using the Nuclear Moisture-Density Gauge

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. AASHTO T 99 for Method of Test for Moisture-Density Relations of Soils
- b. AASHTO T 180 for Method of Test for Moisture-Density Relations of Soils
- c. T 255 WSDOT FOP for AASHTO for Total Moisture Content of Aggregate by Drying
- d. T 272 WSDOT FOP for AASHTO for Family of Curves One Point Method
- e. T 310 WSDOT FOP for AASHTO for In-Place Densities and Moisture Content of Soils and Soil-Aggregate by Nuclear Methods (Shallow Depth)
- f. WSDOT T 606 Method of Test for Compaction Control of Granular Materials

3. 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.

When retesting is required due to a failing test; retest within a 10-foot radius of the original station and offset.

4. 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 in the Materials Testing System (MATS), WSDOT Form 350-074, Field Density Test, or other form approved in writing by the State Materials Engineer.

5. Oversize Determination

a. AASHTO T 99 and WSDOT T 606

A sample weighing a minimum of 4.08 kg (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.

There are two methods for determining the percentage of material retained on the No. 4 sieve:

- 1. Method 1 material that allows for the easy separation of fine and coarse aggregate.
 - a. Dry the sample until no visible free moisture is present (material may still appear damp but will not be shiny).
 - b. Determine and record the mass of the sample to the nearest 0.1 percent of the total mass or better.
 - c. Shake the sample by hand over a verified No. 4 (4.75 mm) sieve taking care not to overload the sieve. Overloading for a No. 4 (4.75 mm) sieve is defined as; A retained mass of more than 800 g (1.8 lbs), on a 12 inch sieve, or 340 g, (0.75 lbs); on an 8 inch sieve after sieving is complete.
 - *Note 1:* If the tester suspects a sieve will be overloaded the sample can be separated into smaller increments and recombined after sieving.
 - d. Determine and record the mass of the material retained on the No. 4 (4.75 mm) sieve to the nearest 0.1 percent of the total mass or better and record.
- 2. Method 2 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.
 - a. Determine and record the mass of the sample to the nearest 0.1 percent of the total mass or better and record.
 - b. Shake sample by hand over a verified No. 4 (4.75 mm) sieve. Do not overload the sieve. (See Section 1a and Note 1 for overload definition and information on how to prevent overloading of a sieve)
 - c. Shake material until no particles are observed passing the No. 4 (4.75 mm) sieve
 - d. Rinse the sample with water
 - e. Continue rinsing the material until it is visibly free of any coating or minus No. 4 material.
 - f. Place the washed material, retained on the No. 4 (4.75 mm) sieve, into a tared container and blot until no visible free moisture is present on the material (material may still appear damp but will not appear shiny).
 - g. Determine and record the mass of the material retained on the No. 4 (4.75 mm) sieve to the nearest 0.1 percent of the total mass or better.

b. AASHTO T 180

- 1. Follow either Method 1 or Method 2 in 5 a. with the following exception; sieve the material over a ¾ in (19.0 mm) sieve.
- 2. Do not overload the ³/₄" (19.0 mm) sieve. Overloading of a ³/₄" (19.0 mm) sieve is defined as: A retained mass of more than 3.2 kg (7.04 pounds) on a 12 inch sieve or 1.4 kg (3.08 pounds) on an 8 inch sieve after sieving is complete.

6. Calculations

a. Calculate the percent retained as follows:

% retained (Pc) =
$$100 \times \frac{\text{mass retained on sieve}}{\text{original mass}}$$
 (round to nearest percent)

b. Calculate percent passing as follows:

$$\%$$
 passing = $100 - \%$ retained

c. Calculate the dry density as follows:

$$d = \frac{100}{100 + W} (m)$$

Where:

d = dry field density of total sample, pcf

m = total field wet density, pcf

W = moisture content of total field sample

d. Calculate the corrected theoretical maximum density as follows:

$$D_{d} = \frac{100 \times (D_{f}) \times (k)}{[(D_{f}) \times (P_{c}) + (k) \times (P_{f})]}$$

Where:

D_d = corrected dry density of combined fine and oversized particles, expressed as lbs/ft³.

 $D_f = dry$ density of fine particles expressed as lbs/ft³, determined in lab.

 P_c = percent of coarse particles, by weight.

 P_f = percent of fine particles, by weight.

k = 62.4 x Bulk Specific Gravity.

Calculate in-place dry density to the nearest 0.1 lbs/ft³.

Note 2: If the specific gravity of the coarse particles has been determined, use this value in the calculation for the "k" value. If the specific gravity is unknown then use 2.67. Either AASHTO T 85 or WSDOT T 606 Test 3 may be used to determine the specific gravity of the coarse particles.

e. Calculate the percent of compaction using the following equation:

% compaction =
$$\frac{\text{Dry Density (lbs/ft}^3)}{\text{corrected theoretical maximum density (lbs/ft}^3)}$$

7. Density Curve Tables

The Materials Testing System (MATS) Density Curve Tables is the WSDOT preferred method for determining the corrected theoretical maximum density.

- a. MATS calculates the corrected theoretical maximum density in accordance with AASHTO T 99 and T 180 ANNEX A1. (Correction of Maximum Dry Density and Optimum Moisture for Oversized Particles) and reports the results in the Density Curve Table.
- b. To determine the corrected theoretical maximum density using the Density Curves Table enter the Table at the line corresponding to the % passing or % retained (T 99 & T 180 requires percent retained, T 606 requires percent passing), read across to the column labeled Max this number is the Corrected Theoretical Maximum Density.

8. Report

- a. Report the results using one or more of the following:
 - Materials Testing System (MATS)
 - WSDOT Form 350-074 and 351-015
 - Form approved in writing by the State Materials Engineer
- b. Report the percent of compaction to the nearest whole number.

Performance Exam Checklist

WSDOT Standard Operating Procedure SOP 615
Determination of the % Compaction for Embankment &
Untreated Surfacing Materials Using the Nuclear Moisture-Density Gauge

Par	ticipant Name	Exam Date		
Pro	ocedure Element		Yes	No
1.	The tester has a copy of the current procedure on hand?			
2.	All equipment is functioning according to the test proced has the current calibration/verification tags present?	ure, and if required,		
	adation Analysis a) Method 1			
1.	Sample Dried to a SSD condition (dried until no visible f mass recorded?	ree moisture present) and		
2.	Sample allowed to cool sufficiently prior to sieving?			
3.	Sample was shaken by hand through the appropriate sieve of time?	e for a sufficient period		
4.	Recorded mass of material retained on the appropriate sie	eve?		
5.	Calculated and recorded percent of material retained and appropriate sieve?	passing the		
3(B) Method 2			
1.	Mass of sample determined prior to washing?			
2.	Material charged with water in suitable container and agi	tated to suspend fines?		
3.	Sample decanted over required sieve for a sufficient amore overloading sieve?	unt of time without		
4.	Retained material dried to SSD condition and mass determined to SSD condition and mas	mined?		
5.	Recorded mass of material retained on appropriate sieve?	ı		
6.	Calculated and recorded percent of material retained and	passing appropriate sieve?		
Co	rrection for Coarse Particles			
7.	Appropriate MATS Density Curve Table used to determine maximum density, based on the percent passing or retained			
8.	All calculations performed correctly?			
Firs	st Attempt: Pass Fail Second	Attempt: Pass Fail		
Sig	nature of Examiner			

Comments:

WSDOT Test Method T 716

Method of Random Sampling for Locations of Testing and Sampling Sites

A. Scope

- 1. 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.
- 2. 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.

B. Summary of Method for Selecting Random Test Location

- Method A Determining a Random Location for Hot Mixture Asphalt (HMA) Density Tests
- Method B Determining Random Test Location for Sampling HMA Mix, Aggregates, and Miscellaneous Materials
- Method C Determining Random Test Location for Portland Cement Concrete
- Appendix A Hot Mix Asphalt Density Test Locations for Irregular Paving Areas

C. Procedure for Determining Random Test/Sampling Location

Method A – Selection of Random Location for HMA Density

1. Stationing

This method outlines the procedure for determining the random location of HMA Density testing sites using stationing.

Calculate the linear foot distance for tons specified per sublot (i.e. 80 or 100 ton sublots).

Equations:

Sublot length (ft) =
$$\frac{Sublot \ quantity \ (tons)}{\left(\frac{width \ (ft) \ x \ depth \ (ft) \ x \ 2.05 \ tons}{27}\right)}$$

a. Use a random number generator (i.e. calculator, computer) or a random number determined by a stopwatch (See Note 1) to enter Table 1. Use the corresponding X value to determine the test station. A new X value is required for every test.

Note 1: To use the stopwatch method, randomly start and stop the stopwatch 10 or more times, then use the decimal part of the seconds as your entry point.

b. Determine the test station as follows:

Test Station = (sublot length \times "X" multiplier) + beginning station of paving (round to the nearest foot)

- c. Use a random number generator (i.e. calculator, computer) or a random number determined by a stopwatch (See Note 1) to enter Table 2. Use the corresponding "Y" multiplier to determine the offset. A new "Y" multiplier is required for every test.
- d. Determine the offset as follows:

Offset = (width of pavement \times "Y" multiplier) (round to the nearest 0.1 ft)

Offset may be figured from the right or left edge of pavement. Tester shall indicate in MATS or approved density form from which edge the offset is measured.

e. If a tester must move a testing location due to an obstruction of other interference, a new random number for the offset and station shall be picked and the location recalculate. Document the new location and the reason the testing location was changed.

Example for a 100 ton sublot:

Given:

Paving width = 12 ft

Paving depth = 0.15 ft

Beginning Station = 10 + 00

Offset from left edge of pavement

Calculations:

Sublot length =
$$\frac{100}{\left(\frac{12 \times 0.15 \times 2.05}{27}\right)}$$
 = 731.7 lf

Ending Station = (Beginning Station + Sublot length) = (1000 + 731.7) = 17 + 32

Random generated number = X=25, Y=10

Beginning Test Location

Enter Table 1 at (25): "X" multiplier = 0.080

Enter Table 2 at (10): "Y" multiplier 0.167

Testing Station = $(732 \times 0.080) + 1000 = 1058.5 = 10 + 59$ (round to the nearest ft)

Offset = $(12 \times 0.167) = 2.00 = 2.0$ ft left of pavement edge (round to the nearest 0.1 ft)

2. Milepost

This method outlines the procedure for determining the random location of HMA Density testing sites using mileposts.

a. Convert to tons per mile using the roadway area based on the roadway width and depth. Equations:

$$Sublot\ length\ (mile) = \frac{Sublot\ quantity\ (tons)}{\left(\frac{width\ (ft)x\ depth\ (ft)x\ 2.05\ tons}{27}\right)x\ 5280\ lf}$$

Round sublot length to the nearest thousandth (0.001) of a mile

Calculate the location of the test site and offset using the same method as described in Method A Stationing except use tons per mile instead of the tons per *lf*.

Test site = (sublot length
$$\times$$
 "X" multiplier) + beginning milepost
Offset = (width \times "Y" multiplier)

Example for 100-ton sublot:

Given:

Paving width = 12 ft

Paving depth = 0.15 ft

Beginning Milepost (MP) = 1.00

Offset determined from right side of pavement

Calculations:

Sublot length =
$$\frac{100}{\left(\frac{12 \times 0.15 \times 2.05}{27}\right) \times 5280} = 0.138$$

Ending MP = (Beginning MP + Sublot length) = (1.00 + 0.138) = 1.138

Random generated number = X=25, Y=90

Beginning Test Location

Enter Table 1 at (25): "X" multiplier = 0.080

Enter Table 2 at (90): "Y" multiplier = 0.060

Testing MP = $(.138 \times 0.080) + 1.00 = 1.011$

Offset = $(12 \times 0.060) = 0.72 = 0.72$ ft right of edge of pavement

Random #	Х						
1	0.794	26	0.526	51	0.304	76	0.617
2	0.500	27	0.519	52	0.167	77	0.584
3	0.393	28	0.446	53	0.308	78	0.591
4	0.427	29	0.219	54	0.570	79	0.563
5	0.165	30	0.780	55	0.322	80	0.482
6	0.821	31	0.574	56	0.491	81	0.499
7	0.562	32	0.730	57	0.349	82	0.227
8	0.284	33	0.435	58	0.681	83	0.476
9	0.704	34	0.338	59	0.858	84	0.258
10	0.988	35	0.515	60	0.716	85	0.227
11	0.692	36	0.751	61	0.521	86	0.364
12	0.491	37	0.063	62	0.568	87	0.186
13	0.769	38	0.269	63	0.168	88	0.791
14	0.675	39	0.357	64	0.460	89	0.985
15	0.205	40	0.555	65	0.708	90	0.562
16	0.187	41	0.837	66	0.453	91	0.753
17	0.238	42	0.699	67	0.778	92	0.097
18	0.400	43	0.456	68	0.484	93	0.723
19	0.263	44	0.730	69	0.609	94	0.214
20	0.545	45	0.314	70	0.949	95	0.215
21	0.230	46	0.179	71	0.575	96	0.428
22	0.700	47	0.152	72	0.263	97	0.647
23	0.616	48	0.334	73	0.192	98	0.794
24	0.179	49	0.284	74	0.845	99	0.154
25	0.080	50	0.819	75	0.095	100	0.964

Random Number - X *Table 1*

Random #	Υ	Random #	Y	Random #	Y	Random #	Υ
1	0.823	26	0.755	51	0.068	76	0.298
2	0.646	27	0.922	52	0.709	77	0.217
3	0.928	28	0.299	53	0.742	78	0.662
4	0.247	29	0.855	54	0.704	79	0.709
5	0.742	30	0.270	55	0.230	80	0.634
6	0.666	31	0.875	56	0.584	81	0.245
7	0.624	32	0.076	57	0.663	82	0.672
8	0.553	33	0.393	58	0.727	83	0.620
9	0.311	34	0.366	59	0.559	84	0.580
10	0.167	35	0.860	60	0.907	85	0.452
11	0.198	36	0.605	61	0.311	86	0.141
12	0.814	37	0.239	62	0.665	87	0.937
13	0.876	38	0.349	63	0.134	88	0.228
14	0.356	39	0.201	64	0.241	89	0.225
15	0.898	40	0.650	65	0.384	90	0.060
16	0.141	41	0.822	66	0.268	91	0.820
17	0.913	42	0.157	67	0.629	92	0.883
18	0.384	43	0.799	68	0.227	93	0.528
19	0.815	44	0.340	69	0.187	94	0.749
20	0.761	45	0.479	70	0.167	95	0.441
21	0.370	46	0.925	71	0.127	96	0.221
22	0.156	47	0.494	72	0.288	97	0.863
23	0.397	48	0.833	73	0.436	98	0.082
24	0.416	49	0.128	74	0.913	99	0.467
25	0.705	50	0.294	75	0.665	100	0.828

Random Number - Y *Table 2*

Method B – Hot Mix Asphalt (HMA) Pavement Mixture or Aggregates

- 1. Determine the sublot increment of the material.
- 2. Use a random number generator (i.e. calculator, computer, etc) or a random number determined by a stopwatch (See Note 1) to enter Table 1. Use the corresponding X multiplier to determine the offset.
- 3. A new X multiplier is required for every sublot.
- 4. Random sample tonnage may be adjusted per sublot to accommodate field testing. Adjustments to random sample tonnage must be documented.
- 5. Calculate the location of the sampling site as follows:

Equations:

```
First Sample Site = Sublot increment × "X" multiplier (Table 1)
Subsequent Sites= (sublot increment + (Sublot increment × "X" multiplier)
```

Aggregate Sample Example:

Given: Crushed Surfacing Base Coarse

Random sample frequency per 9-3.7 =1 per 2,000 tons.

Calculate the location of the first random sample site as follows:

The computer-generated number is 22.

Sublot Increment (Frequency of sampling) = 2,000 tons

Enter Table 1 at (22) "X" = 0.700

Sampling Site = $2000 \times 0.700 = 1400 \text{ tons}$

Calculate subsequent sample sites as follows:

The computer-generated number is (53).

Sublot Increment (Frequency of sampling) = 2,000 tons

Enter Table 1at 53 "X" = 0.308

Sampling Site = $2000 + (2000 \times 0.308) = 2616$ tons

Method C Portland Cement Concrete (PCC)

- 1. Determine subsequent random sampling locations as follows:
 - a. Example for less than 100 cubic yards remaining after reducing frequency:
 - (1) Determine amount of pour remaining this will be the sublot increment
 - (2) Use a random number generator (i.e. calculator, computer) or a random number determined by a stopwatch (See Note 1) to enter Table 1. Use the corresponding X multiplier to determine the test station. A new X multiplier is required for every test.
 - (3) Determine the sample location as follows:

```
Sampling Location = Concrete remaining \times "X" multiplier (Table 1)
```

Given:

```
Total cubic yards (cy) of concrete placement = 80 cy
Truckload = 10 cy
Given: First truck is in specification = 10 cy
Remaining cubic yards = 80 cy - 10 cy = 70 cy < 100 cy
Sublot increment = 70 cy
Random number = 30 "X" = 0.780
Sampling Location = 70 cy × 0.780 = 54.6 = 55 cy or 7th truck
```

- b. Example for greater than 100 cubic yards remaining after reducing frequency
 - (1) *Given*:

```
Pour = 130 cy
Each truck carries 8 cy of concrete
First truck is in specification = 8 cy
Remaining cubic yards = 130 - 8 = 122 > 100 cy
Sublot Increment = 100 cy
```

- (2) Use a random number generator (i.e. calculator, computer) or a random number determined by a stopwatch (See Note 1) to enter Table 1. Use the corresponding X value to determine the test station. A new X value is required for every test.
- (3) Determine the sample location as follows:

```
Sampling Location = Sublot increment \times "X" multiplier (Table 1)
```

Example:

```
Random number = 15 "X"= 0.205
Sample location = 100 cy \times 0.205 = 20.5
Determine where the first sample will be taken:
Testing location = (accumulated cy of last truck sampled) + sample yardage
```

Example:

```
First Sample Location:
Accumulated cy first truck = 8
Sample location = 8 cy + 20.5 cy = 28.5 cy
Truck load = 28.5/8 = 4
Sampling = second half of 4th truck
```

Determine subsequent sampling locations as follows:

```
Sublot increment = total pour – (initial loads in specification)-(first sublot increment) Sublot increment = 130 \text{ cy} - (8 \text{ cy}) - (100 \text{ cy}) = 22 \text{ cy} Random number = 52 \text{ "X"} = 0.167 Testing location = (initial load in specification) + (first sublot increment) + (testing location within the second sublot) Testing location = (8 \text{ cy})+(100 \text{ cy})+(0.167 \times 22 \text{ cy}) Testing location = 111.7 \text{ cy} or 111.7/8 \text{ cy} per truck = 14.0 = 14 \text{th} truck
```

3. Report

- a. Report the random number used to determine station and offset
- b. Document any changes in station or offset of random testing location
- c. Use one of the following to report random location information:
 - Materials Testing System (MATS)
 - Form approved in writing by the State Materials Engineer

Appendix A

Hot Mix Asphalt Density Test Locations for Irregular Paving Areas

- A. Track tonnage placed in the irregular shaped area until specified tons are placed, note the stationing.
- B. Measure back to the beginning of the paving or end of the previous lot to obtain the length (this is also your beginning station).
- C. Use a computer-generated random number or a random number determined by a stopwatch (See Note 1) to enter Table 1. Use the corresponding X value to determine the test station. A new X value is required for every test.
- D. Multiply the length by the "X" value and add to the beginning station to locate your testing site.
- E. Use a computer-generated random number or a random number determined by a stopwatch (See Note 1) to enter Table 2. Use the corresponding Y value to determine the offset. A new Y value is required for every test.
- F. Measure the width at the testing station and multiply the width time the "Y" value to determine the offset of the testing site.
- G. Make a sketch of the area to document the test location in the event a retest is required.

Example:

Paving began at Station 101 + 00.

The tester determined Station 105 + 75 was the end of the 100 ton lot.

The width of the pavement began at 0 and transitioned to 12.

Testing Station

```
Sta 105 + 75 – Sta 101 + 00 = 475 ft
Random number = 45, "X" value = 0.314
475 ft × 0.314= 149.15= 149 Testing station = 10100 + 149 = 102 + 49
```

Testing Offset

```
Measure width at station 102 + 49
Width = 3.76
Random # 65 "Y" value = 0.384
Offset = 3.76 \times 0.384 = 1.44 = 1.4 ft 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 26 mm Hg and large enough to accommodate test specimens 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 specimens.
- e. Air-bath freezer, maintained at 0 ± 5 °F.
- f. Water bath maintained at 55 ± 1 °F.
- g. Testing machine A compression testing machine having a minimum capacity of 10,000 lbf and capable of producing a uniform vertical movement of 0.065 inches per minute.
- h. Equipment for preparing and compacting specimens for WSDOT FOP for AASHTO T 312.
- i. 100 ± 0.10 mm gyratory specimen mold and 99.50 to 99.75mm top/bottom plates which meet WSDOT FOP for AASHTO T 312 section 4.2 (excluding inside diameter measurements) and section 4.3 (excluding diameter measurement).

3. Preparation of Laboratory-Mixed, Laboratory-Compacted Specimens for Mix Designs

- a. Mix specimens 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 specimens per asphalt binder supplier, two samples with 0% anti-strip additive and the other specimens with varying amounts of anti-strip additive (Note 1).
 - *Note 1:* Liquid anti-strip agents added directly to the asphalt binder shall be added by weight of asphalt at levels of ½%, ½%, ¾% and 1% or levels not exceeding 1% which test an even progression of anti-strip additive per manufacture recommendation. Latex anti-strip agents shall be added to the aggregate in a Saturated Surface Dry (SSD) condition at levels of 0.08%, 0.17%, 0.33% and 0.50% by weight of dry aggregate.
- c. Condition and compact the 100 mm specimens per WSDOT FOP for AASHTO T 312 sections 8.5 through 9.8.

Preconditioning of Test Specimens

- a. Once the set of six specimens have been compacted and cooled to room temperature, set one of the specimens mixed with 0% anti-strip aside to be stored at room temperature, this will be the referee specimen.
- b. Test remaining set of specimens per AASHTO T 166 Method A. Calculate the air void level of the specimen using mix design Theoretical Maximum Specific Gravity value.
- c. Place the specimens in the vacuum container. The container must be filled with potable water at room temperature $(77 \pm 9^{\circ}F)$ so that the specimens have at least 1 inch of water above their surface. Apply a vacuum for a short amount of time, suitable to saturate the specimens air voids between 60 and 80 percent.
- d. Determine the mass of the saturated, surface-dry specimen after partial vacuum saturation per AASHTO T 166 Method A.
- 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.

$$S = \frac{100J}{Va}$$

Where:

S = Degree of saturation, percent.

Va = Volume of air voids

Determine the Volume of air voids using the following equation:

$$Va = \frac{Pa \times E}{100}$$

Where:

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 specimen has been damaged and must be discarded.
- h. After saturation is achieved place each specimen in a plastic bag, seal the bag and place specimen in a freezer at a temperature of $0 \pm 5^{\circ}F$ for a minimum of 16 hours.

- i. Remove specimens from the freezer, remove plastic bags and place them in a water bath maintained at $140 \pm 2^{\circ}$ F for 24 ± 1 hour (Note 2).
 - **Note 2:** Some specimens 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 coldwater bath.
- j. After 24 ± 1 hours in the $140 \pm 2^{\circ}F$ water bath, remove the specimens and place them into the cold water bath maintained at $55 \pm 1^{\circ}F$. At this time the referee specimen shall be placed into the cold water bath with the conditioned specimens. Testing must begin within $2 \text{ hours} \pm 10 \text{ minutes}$ after specimens have been placed into the cold water bath.

4. Testing

- a. After 2 hours \pm 10 minutes in the cold water bath, remove and test one specimen 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 specimen.
- b. Continue to load specimen until specimen can be easily broken open.
- c. Remove specimen from machine, break specimen in half by hand for visual inspection. Record the visual condition of each specimen as to stripping action: none, slight, moderate, or severe.
- d. Determine the Tensile Strength Ratio (TSR) of each specimen by comparing the load needed to break the testing specimen to the load needed to break the referee specimen, using the following equation:

$$TSR = \left(\frac{S_1}{S_2}\right) \times 100$$

Where:

 S_1 = tensile strength of the conditioned specimen

 S_2 = tensile strength of the unconditioned specimen

5. Visiual Condition Definitions

- None The specimen condition is solid with no evidence of asphalt binder withdrawing from aggregate. After the specimen has air-dried, the appearance is black.
- Slight The specimen condition is solid to slightly soft with evidence of the asphalt binder beginning to withdraw from edges and surfaces of the aggregates. After the specimen has air-dried, the appearance remains black.
- Moderate The specimen condition is soft, easily broken in half, with partial to completely exposed aggregates. After the specimen has air-dried, the appearance is slightly gray.
- Severe The specimen condition is soft to falling apart with the majority of coarse aggregate completely exposed and asphalt binder almost nonexistent. After the specimen has air-dried, the appearance is gray.

6. Report

The report shall include the following: Visually estimated moisture damage (stripping) and Tensile Strength Ratio (TSR) of the specimens.

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 in the Materials Testing System (MATS)

Standard Operating Procedure for Submitting Hot Mix Asphalt (HMA) Mix Designs for Verification

1. Scope

- 1.1 This standard covers the procedural steps required for submitting a HMA mix design for verification to the Bituminous Materials Section of the State Materials Laboratory.
- 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. Approval of Material

- 2.1 Approvals of the material for HMA are required prior to use per *Standard Specifications* Section 1-06.1.
- 2.2 A HMA mix design is required for each contract.

3. Referenced Documents

- 3.1 WSDOT Standards
 - T 2 Standard Practice for Sampling Aggregates
 - T 724 Method of Preparation of Aggregate for Hot Mix Asphalt (HMA) Mix Designs *Standard Specifications* M 41-10

4. Procedure

- 4.1 The Contractor shall determine a design aggregate structure and asphalt binder content in accordance with WSDOT Standard Operating Procedure 732.
- 4.2 Once the design aggregate structure and asphalt binder content have been determined, the Contractor shall submit the HMA mix design on WSDOT form 350-042 demonstrating that the design meets the requirements of *Standard Specifications* Section 9-03.8(2) and 9-03.8(6). For mix designs that contain > 20% RAP and any amount of RAS, the contractor shall include test results for asphalt content and gradation per GSP 5-04.2OPT8.GR5, along with a statement certifying the tonnage of the RAP and/or RAS stockpile(s) to be used in the HMA production.
- 4.3 For mix designs that contain ≤ 20% RAP and no amount of RAS, the Contractor shall obtain representative samples of aggregate per WSDOT FOP for AASHTO T 2 that will be used in the HMA production.

- 4.4 For mix designs that contain > 20% RAP and any amount of RAS, the contractor shall obtain representative samples of aggregate, RAP and/or RAS per WSDOT FOP for AASHTO T 2 that will be used in the HMA production. Additionally, the contractor will submit 100 grams each of recovered asphalt residue from the RAP and/or RAS that are to be used in the HMA production.
- 4.5 The Contractor shall submit representative samples of aggregate, RAP and RAS (if required), totaling 700 pounds proportioned to match the Contractor's proposal to the State Material's Laboratory for testing.

For example, if the Contractor's proposal consists of five stockpiles with the following blending ratio:

Material	Ratio		
³⁄₄" – #4	20%		
1/2" – #8	30%		
#4 – 0	30%		
RAP	15%		
RAS	5%		

Calculate the amount of aggregate needed from each stockpile in the following manner.

Material		Pounds of Aggregate Needed Per Stockpile
³ ⁄ ₄ " – #4	700 lbs x 0.20	140 pounds
1/2" – #8	700 lbs x 0.30	210 pounds
#4 – 0	700 lbs x 0.30	210 pounds
RAP	700 lbs x 0.15	105 pounds
RAS	700 lbs x 0.05	35 pounds

5. Shipping Samples

- 5.1 Transport aggregate 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 or container of aggregate is 30 pounds maximum.
- 5.2 Each aggregate bag or container shall be clearly marked or labeled with suitable identification including the contract number, aggregate source identification and size of stockpile material. Aggregate bags or containers submitted to the State Materials Laboratory shall be accompanied by a completed transmittal for each stockpile used in the HMA mix design and a completed copy of DOT Form 350-042.

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 $325 \pm 25^{\circ}$ F ($163 \pm 14^{\circ}$ 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 M 231.
- 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 350° 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

Partici	ipant Name		Exam I	Date			
Proce	dure Element					Yes	No
1. T	he tester has a copy	of the current p	rocedure on hand?				
	all equipment is fundarrent calibration/ve	•	ng to the test procedure, and i resent?	if required	, has the		
3. R	epresentative samp	le(s) of the prod	uction aggregates obtained.				
4. A	ggregate dried in a	n oven to a cons	tant mass?				
5. A	ggregate sieved ov	er designated sie	eves for class of mix being tes	sted?			
6. M	laterial retained on	each sieve place	ed in separate containers?				
	eparated aggregates eve, in accordance	•	the portion passing the No. 2 ASHTO T27/T11?	200 (0.075	mm)		
8. W	Vashed aggregate sa	imples dried in a	n oven to a constant mass?				
9. A	ggregate recombine	ed to match the	grading of the job mix formul	a?			
10. Sa	ample size determin	ned by the specif	fic test procedure to be perform	med?			
First A	Attempt: Pass	Fail	Second Attempt:	Pass	Fail		
Signat	ture of Examiner						
Comme	ents:						

WSDOT Test Method T 726

Mixing Procedure for Hot Mix Asphalt (HMA)

1. Scope

This is the mixing procedure for laboratory prepared samples of asphalt concrete, asphalt treated base, or open graded asphalt products mixtures. 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 sphalt.
- 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.
- h. Thermometer- Armored glass or dial-type thermometric devices with metal stems or probe for determining the temperature of aggregates, binder, and HMA between 18θ and 418θ ° F (10θ and 232θ ° C).

3. Procedure

- a. Heat asphalt binder, aggregate sample(s), and mixing bowl(s) in a preheated oven to the mixing temperature specified by the supplier of asphalt binder or as indicated on mix design report.
- b. 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 verification report.
- c. After the materials are heated place mixing bowl on balance and tare.
- d. Place heated aggregate sample in the tared mixing bowl and determine the mass of the aggregate sample. Use this mass to calculate the mass of asphalt binder required to produce a sample of HMA at the Job Mix Formula (JMF) asphalt binder content (See calculation below).
- e. Form a crater in the aggregate sample and weigh in asphalt binder as determined above.
 - *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.

f. Mix aggregate sample and asphalt binder for approximately 3 minutes or until aggregate sample is completely coated with asphalt binder. This can be accomplished by hand mixing or by mechanical mixer.

Note: Reheating of the HMA for a short period of time may be necessary to assure complete coating of the aggregate.

- g. Transfer mixed HMA to the proper container for other testing as required.
- h. Repeat steps A thru H for each sample to be mixed.

Calculation for Mass of Asphalt Binder:

Designated Mass of Asphalt Binder =
$$\frac{\text{(A) D}}{\text{(1 - A)}}$$

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 =
$$\frac{(0.053) \ 1567.1}{(1 - 0.053)} = \frac{83.1}{0.947} = 87.7g$$

Performance Exam Checklist

Mixing Procedure for Hot Mix Asphalt (HMA) WSDOT Test Method T 726

Part	ticipant Name Exam Date		
Pro	cedure 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.	Aggregate samples prepared as per WSDOT Test Method T 724?		
4.	Mixing bowl(s), aggregate and asphalt binder heated to appropriate mixing temperature?		
5.	Asphalt binder stirred and temperature confirmed by thermometer?		
6.	Heated mixing bowl placed on scale and scale then tared?		
7.	Heated aggregate sample placed in bowl and scale then tared?		
8.	Crater formed into center aggregate, weigh in asphalt binder in accordance with mix design information?		
9.	Mix aggregate and asphalt for approximately 3 minutes or until aggregate is completely coated?		
10.	When mixing is complete carefully scrape off mixing apparatus, tools and bowl is dumped into correctly marked pan?		
11.	Repeat steps 4 - 8 for each sample to be mixed?		
12.	All calculations performed correctly?		
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	nature of Examiner		

Comments:



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. Dry sample to a constant mass, not to exceed 325 \pm 25°F (163 \pm 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.



Determination of the Moving Average of Theoretical Maximum Density (TMD) for HMA

1. Scope

This procedure covers the process for obtaining the moving average of the Theoretical Maximum Density (TMD) for calculating pavement compaction in accordance with WSDOT FOP for WAQTC T 355. The TMD is to be determined in accordance with WSDOT FOP for AASHTO T 209.

2. Procedure

The procedure for determining the moving average of TMD is as follows:

- a. On the initial day of production of a new Job Mix Formula, two determinations shall be made to establish an initial average value. The samples shall not be from the same truck. Average the two TMDs and report the result to the Moisture Density Gauge Operator. The TMD value from the Mix Design Verification Report shall not be included in the average. If the two TMDs determined on the initial day do not agree within 1.5 lb/ft³ (24 kg/m³), a third determination shall be made. The initial average density shall be based on the two closest results.
- b. A TMD test shall be taken with each mix sample. The moving average is defined as the average of the last five TMD values for the HMA being placed. Until five TMD values have been determined, the moving average will consist of all previous TMD values plus the first TMD value for the current production shift. When five TMD values have been determined, the moving average for each shift will include the last four TMD values plus the first TMD value for the current paving shift. This new moving average value will be used for the entire paving shift.
- c. Each TMD shall be compared with the previously computed moving average. If a TMD deviates from the moving average by more than 1.5 lb/ft³ (± 24 kg/m³), a second test shall be made on another portion of the same sample. If the second TMD agrees within 1.5 lb/ft³ (± 24 kg/m³) of the moving average then the first TMD will be discarded and the second TMD will be included in the moving average. If the second TMD is not within 1.5 lb/ft³ (± 24 kg/m³) of the moving average but is within 1.5 lb/ft³ (± 24 kg/m³) of the first TMD, 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.
- d. A moving average will be sent to the Moisture Density Gauge operator once per production shift, unless two tests during a shift are not within 1.5 lb/ft³ (± 24 kg/m³), then a new moving average will be calculated in accordance with "c" of this procedure and sent to the Moisture Density Gauge operator as the new moving average for the shift. The Moisture Density Gauge Operator will continue to use the previous moving average until a new moving average is available.

3. Report

The gauge operator will record the average TMD received from the tester at the HMA plant on WSDOT Form 350-092 and 350-157 or in the MATS database. The average TMD will be used in WSDOT FOP for WAQTC T 355 to calculate the percent of compaction for statistical evaluation.



Correlation of Nuclear Gauge Densities With Hot Mix Asphalt (HMA) Cores

- 1. When evaluating HMA compaction:
 - 1.1 A gauge correlation is required:
 - a. For each combination of gauge and HMA Mix Design (initial JMF).
 - b. When gauge mode changes (i.e., back scatter to thin layer).
 - c. When wearing course lift thickness changes per Note 1.
 - d. When a gauge is recalibrated.
 - **Note 1:** For density determined with "Thin Layer Mode," a layer thickness change of greater than 0.08 feet requires a new correlation. For density determined with "Back Scatter Mode," a layer thickness change of greater than 0.15 feet requires a new gauge correlation.
 - 1.2 A gauge correlation is not required but may be considered by the Regional Materials Engineer when:
 - a. Base material changes from the original correlation base (i.e., from a surfacing base to an asphalt base).
 - b. The same gauge HMA Mix Design (Reference Mix Design) combination are used on a different contract within the same construction year.
 - c. When JMF has been adjusted in accordance with *Standard Specifications* Section 9-03.8(7)A.
- 2. Gauge correlation is based on ten in-place HMA densities and ten cores taken at the same location as the in-place density.
 - 2.1 In-place HMA densities shall be determined in accordance with WSDOT FOP for WAQTC T 355.
 - 2.2 Cores should be taken no later than the day following paving and before traffic has been allowed on roadway. Correlation cores are not required to be taken at record density locations. Therefore, a site outside the traveled way should be considered for worker safety, as long as the lift thickness matches that of the plan lift thickness of the record density locations.
 - *Note 2:* If a core becomes damaged, it shall be eliminated from the average.
 - **Note 3:** Cores may be taken sooner than the day after paving if the HMA is cooled to prevent damage during coring and removal of cores. Water, ice, or dry-ice may be used to cool the pavement. Another method of cooling that may be used is substitution of nitrogen gas or CO_2 for drilling fluids.

- 3. Obtain a pavement core from each of the test sites in accordance with WSDOT SOP 734. The core shall be taken in the nuclear gauge footprint.
 - 3.1 For either gauge mode, back scatter or thin layer, locate the core in the approximate center of the nuclear gauge footprint. If the core thickness exceeds the plan pavement thickness by more than 0.04 feet, then the core shall be saw cut to the plan thickness prior to performing density testing. If a core thickness is less than the plan thickness by more than 0.04 feet, it shall be eliminated from the average.
- 4. Bulk Specific Gravity (G_{mb}) of core shall be determined in conformance with WSDOT FOP for AASHTO T 166 Bulk Specific Gravity of Compacted Hot Mix Asphalt (HMA) Using Saturated Surface-Dry Specimens.

Calculate core density as follows, round to the nearest 0.1 pcf:

Core Density =
$$G_{mb} \times 62.245$$
 pcf

Calculate gauge correlation factor as follows:

Density Ratio =
$$\frac{(core\ density)}{(nuclear\ gauge\ density)}$$

Round Density Ratio to the nearest 0.001

Gauge correlation factor =
$$\frac{(Sum \ of \ ratios)}{(number \ of \ cores)}$$

- 5. Gauge Correlation Factor shall be determined to 0.001.
- 6. Report the Gauge Correlation Factor using MATS or DOT Form 350-112.

Method for Determining Volumetric Properties of Hot Mix Asphalt

1. Scope

This procedure covers the determination of volumetric properties of Hot Mix Asphalt, 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

- T 329 WSDOT FOP for AASHTO Moisture Content of Hot Mix Asphalt (HMA) by Oven Method
- T 27/11 WSDOT FOP for WAQTC/AASHTO Sieve Analysis of Fine and Coarse Aggregates
- T 166 WSDOT FOP for AASHTO Bulk Specific Gravity of Compacted Hot Mix Asphalt Using Saturated Surface-Dry Specimens
- T 168 WSDOT FOP for WAQTC/AASHTO Sampling of Hot Mix Asphalt Paving Mixtures
- T 209 WSDOT FOP for AASHTO Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt Paving Mixtures
- T 308 WSDOT FOP for AASHTO Determining the Asphalt Binder Content of Hot Mix Asphalt (HMA) by the Ignition Method
- T 312 WSDOT FOP for AASHTO Preparing Hot Mix Asphalt (HMA) Specimens by Means of the Superpave Gyratory Compactor
- T 712 WSDOT Test Method 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 T 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 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 25° F above the compaction temperature (Note 1) as soon as possible to reduce sample cooling. The gyratory test is temperature sensitive. The sample should be heated five degrees above the compaction temperature as shown on the mix design verification report.
 - **Note 1:** 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 350°F for a minimum of 60 minutes prior to the estimated beginning of compaction. Subsequent uses of a conditioned mold will require 5 minutes of reheating.
- 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 in accordance with step 4 (c).
 - **Note 2:** 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 T 312 Section 9
- d. Determine theoretical maximum 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 T 27/11.
- 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.

Note 3: For repeatability between operators the retest sample should be cooled for the same amount of time at room temperature as the original specimen. When sending retest samples to the Region or State Laboratory, note the time the original sample was cooled at room temperature in the remarks section of the transmittal.

6. Volumetric Calculations

Calculations

a. Calculate $\%G_{mm}$ @ N_{design} as follows:

Where:

 $\%G_{mm}@N_{design}=\%$ theoretical maximum specific gravity @ N_{design} = Bulk specific gravity of the compacted specimen G_{mm} = Maximum specific gravity of the paving mixture

 N_{design} = Number of design gyrations

b. Calculate %G_{mm} @N_{ini} as follows:

Example:

$$\%G_{mm}@N_{ini} = 100 \times \left(\frac{G_{mb} \times h_d}{G_{mm} \times h_i}\right)$$

$$\%G_{mm}@N_{ini} = 100 \times \left(\frac{G_{mb} \times h_d}{G_{mm} \times h_i}\right) \qquad \qquad \%G_{mm}@N_{ini} = 100 \times \left(\frac{2.383 \times 110.0}{2.493 \times 123.1}\right) = 85.4\%$$

Where:

 G_{mm} N_{ini} = Percent theoretical maximum specific gravity $N_{initial}$

= Height of specimen at design gyration level

= Height of specimen at initial design gyration level

= Number of initial gyrations

c. Calculate Air Voids (V_a) as follow:

Example:

$$V_a = 100 \, \times \left(1 - \left(\frac{G_{mb}}{G_{mm}}\right)\right)$$

$$V_a = 100 \times \left(1 - \left(\frac{2.383}{2.493}\right)\right) = 4.4\%$$

Where:

V_a = Percent air voids

d. Calculate Voids in Mineral Aggregate (VMA) as follows:

Example:

$$VMA = 100 - \left(\frac{(G_{mb} \times P_s)}{G_{sb}}\right)$$

$$VMA = 100 - \left(\frac{(2.383 \times 94.8)}{2.630}\right) = 14.1\%$$

Where:

 P_s = Percent of aggregate in the mixture (100- P_b)

Example:

100% mix - 5.2% asphalt = 94.8% aggregate

G_{sb} = Bulk specific gravity of the combined aggregate VMA = Voids in Mineral Aggregate, percent

e. Calculate Voids Filled with Asphalt (VFA) as follows:

Example:

$$VFA = 100 \times \left(\frac{VMA - V_a}{VMA}\right)$$

$$VFA = 100 \times \left(\frac{14.1 - 4.4}{14.1}\right) = 68.8\%$$

Where:

VFA = Voids Filled with Asphalt, percent

f. Calculate Gravity Stone Effective (G_{se}) as follows:

$$G_{se} = \frac{100 - P_b}{\left(\frac{100}{G_{mm}} - \frac{P_b}{G_b}\right)}$$

$$G_{\text{se}} = \frac{100 - 5.2}{\left(\frac{100}{2.493} - \frac{5.2}{1.025}\right)} = 2.706$$

Where

G_{se} = Gravity Stone Effective (specific gravity of aggregates, excluding voids permeable to asphalt)

 P_b = Percent of binder 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:

Example:

$$P_{be} = P_{b} - \left(\frac{(P_{s} \times G_{b})(G_{se} - G_{sb})}{(G_{se} \times G_{sb})}\right) \qquad P_{be} = 5.2 - \left(\frac{(94.8 \times 1.025)(2.706 - 2.630)}{(2.706 \times 2.630)}\right) = 4.2$$

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 of 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_{h} = Percent of binder

h. Calculate dust-to-binder ratio (P_{200}/P_{be}) as follows:

Example: $5.0 \div 3.6 = 1.4$

$$P_{200}/P_{be} = P_{200} \div P_{be}$$

Where

 P_{200}/P_{be} = Dust-to-binder ratio

 P_{200} = Percent of aggregate passing the No. 200 sieve

7. Report

Report the results using one or more of the following of the following:

- Materials Testing System (MATS)
- WSDOT Form 350-560 for asphalt content, gradation, and moisture content
- WSDOT Form 350-162 for volumetric properties
- Form approved in writing by the State Materials Engineer

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).
- 1.3 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
R 76	Reducing Samples of Aggregate to Testing Size

¹This Standard Operating procedure is based on AASHTO T 323-04

	T 275	Bulk Specific Gravity of Compacted Hot Mix Asphalt (HMA) Using Paraffin-Coated Specimens
	T 283	Resistance of Compacted Asphalt Mixture to Moisture-Induced Damage
	T 304	Uncompacted Void Content of Fine Aggregate
	T 312	Preparing and Determining the Density of the Hot-Mix Asphalt (HMA) Specimens by Means of the Superpave Gyratory Compactor
2.2	Asphalt In	astitute
2.3	ASTM Sta	andards
2.4	WSDOT S	Standards
	Construct	ion Manual M 41-01
		Specifications for Road, Bridge, and Municipal Construction Specifications) M 41-10
	Materials	Manual M 46-01
	SOP 731	Method for Determining Volumetric Properties of Hot-Mix Asphalt (HMA)
	T 2	WSDOT FOP for AASHTO for Standard Practice for Sampling Aggregate
	T 27/11	WSDOT FOP for WAQTC/AASHTO for Sieve Analysis of Fine and Coarse Aggregates
	T 113	Method of Test for Determination of Degradation Value
	T 166	WSDOT FOP for AASHTO for Bulk Specific Gravity of Compacted Hot Mix Asphalt Using Saturated Surface-Dry Specimens
	T 176	WSDOT FOP for AASHTO for Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test
	T 209	WSDOT FOP for AASHTO for Method of Test for Maximum Specific Gravity of Hot Mix Asphalt Paving Mixtures "Rice Density"
	R 76	WSDOT FOP for AASHTO for Reducing Samples of Aggregates to Testing Size
	T 304	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 335	WSDOT FOP for AASHTO T 335 Determining the Percentage of Fracture in Coarse Aggregate
	T 718	Method of Test for Determining Stripping of Hot Mix Asphalt

Method of Preparation of Aggregate for HMA Mix Designs

Mixing Procedure for Hot-Mix Asphalt (HMA)

T 724

T 726

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-year period. For pavements designed for more or less than 15 years, determine the design ESALs for 15 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_{ba}) 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/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.

- 3.13 N_{initial}, N_{design}, N_{maximum} the number of gyrations defined in WSDOT *Standard Specification* 9-03.8(2).
- 3.14 Effective Asphalt Content (P_{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_{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 (Standard Specifications) for V_a, VMA, VFA, Dust/Asphalt Ratio at N_{design}, and relative density at N_{intial}.
 - **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%. The design binder content is selected on the basis of satisfactory conformance with the requirements of Section 9-03.8(2) of the *Standard Specifications* for V_a, VMA, VFA, and Dust/Asphalt Ratio (P₂₀₀/P_{be}) at N_{des}, and the relative density at N_{ini} and N_{max}. For WSDOT projects, 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: compacted to approximately 4.0% air voids in accordance with WSDOT FOP for AASHTO T 312, and evaluated according to WSDOT T 718. The design shall meet the tensile strength ratio requirement of 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 The asphalt binder grade will be indicated in WSDOT Contract Plans.
- 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 *Standard Specifications*.
- 6.4 Reduce the samples of aggregate fractions according to WSDOT FOP for AASHTO R 76 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:

$$P = Aa + Bb + Cc, etc. (1)$$

Where:

P = Percentage of material passing a given sieve for the combined aggregates A, B, C, etc.

A, B, C, etc. = Percentage of material passing a given sieve for aggregates A, B, C, etc. a, b, c, etc. = proportions of aggregates A, B, C, etc. used in the combination, and where the total = 1.00.

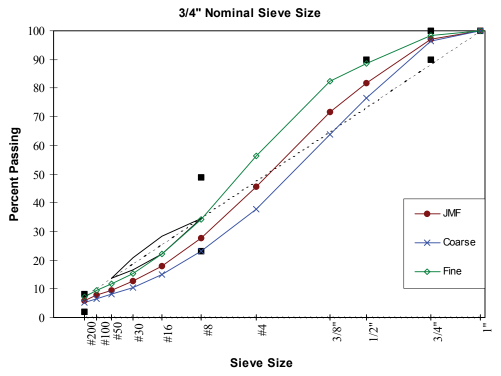
6.8 Prepare a minimum of three trial aggregate blend gradations; plot the gradation of each trial blend on a 0.45-power gradation analysis chart, and confirm that each trial blend meets the Aggregate Gradation Control Points in Section 9-03.8(6) of the *Standard Specifications* Gradation control is based 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						
Sieve Size	³⁄₃ in	½ in	³⁄₄ in	1 in		
1½"				X		
1"			X	X		
3/4"		X	X	X		
1/2"	X	X	X	X		
3/8"	X	X	X	X		
No. 4	X	X	X	X		
No. 8	X	X	X	X		
No. 16	X	X	X	X		
No. 30	X	X	X	X		
No. 50	X	X	X	X		
No. 100	X	X	X	X		
No. 200	X	X	X	X		

X = indicates sieve is required for gradation determination

6.9 Obtain a test specimen from each of the trial blends according to WSDOT FOP for AASHTO R 76, and conduct the quality tests specified in Section 9-03.8(2) subsections 1, 2, 3, and 4 of the *Standard Specifications* to confirm that the aggregate in the trial blends meets the minimum quality requirements specified in Section 9-03.8(2) of the *Standard Specifications*.

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.



Evaluation of the Gradations of Three Trial Blends (Example) Figure 1

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 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 $N_{\rm design}$ 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 shall not differ by more than 0.020.

Design ESALs ^a	Compaction Parameters			Typical Roadway Application ^b		
(million)	N _{initial} N _{design} N _{max}		N _{max}	,. ,		
< 0.3	6	50	75	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.		
0.3 to < 3	7	75	115	Applications include many collector roads or access streets. Medium-trafficked city streets and the majority of county roadways may be applicable to this level.		
3 to < 30	8	100	160	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.		
≥ 30	9	125	205	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.		

^aThe anticipated project traffic level expected on the design lane over a 15-year period. Regardless of the actual design life of the roadway, determine the design ESALs for 15 years.

Superpave Gyratory Compaction Effort Table 1

- 8.4 Determine the theoretical maximum specific gravity (G_{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 shall not differ by more than 0.011.

^bAs defined by A Policy on Geometric Design of Highways and Streets, 2001, AASHTO.

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 *Standard Specifications*.
- 9.2 Calculate V_a and VMA at N_{design} for each trial mixture using equations 2 and 3:

$$V_a = 100 \times \left(1 - \left(\frac{G_{mb}}{G_{mm}}\right)\right) \tag{2}$$

$$VMA = 100 - \left(\frac{G_{mb}P_{s}}{G_{sb}}\right) \tag{3}$$

Where:

 $G_{\rm mh}$ = Bulk specific gravity of the extruded specimen

 G_{mm}^{into} = Theoretical maximum specific gravity of the mixture P_s = Percent of aggregate in the mixture (100- P_b) G_{sh} = 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%, the actual air void content of the compacted specimen is unlikely to be exactly 4.0%. Therefore, the change in binder content needed to obtain a 4.0% 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%.

- 9.3 Estimate the volumetric properties at 4.0 percent air voids for each compacted specimen. On WSDOT projects, the gyration level will be specified in the Contract Provisions.
 - Determine the difference in average air void content at $N_{\rm design}$ (ΔV_a) of each aggregate trial blend from the design level of 4.0% using Equation 4:

$$\Delta V_a = 4.0 - V_a \tag{4}$$

9.3.2 Estimate the change in binder content (ΔP_h) needed to change the air void content to 4.0% using Equation 5:

$$\Delta P_b = -0.4 \, (\Delta V_a) \tag{5}$$

9.3.3 Estimate the change in VMA (Δ VMA) caused by the change in the air void content (ΔV_a) determined in Section 9.3.1 for each trial aggregate blend gradation, using Equations 6 or 7.

$$\Delta VMA = 0.2(\Delta V_a) \text{ if } V_a > 4.0 \tag{6}$$

$$\Delta VMA = -0.1(\Delta V_a) \text{ if } V_a < 4.0 \tag{7}$$

Note 13: A change in binder content affects the VMA through a change in the bulk specific gravity of the compacted specimen (G_{mb}) .

9.3.4 Calculate the VMA for each aggregate trial blend at N_{design} gyrations and 4.0% air voids using Equation 8:

$$VMA_{design} = VMA_{trial} + \Delta VMA \tag{8}$$

Where:

VMA_{design} = VMA estimated at a design air void content of 4.0% VMA_{trial} = VMA determined at the initial trial binder content

9.3.5 Using the values of ΔV_a determined in Section 9.3.1 and Equation 9, estimate the relative density of each specimen at $N_{\rm initial}$ when the design air void content is adjusted to 4.0 percent at $N_{\rm design}$:

$$\%G_{mm_{initial}} = 100 \times \left(\frac{G_{mb}h_d}{G_{mm}h_i}\right) - \Delta V_a$$
(9)

Where:

 $\%G_{mm_{\text{initial}}}$ = relative density at N_{initial} gyrations at the adjusted design binder content

 h_d = Height of the specimen after N_{design} gyrations, from the Superpave gyratory compactor, mm

 h_i = Height of the specimen after $N_{initial}$ gyrations, from the Superpave gyratory compactor, mm

9.3.6 Estimate the percent of effective binder (P_{be}) and calculate the Dust/Asphalt Ratio (P_{200}/P_{be}) for each trial blend using Equations 10 and 11:

$$P_{be_{est}} = -(P_s \times G_b) \frac{(G_{se} - G_{sb})}{(G_{se} \times G_{sb})} + P_{b_{est}}$$

$$\tag{10}$$

Where:

 $P_{be_{est}}$ = Estimated effective binder content

 P_s = Percent of aggregate in the mixture (100- P_b)

G_b = Specific gravity of the binder

 G_{se} = Effective specific gravity of the aggregate

 $G_{\rm sh}$ = Bulk specific gravity of the combined aggregate

 P_{bast} = Estimated binder content

Dust/Asphalt Ratio =
$$\frac{P_{200}}{P_{be}}$$
 (11)

Where:

 P_{200} = Percent passing the 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 *Standard Specifications*. 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.

	Trial Mixture (¾ Incl Projec	Outtour!			
	1	2	3	Criteria	
Volumetric Property	At the	Initial Trial Binder C	ontent		
P _b (trial)	4.4	4.4	4.4		
%G _{mm initial} (trial)	88.1	87.8	87.1		
%G _{mm design} (trial)	95.9	95.3	94.7		
$V_{\rm a}$ at $N_{\rm design}$	4.1	4.7	5.3	4.0	
VMA _{trial}	12.9	13.4	13.9		
	Adjustments	s to Reach Design Bin	der Content ($V_a = 4.0^{\circ}$	% at N _{design})	
ΔV_a	-0.1	-0.7	-1.3	Ğ	
ΔP_b	0.0	0.3	0.5		
ΔVMA	0.0	-0.1	-0.3		
	At the Estimated Design Binder Content ($V_a = 4.0 \%$ at N_{design})				
Estimated P _b (design)	4.4	4.7	4.9	<u> </u>	
VMA (design)	12.9	13.3	13.6	≥ 13.0	
%G _{mm initial} (design)	88.2	89.5	88.4	≤ 89.0	

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 $TV_a = 4.0$ percent, 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 (ΔP_b) and VMA (Δ 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% versus a requirement of ≥ 13.0%). Trial blend No. 2 exceeds the criterion for relative density at N_{initial} gyrations (89.5% versus requirement of ≤ 89.0%). Trial No. 3 meets the requirement for relative density and VMA and, in this example, is selected as the design aggregate structure.

Selection of a Design Aggregate Structure (Example) *Table 2*

10. Selecting the Design Binder Content

- 10.1 Prepare replicate mixtures (Note 8) containing the selected design aggregate structure at each of the following three binder contents: (1) the estimated design binder content, $P_{b \text{ (design)}}$; (2) 0.5% below $P_{b \text{ (design)}}$; and (3) 0.5% 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 $N_{\rm design}$ 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_{mm}) according to WSDOT FOP for AASHTO T 209 of each of the three 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_{design} gyrations using the following steps:
 - 10.5.1 Calculate V_a , VMA, and VFA at $N_{\rm design}$ using Equations 2, 3 and 12: The volumetric properties are determined for each specimen and then averaged for each replicate mixture.

$$VFA = 100 \times \left(\frac{VMA - V_a}{VMA}\right) \tag{12}$$

10.5.2 Calculate the Dust/Asphalt Ratio, using Equation 13.

Dust/Asphalt Ration
$$\frac{P_{200}}{P_{be}}$$
 (13)

Where:

 P_{be} = Effective binder content

10.5.3 For each of the three mixtures, determine the average corrected specimen relative densities at $N_{\rm initial}$ (%), using Equation 14.

$$\%G_{mm_{initial}} = 100 \times \left(\frac{G_{mb}h_d}{G_{mm}h_i}\right) \tag{14}$$

- 10.5.4 Plot the average V_a , VMA, VFA, and relative density at N_{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 $N_{\rm design}$.
- 10.5.6 By interpolation (Figure 2), verify that the volumetric requirements specified in Section 9-03.8(2) of the *Standard Specifications* are met at the design binder content.

- 10.6 Compare the calculated percent of maximum relative density with the design criteria at N_{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, $%G_{mm_x}$, 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 ΔV_a .
 - 10.6.3 Using Equation 14, determine the average corrected specimen relative densities at N_{initial} . Confirm that satisfies the design requirements in Section 9-03.8(2) of the *Standard Specifications* 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 $%G_{mm}$ satisfies the design requirements in Section 9-03.8(2) of the *Standard Specifications*.
 - 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_{max} , from Section 9-03.8(2) of the *Standard Specifications*.
 - 10.7.2 Determine the average specimen relative density at N_{max} , % $G_{mm_{\text{max}}}$, by using Equation 15, and confirm that satisfies the volumetric requirement in Section 9-03.8(2) of the *Standard Specifications*.

$$\%G_{mm_{max}} = 100 \times \frac{G_{mb}}{G_{mm}} \tag{15}$$

Where:

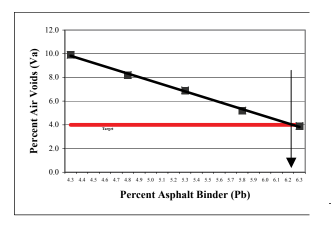
 $\%G_{mm_{max}}$ = Relative density at N_{max} gyrations at the design binder content

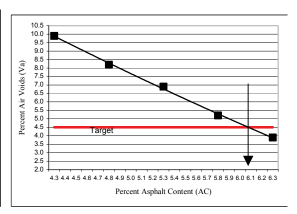
P _b (%)	V _a (%)	VMA (%)	VFA (%)	Maximum Density at N _{design} (G _{mm})	Density at N _{design} lbs/ft ³
4.3	9.9	17.0	41.8	2.660	165.6
4.8	8.2	16.7	50.9	2.636	164.1
5.3	6.9	16.6	58.5	2.617	162.9
5.8	5.2	16.5	68.5	2.585	160.9
6.3	3.9	16.2	76.0	2.574	160.2

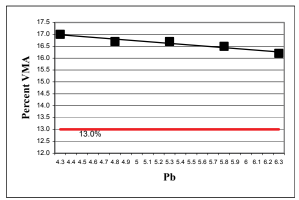
In this example, the estimated design binder content is 4.8 percent; the minimum VMA requirement for the design aggregate structure (¾ in nominal maximum size) is 13.0 percent, and the VFA requirements is 65 to 78 percent. Entering the plot of percent air voids versus percent binder content at 4.0 percent air voids, the design binder content is determined as 6.2 percent.

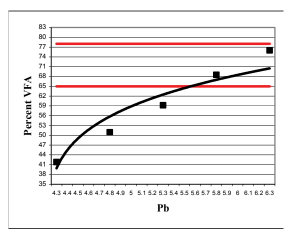
Entering the plots of percent VMA versus percent binder content and percent VFA versus percent binder content at 6.2 percent binder content, the mix meets the VMA and VFA requirement.

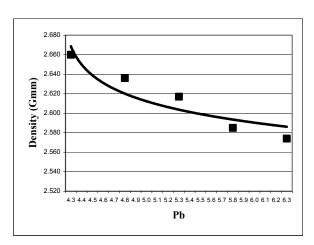
Sample Volumetric Design Data at N_{des} Figure 2











Sample Densification Curve Figure 3

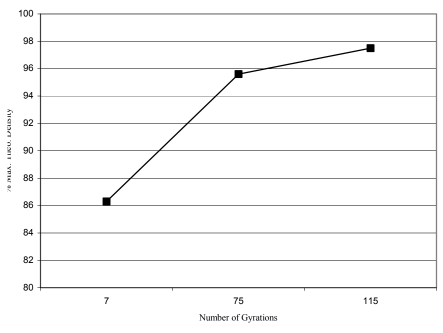


Figure 4

11. Evaluating Moisture Susceptibility

- 11.1 Prepare six mixture specimens composed of the design aggregate structure at the design binder content. Prepare the specimens according to WSDOT T 726, and compact the specimens to approximate 4.0% air voids in accordance to WSDOT FOP for AASHTO T 312. 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 WSDOT T 718.

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 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 No. 200 (0.075 mm) sieve of the mix will typically increase the VMA. If the percent passing the 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% 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 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.

13. Report

- 13.1 The report shall include the identification of the project number, mix class designation, and mix design number.
- 13.2 The report shall include information on the design aggregate structure including the source of aggregate, 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, V_a , and Dust/Asphalt Ratio P_{be} , G_{mm} , G_{mb} , G_{sb} and G_{se} of the aggregate blend, G_{sb} of the fine aggregate, and G_{b} .
- 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

A1.1Calculate 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:

$$G_{sb} = \frac{P_1 + P_2 + \dots + P_n}{\frac{P_1}{G_1} + \frac{P_2}{G_2} + \dots + \frac{P_n}{G_n}}$$
(16)

$$G_{sa} = \frac{P_1 + P_2 + \dots + P_n}{\frac{P_1}{G_1} + \frac{P_2}{G_2} + \dots + \frac{P_n}{G_n}}$$
(17)

Where:

 Bulk specific gravity for the combined aggregate
 Apparent specific gravity for the combined aggregate $G_{\operatorname{\mathsf{eh}}}$

 P_1 , P_2 , P_n = Percentages by mass of aggregates 1, 2, n G_1 , G_2 , G_n = Bulk specific gravities (Equation 16) or apparent specific gravities (Equation 17) of aggregates 1, 2, n.

A1.2Estimate the effective specific gravity of the combined aggregate in the aggregate trial blend using Equation 18:

$$G_{se} = G_{sb} + 0.8(G_{sa} - G_{sb}) ag{18}$$

Where:

 $G_{\rm se}={\rm Effective}$ specific gravity of the combined aggregate $G_{\rm sb}={\rm Bulk}$ specific gravity of the combined aggregate

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.

A1.3Estimate the volume of binder absorbed into the aggregate, V_{ba} , using Equations 19 and 20:

$$V_{ba} = W_s \left(\frac{1}{G_{sb}} - \frac{1}{G_{se}} \right) \tag{19}$$

 W_s = The mass of aggregate in 1 cm³ of mix, g, is calculated as

$$W_{S} = \frac{P_{S}(1 - V_{a})}{\frac{P_{b}}{G_{b}} + \frac{P_{S}}{G_{Se}}}$$
(20)

and Where:

 P_b = Percent of binder, in decimal equivalent, assumed to be 0.05

 P_s = Percent of aggregate in mixture, in decimal equivalent, assumed to be 0.95

 G_b = Specific gravity of the binder

 V_a = Volume of air voids, assumed to be 0.04 cm³ in 1 cm³ 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%.

A1.4Estimate the volume of effective binder using Equation 21:

$$V_{be} = 0.176 - (0.0675 \log (S_n)) \tag{21}$$

Where:

 V_{be} = Volume of effective binder, cm³ S_n = 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: $V_{be} = VMA - V_a =$ 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_{hi}) content for the aggregate trial blend gradation using Equation 22:

$$P_{bi} = 100 \times \left(\frac{G_b(V_{be} + V_{ba})}{\left(G_b(V_{be} + V_{ba}) \right) + W_s} \right)$$
 (22)

Where:

 P_{bi} = Estimated initial trial binder content, percent by weight of total mix

Determination of Pavement Density Differentials Using the Nuclear Density Gauge

1. Scope

This test method describes the procedure for locating and testing areas of suspected low cyclic density. Lower pavement density has been related to temperature differentials and areas of "spots, streaks" or visual pavement irregularities. This method uses infrared detection devices and visual inspection to identify areas of potentially low cyclic density.

2. Definitions

- a. Temperature Differential Area- Any area where the temperature of the newly placed HMA pavement is greater than 25° F different than the surrounding area.
- b. Aggregate segregation- "Spots, streaks" or visual pavement irregularities in the newly placed HMA pavement that has a significant difference in texture when compared to the surrounding material.
- c. Systematic Density Testing the testing of temperature differential areas or areas of aggregate segregation to determine if there is a pattern of low cyclic density.

3. 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 moisture-density gauge.
- c. Tape measure.
- d. A can of spray paint for marking test locations.
- e. Required report form.

4. Testing Criteria

- a. Where temperature differentials are 25° F or greater a systematic HMA compaction test is required.
- b. Where temperature differentials are less than 25° F a systematic HMA compaction test is not required unless, an area shows signs of visual pavement irregularities, surface segregation or a significantly different texture.

5. Determination of Systematic Density Testing Locations

Use either and infrared camera or a handheld non-contact infrared device to locate temperature differential areas as follows:

5.1 Infrared Camera

- a. Delineate a 500 ft section of pavement and systematically check the area for temperature differentials within one minute of HMA placement and prior to any compaction of the pavement.
- b. 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.
- c. Focus the camera on the freshly placed HMA pavement prior to compaction. Adjust the camera to show the high and low temperatures.
- d. Viewing should occur from the side of the paved lane approximately 15 to 20 feet back from the paver looking toward the paver.
- e. The "spot" function on the camera should be used to obtain the temperature of the cool area and the surrounding HMA to assess for temperature differentials.
- f. 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 HMA compaction 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 on DOT form 350-170 and in the MATS database.
- g. 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.

5.2 Handheld Noncontact Infrared Device

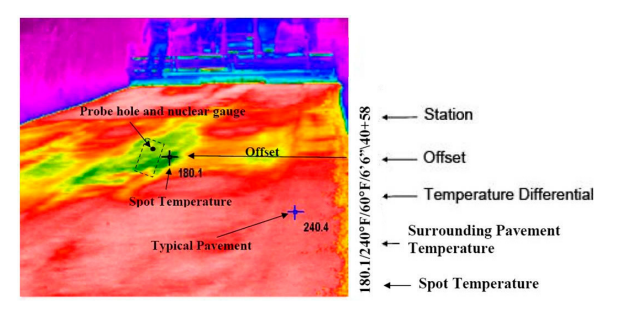
- a. Delineate a 500 ft section of pavement and systematically check the area for temperature differentials within one minute of HMA placement and prior to any compaction of the pavement.
- b. No temperature profiles shall be performed within the first or last 25 tons of production each day or within 25 feet of any transverse joint.
- c. Perform a longitudinal scan of the pavement by standing 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 500 ft of HMA placement.
- d. 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 can be captured with the longitudinal scan.

- e. Perform a transverse scan after completion of the longitudinal scan, making sure to scan the entire width of the paved lane excluding the outer 18 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.
- f. 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 testing 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 on DOT form 350-170 or in the MATS database.
- g. 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.

6. Systematic Density Testing Procedure

- a. Systematic density testing shall begin after finish rolling is completed.
- b. All systematic density testing shall be performed in accordance with WSDOT FOP for WAOTC T 355.
- c. Systematic density testing shall be performed at all the locations recorded in 5.1f and 5.2f of this procedure. Gauge probe shall be placed at the station and offset determined above as the center of the temperature differential area.
- e. If any temperature differentials are found in the initial assessment of the paving operations, at least one temperature profile shall be taken for every subsequent 500 ft of paving operation.
- d. If no temperature differentials or streaks greater than 25° F are found or if there are no more than 2 density readings lower than 90 percent found in a 500 ft section, the testing frequency may be reduced. Random checks however, should continue to be made throughout the day and the results recorded.
- e. 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.
- f. If it is found that the paving machine is creating areas that are significantly different in texture from the surrounding pavement, systematic density tests should be performed to determine if these are areas of low cyclic density.



Marking Location of Temperature Differential Figure 1

Sampling Hot Mix Asphalt After Compaction (Obtaining Cores)

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.
- When cores are used for forensic testing of HMA, refer to SOP 737 "Procedure for the Forensic Testing of HMA Field Cores" to determine the required number and size of cores.

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.

5. 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.

6. 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 1:** 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 2:** 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.

7. 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 Hot Mix Asphalt is available and used, it shall be compacted into the hole. A ready mix concrete or 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 driving surface.
- Prior to backfilling a core hole on a bridge deck, ensure that the hole and sidewalls are dry enough to bond with the sealant before applying.
- Acceptable sealants include; asphalt binder or any waterproof sealant designed for asphalt applications as stated by the manufacturer.
- Apply sealant to bottom surface and side walls of core hole as needed.
- Backfill the core hole with Hot Mix Asphalt, cold mix asphalt, ready mix concrete or grout and compact as needed.

8. 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.

9. 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 4: Lift lines are often more visible by rolling the core on a flat surface and/or surface drying the core.

10. Length Determination

Measure the thickness of the designated lift to the nearest 0.01' or 1/8" according to WSDOT Test Method 720.

11. 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 ½ ")
- Average Theoretical Maximum Density

SOP 734	Sampling Hot Mix Asphalt After Compaction (Obtaining Cores)
	This page intentionally left blank.
	This page intentionally left blank.

Standard Operating Procedure for Longitudinal Joint Density

1. General Scope

- a. This procedure describes the method for determining the location of a longitudinal joint density test.
- b. Longitudinal joint density tests are performed in addition to Quality Assurance (QA) density tests.
- c. One longitudinal joint density test will be performed on the confined or unconfined edge at each longitudinal joint.

2. Longitudinal Joint Testing

- a. The longitudinal joint density test will be conducted in accordance with WSDOT FOP for WAQTC T 355, except "Test Site Location, Section 1, subsection c, which is modified by this procedure to read "No closer than 18 in (450mm) to any vertical mass, or less than 6 in (152 mm) from a vertical pavement edge," making sure the gauge will sit flush with the hot-mix asphalt (HMA). See Figure 1.
- b. A longitudinal joint density will be required on the lane edge side of a shoulder if the shoulder is required to meet the same QA density requirements as the traveled lane.

Note: Hot lap joints are not included in longitudinal joint testing.

3. Number of Longitudinal Joint Tests

- a. For projects requiring 400 tons sublot with 5 sublots One reading, at each longitudinal joint to be tested, will be taken within each compaction lot at the same station location as the third sublot.
- b. For projects requiring 80 ton sublots One reading, at each longitudinal joint to be tested, will be taken every four hundred tons or at every fifth sublot tested.

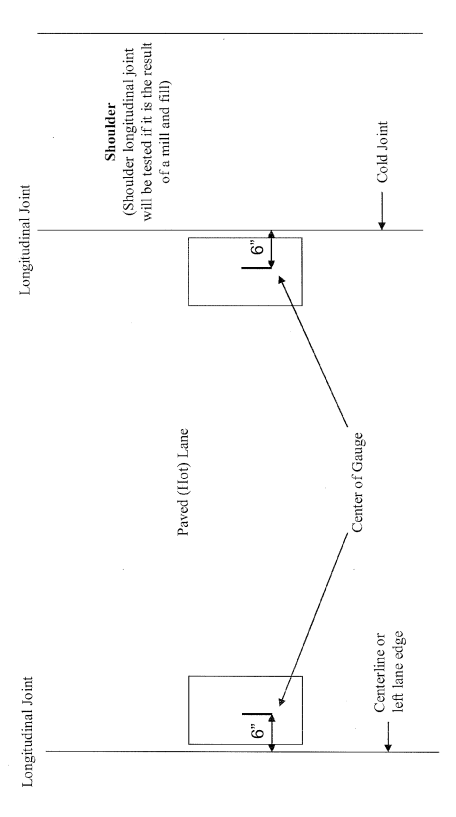
4. Calculation of Results

a. Calculate the Longitudinal Joint density in accordance WSDOT SOP 729.

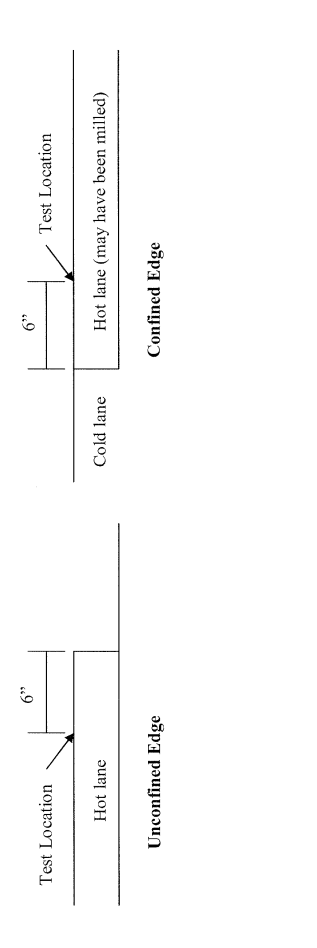
5. Report

- a. Report the results using one or more of the following:
 - Materials Testing System (MATS)
 - WSDOT Form 350-095
 - Form approved in writing by the State Materials Engineer

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.



Longitudinal Joint Testing Locations *Figure 1*



Longitudinal Test Location Examples Figure 2

In-Place Density of Bituminous Mixes Using Cores

1. Scope

This test method describes the procedure for determining the relative density of bituminous mixes by means of roadway cores.

2. Procedure

Cores for densities will be taken in accordance with WSDOT SOP 734 Sampling Hot Mix Asphalt After Compaction (Obtaining Core)

The bulk specific gravity (Gmb) of the core will be determined in accordance with WSDOT FOP for T 166 Bulk Specific Gravity of Compacted Hot Mix Asphalt (HMA) Using Saturate Surface-Dry Specimens.

The Theoretical maximum density of the mix will be determined in accordance with WSDOT FOP for AASHTO T 209 Theoretical Maximum Specific Gravity and Density of Hot-Mix Asphalt Paving Mixtures.

Determine the average theoretical maximum density in accordance with WSDOT SOP 729 Determination of the Moving Average of Theoretical Maximum Density (TMD) for HMA

3. Calculation of Percent of Compaction

The percent compaction is determined by comparing the density of the roadway core to the theoretical maximum density.

Calculate core density to the nearest 0.1 pcf as follows:

Core Density =
$$Gmb \times 62.245 pcf$$

Calculate percent compact (round to the nearest 0.1 percent) as follows:

Percent Compaction = (Core Density)/(Average Theoretical Maximum Density) \times 100

Procedure for the Forensic Testing of HMA Field Cores

1. Scope

This method describes the process for testing Hot Mix Asphalt (HMA) field cores for asphalt content, gradation, volumetric analysis, Hamburg Wheel-Test, Indirect Tensile Strength and asphalt binder grade determination.

- 1.1 This standard covers the procedural steps required for forensic testing of HMA field cores. Cores for forensic testing may range in size from 4-12 inches, although many specific test procedures require the core specimen to be six inches.
- 1.2 The values stated in English units are to be regarded as the standard.

2. Significance And Use

- 2.1 Approvals of the material for HMA are required prior to use per *Standard Specifications* Section 1-06.1.
- 2.2 Samples obtained in accordance with this procedure, shall be obtained using WSDOT SOP 734, "Sampling Hot Mix Asphalt after Compaction (Obtaining Cores)".

3. Reference Documents

Refer to applicable test methods within this procedure.

4. Apparatus

Refer to applicable test methods within this procedure.

5. 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.

6. Test Site Location

The sample location and quantity of cores to be obtained shall be determined by the test procedure to be performed or agency requirements.

7. Procedures

Perform procedures as needed to obtain desired test results:

7.1 Obtain cores per WSDOT SOP 734, "Sampling Hot Mix Asphalt after Compaction". The required quantity and size of cores for each procedure shall be as shown in Table 1:

Procedure	Size	Number of Cores	Special Instructions
AASHTO T331, "Standard Method of Test for Bulk Specific Gravity (Gmb) Density of Compacted Hot Mix Asphalt (HMA) Using Automatic Vacuum Sealing Method"	4" or 6"	1	·
WSDOT FOP for AASHTO T209, "Theoretical Maximum Specific Gravity Density of Hot Mix Asphalt Paving Mixtures"	4" or 6"	1	
WSDOT FOP for AASHTO T308, "Determining the Asphalt Binder Content of Hot Mix Asphalt by the Ignition Method"	6"	1	
WSDOT FOP for AASHTO T27/11, "MechanicalAnalysis of Extracted Aggregate"	6"	1	
WSDOT FOP for AASHTO T324, "Hamburg Wheel-Track Testing of Compacted Hot Mix Asphalt	6"	2	Obtain cores 6" apart for each determination
WSDOT FOP for ASTM D 6931, "Standard Test Method for Indirect Tensile Strength of Bituminous Mixtures"	6"	3	Obtain cores 6" apart for each determination
AASHTO R29, "Standard Practice for Grading or Verifying the Performance Grade (PG) of an Asphalt Binder"	6"	2	Obtain cores 6" apart for each determination

Table 1

- 7.2 Remove moisture from cores per AASHTO R 79, "Vacuum Drying Compacted Asphalt Specimens".
- 7.3 Determine core density per AASHTO T331, "Standard Method of Test for Bulk Specific Gravity (Gmb) and Density of Compacted Hot Mix Asphalt (HMA) Using Automatic Vacuum Sealing Method", and WSDOT FOP for AASHTO T209, "Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt Paving Mixtures". Theoretical Maximum Specific Gravity and Density of Hot Mix Asphalt Paving Mixtures data from corresponding field testing may be substituted in lieu of testing core material.
 - *Note 1*: AASHTO T331 shall be performed prior to WSDOT FOP for AASHTO T 209. Before performing T 209 all shaved or bare aggregate surfaces either from coring, surface wear or handling of the specimen shall be removed and separated from the specimen by carefully picking them from the specimen using a sharp tipped tool. Care must be taken not to remove fully coated aggregate. Removed particles shall be discarded and not included with the WSDOT FOP for AASHTO T209 test specimen.

- 7.4 Determine asphalt content per WSDOT FOP for AASHTO T308, "Determining the Asphalt Binder Content of Hot Mix Asphalt by the Ignition Method", if an ignition furnace correction factor (IFCF) is available. Otherwise, perform AASHTO T 164, "Standard Method of Test for Quantitative Extraction of Asphalt Binder from Hot Mix Asphalt".
- 7.5 Determine aggregate sieve analysis per WSDOT FOP for AASHTO T 27/11, "Mechanical Analysis of Extracted Aggregate". WSDOT FOP for AASHTO T27/11 shall be performed following binder extraction per WSDOT FOP for AASHTO T 308, "Determining the Asphalt Binder Content of Hot Mix Asphalt by the Ignition Method" or AASHTO T164, "Standard Method of Test for Quantitative Extraction of Asphalt Binder from Hot Mix Asphalt".
- 7.6 Determine rutting and moisture-susceptibility of HMA per WSDOT FOP for AASHTO T324, "Hamburg Wheel-Track Testing of Compacted Hot Mix Asphalt".
- 7.7 Determine Indirect Tensile Strength (IDT) per WSDOT FOP for ASTM D6931, "Standard Test Method for Indirect Tensile Strength of Bituminous Mixtures".
- 7.8 Determine grade of asphalt per AASHTO R29, "Standard Practice for Grading or Verifying the Performance Grade (PG) of an Asphalt Binder". Extract the binder in accordance with AASHTO R59, "Recovery of Asphalt Binder from Solution by Abson Method" or ASTM D1856, "Standard Test Method for Recovery of Asphalt from Solution by Abson Method", for each asphalt grade determination.
 - *Note 2*: Binder specimens for AASHTO R29, Standard Practice for Grading or Verifying the Performance Grade (PG) of an Asphalt Binder may be obtained in conjunction with AASHTO T164, Standard Method of Test for Quantitative Extraction of Asphalt Binder from Hot Mix Asphalt.

WSDOT Test Method No. 802

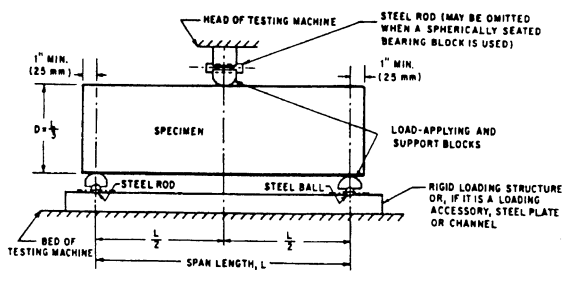
Method of Test for Flexural Strength of Concrete (Using Simple Beam With Center-Point Loading)

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 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).



Note-Apparatus may be used inverted.

Diagrammatic View of Apparatus for Flexure Test of Concrete be Center-point Loading Method

Figure 1

3. Test Specimen

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 12-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:

$$R = \frac{3P1}{2bd^2}$$

Where:

R= Modulus of rupture in psi or MPa

P = Maximum applied load indicated by the testing machine in lb•f or N

1 = 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 (0.03MPa),
 - (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

Pai	rticipant Name Exam Date				
Pro	ocedure Element				
Pre	eparation epitation	Yes	No		
1.	Copy of current procedure available at test site?				
2.	In the field, Jack properly calibrated?				
3.	Beam turned on its side with respect to its position when molded, and centered on the supporting bearing blocks?				
4.	Load applying block brought into contact with the beam at the center line between the supports?				
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?				
6.	Load applied to the beam at the proper uniform rate?				
Eq	uipment				
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?				
Fir	st Attempt: Pass Fail Second Attempt: Pass Fail				
Sig	gnature of Examiner				
Cor	nments:				

WSDOT FOP for C 8051

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 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. Significance and Use

3.1 This test method is not intended as the basis for acceptance or rejection of concrete because of the inherent uncertainty in the estimated strength.

4. Apparatus

- 4.1 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:* Use type N rebound hammers that are commercially available to accommodate testing of various sizes and types of concrete construction.
- 4.2 Abrasive Stone Consisting of medium-grain texture silicon carbide or equivalent material.
- 4.3 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 method ASTM E 18. An instrument guide is provided to center the rebound hammer over the impact area and keep the instrument perpendicular to the surface.

¹This FOP is based on ASTM C 805 and has been modified per WSDOT standards. To view the redline modifications, contact the WSDOT Quality Systems Manager at 360-709-5412.

- 4.4 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 Section 4.3. 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 Section 4.3. 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.

5. Test Area and Interferences

- 5.1 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. Troweled surfaces generally exhibit higher rebound numbers than screeded or formed finishes. If possible, test structural slabs from the underside to avoid finished surfaces.
- 5.2 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 Section 4.2. Smooth-formed or troweled surfaces do not have to be ground prior to testing. Do not compare results from ground and unground surfaces.
- 5.3 Do not test frozen concrete.
 - **Note 3:** 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 two or three units (1 unit = 1 whole number).
- 5.4 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.
- 5.5 Do not conduct tests directly over reinforcing bars with cover less than 0.75 in (20 mm).
 - **Note 4:** The location of reinforcement may be established using reinforcement locators or metal detectors. Follow the manufacturer's instructions for proper operation of such devices.

6. Procedure

6.1 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.

7. Calculation

7.1 Discard readings differing from the average of ten readings by more than six units and determine the average of the remaining readings. If more than two readings differ from the average by six units, discard the entire set of readings and determine rebound numbers at ten new locations within the test area.

8. Report

- 8.1 Report the following information for each test area:
 - 8.1.1 Date and time of testing.
 - 8.1.2 Identification of location tested in the concrete construction and the type and size of member tested.
 - 8.1.2.1 Description of the concrete mixture proportions including type of coarse aggregates if known.
 - 8.1.2.2 Design strength of concrete tested.
 - 8.1.3 Description of the test area including:
 - 8.1.3.1 Surface characteristics (trowelled, screeded) of area.
 - 8.1.3.2 If surface was ground and depth of grinding.
 - 8.1.3.3 Type of form material used for test area.
 - 8.1.3.4 Curing conditions of test area.
 - 8.1.3.5 Type of exposure to the environment.
 - 8.1.4 Hammer identification and serial number.
 - 8.1.4.1 Air temperature at the time of testing.
 - 8.1.4.2 Orientation of hammer during test.
 - 8.1.5 Average rebound number for test area.
 - 8.1.5.1 Remarks regarding discarded readings of test data or any unusual conditions.

10. Precision and Bias

See ASTM C 805 precision and bias.

Performance Exam Checklist

Rebound Hammer Determination of Compressive Strength of Hardened Concrete FOP for ASTM C 805

Participant Name	Exam Date
Procedure Element Preparation	Yes No
1. Copy of current procedure available at test site?	
2. Hammer properly serviced and calibrated or verified?	
3. Test location properly prepared?	
4. Test location meets minimum size requirement?	
5. Ten acceptable readings taken in each test area?	
6. Readings properly spaced in test area?	
7. Test readings properly converted to estimated strength?	
8. Test information properly recorded?	
9. All calculations performed correctly?	
Equipment	
10. Are calibration/verifications tags present on equipment us	sed in this procedure?
11. All equipment functions according to the requirements of	this procedure?
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 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 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 longitudinally and the actual change in elevation 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.

3. Procedure

a. Equipment – To determine the Profile Index, use a plastic scale 1.70 in wide and 1.76 LF long representing a pavement length of 528 LF at a scale of 1 in = 25 LF. A plastic scale for the Profilograph may be obtained by the regions from the State Materials Laboratory. Near the center of the scale is an opaque band 0.2 in wide extending the entire length of 21.12 in. On either side of this band are scribed lines 0.1 in 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."

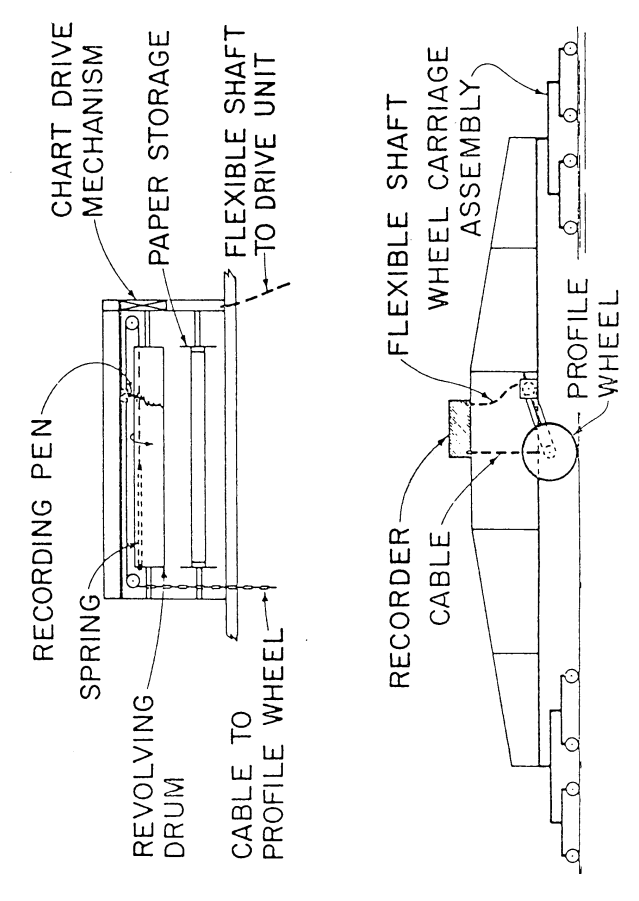


Figure 1

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, the 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 superelevated 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 (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 or more and extend longitudinally for 2 LF (0.08 in 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 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. If not, its length should be scaled. An example follows:

	Section Length, miles	Counts, tenth of an inch
	0.10	5.0
	0.10	4.0
	0.10	3.5
400 ft =	0.076	2.0
Total	0.376	14.5

The Profile Index is determined as "inches per mile in excess of the 0.2 in 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., total count = 14.5 tenths of an inch

Profile Index =
$$\frac{1 \text{ mile}}{\text{length of profiles in miles}} \times \text{a total count in inches}$$

Pr I = $\frac{1 \text{ mile}}{0.376 \text{ mile}} \times 1.45 = 3.9$

(Note that the formula uses the count in inches rather than tenths of an inch and is obtained by dividing the count by ten.)

PROFILOGRAMS DERIVING PROFILE INDEX FROM OF SHOWING METHOD EXAMPLE

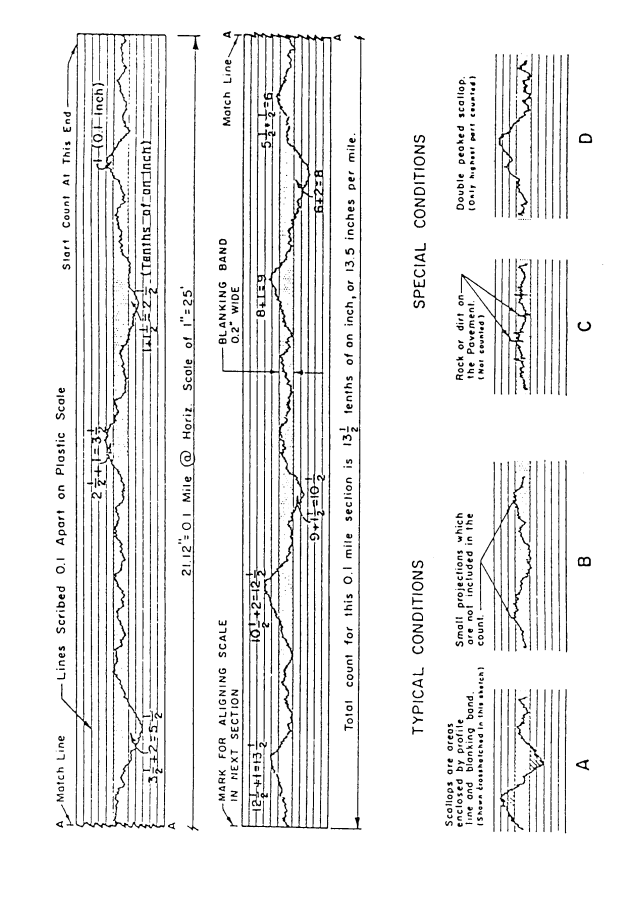


Figure 2

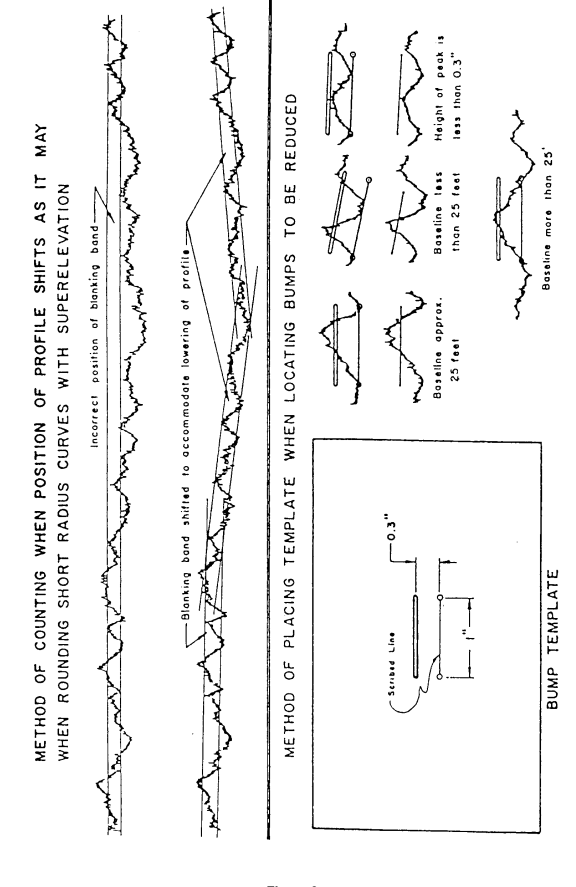


Figure 3

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:

		Counts, tenths of an inch		
	Section Length, miles	Left wheel track	Right wheel track	
	0.10	5.0	4.5	
	0.10	4.0	5.0	
	0.10	3.5	3.0	
400 LF =	0.076	2.0	1.5	
Total	0.376	14.5	14.0	
Prl (by formula)		3.9	3.7	
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 Mile 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 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.

Part III Determination of High Points in Excess of 0.3 in

4. Procedure

- a. Equipment Use a plastic template having a line 1 in long scribed on one face with a small hole or scribed mark at either end, and a slot 0.3 in from and parallel to the scribed line (see Figure 3). (The 1 in line corresponds to a horizontal distance of 0.3 in on the horizontal scale of the profilogram.) The plastic template may be obtained from the State Materials Laboratory.
- b. Locating High Points in Excess of 0.3 in 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.

There may be instances where the distance between easily recognizable low points is less than 1 in (25 LF). 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 25 LF as possible, but in no case to exceed this value. When the distance between prominent low points is greater than 25 LF, 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.

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, $6 \text{ in} \times 6 \text{ in} \times 21 \pm \frac{1}{2} \text{ in} (150 \text{ mm} \times 150 \text{ mm} \times 550 \pm 13 \text{ mm})$ or $8 \text{ in} \times 8 \text{ in} \times 26 \pm \frac{1}{2} \text{ in} (200 \text{ mm} \times 200 \text{ mm} \times 670 \pm 13 \text{ mm})$.
- b. Vibrator, capable of 7,000 vibrations per minute with a diameter not less than $\frac{3}{4}$ in (19.0 mm) or greater than $\frac{1}{2}$ in (38.1 mm).
- c. Tamping Rod The tamping rod is a round, straight steel rod \(\frac{5}{8} \) in (16.0 mm) diameter and approximately 24 in (610 mm) long, having the tamping end rounded to a \(\frac{5}{8} \) in (16.0 mm) diameter hemispherical tip.
- d. Mallet A mallet with a rubber or rawhide head weighing 1.25 ± 0.50 lb $(0.57 \pm 0.23$ 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 ft³ (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 in² (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 $73.4^{\circ} \pm 3^{\circ}$ F ($23^{\circ} \pm 2^{\circ}$ 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 $73.4^{\circ} \pm 5^{\circ}$ F ($23^{\circ} \pm 2.8^{\circ}$ 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.

Method for Making Flexural Test Beams WSDOT T 808

Par	ticipant Name Exam Date		
Pro	ocedure Element	Yes	No
1.	Copy of current procedure available at test site?		
2.	Making of test specimens begins within 15 minutes for sampling?		
3.	Assemble of molds on a rigid surface free from vibration and other disturbances?		
4.	Concrete remixed to a uniform appearance?		
5.	When method of concrete consolidation is by rodding: a. Mold filled in two layers?		
	b. Each layer rodded one time for each 2 in ² (1290 mm) of mold surface area?		
	c. Rodding, evenly distributed over the entire surface area?		
	d. After rodding each layer, mold tapped lightly 10-15 times with mallet?		
6.	When method of concrete consolidation is by internal vibrators: a. Mold filled in a single layer?		
	b. Vibrator inserted at intervals not to exceed 6 in (150 mm) along the centerline of the long dimension?		
	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?		
	d. Mold tapped lightly 10-15 times with mallet?		
7.	Top of mold properly finished?		
8.	Top of mold properly treated to prevent moisture loss?		
Eq	uipment		
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?		
Firs	st Attempt: Pass Fail Second Attempt: Pass Fail		
Sig	nature of Examiner		

Comments:

Method of Test for Determination of the Density of Portland Cement Concrete Pavement Cores

1. Scope

a. This method of test is intended for use in determining the density of Portland cement concrete pavement cores. The object of this test is to determine the in place density of the concrete as it exists. This density is the value desired for comparison to the density of freshly mixed portland cement as determined by AASHTO T 121 or by the densities on the mix design.

2. Equipment

- a. Balance Capacity sufficient for the masses required by the test procedure, accurate to 0.1 percent of the sample mass or better and conforms to the requirements of AASHTO M 231.
- 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. Thermometer The Thermometer 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 and 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 24 hours after coring.

Temperature °F	Pounds Per Cubic Foot	Temperature °F	Pounds Per Cubic Foot
65	62.336	74	62.269
66	62.329	75	62.261
67	62.322	76	62.252
68	62.315	77	62.243
69	62.308	78	62.234
70	62.301	79	62.225
71	62.293	80	62.216
72	62.285		
73	62.277		

Unit Mass of Water Table 1

- 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:

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_w = Density of the water at the test temperature (see Table 1)

Calculate the density to the nearest 0.1 lb per ft³ (1 kg per m³).

5. Reproducibility of Results

a. Duplicate determinations should check to within 0.1 lb per ft³ (3 kg per m³).

6. Reports

a. The test results will be reported on the appropriate test data sheet.

Method of Test for Determination of the Density of PCC Pavement Cores (WSDOT TM 810)

Part	icipant Name				Exam D	ate		
Pro	cedure Elemer	nt						Yes No
1.	The tester has	a copy	of the cur	rent procedure	on hand?			
2.			_	ecording to the ication tags pro	test procedure, and i esent?	f required	d,	
3.	Finished pave	ment co	red after a	a minimum of	24 hours of curing?			
4.	Core's moistu	re conte	nt preserv	ved in bags or b	y immersion?			
5.	Density deterr	mined w	ithin 24 h	ours of coring	?			
6.	Core washed to	thorougl	nly?					
7.	Weight in wat	er deter	mined to r	nearest gram?				
8.	Temperature of	of water	determine	ed to nearest de	egree?			
9.	Core rolled or appearing dan		ent towel	removing visib	ole films of water but	t still		
10.	Core not over-	-dried or	r allowed	to evaporate?				
11.	Weight of surf	face-dry	core dete	rmined to near	est gram?			
12.	All calculation	ns perfo	rmed corr	ectly?				
Firs	t Attempt: Pa	ss]	Fail		Second Attempt:	Pass	Fail	
Sign	nature of Exam	iner _						
Con	nments:							

Method of Test for Measuring Length of Drilled Concrete Cores

1. Scope

a. This method is for the WSDOT ten point callipering device it is similar to AASHTO T- 48 and covers the procedure for determining the length of a core drilled from a concrete structure, and from Portland cement concrete pavement. It is acceptable to use the commercially available nine point callipering device defined in AASHTO T 148.

2. Apparatus

- a. The apparatus shall be a callipering 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 ½ inch more than ½ 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 callipering 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 (See Note 1) 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.
 - **Note 1:** Commercially available nine point callipering device is acceptable.
- 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 ½ 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 shall be obtained per AASHTO T 24. 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 (See Note 2) 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.
 - **Note 2:** For commercially available callipering devices nine measurements is allowed.
- 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 (See Note 3), expressed to the nearest 0.01 foot, Measurements will be reported in the Materials Tracking System (MATS) database.
 - **Note 3:** For commercially available callipering devices average nine measurements.

Method T 812 Checklist Measuring Length of PCC Cores

Participant Name	Exam Date	_
Procedure Element		Yes No
1. Only concrete measured?		
2. Damaged cores not measured?		
3. Apparatus calibrated?		
4. Smooth (top) end of core set on pins?		
5. Center probe located at center of core?		
6. Ten measurements taken?		
7. Measurements read to 0.10 in directly?		
8. Measurements read indirectly to 0.01 in?		
9. Measurements recorded to 0.01 in?		
10. Averaged and reported to 0.01 foot?		
First Attempt: Pass Fail	Second Attempt: Pass	Fail
Signature of Examiner		

Comments:

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 Cı	ıbe 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 Compartment	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 Faces*	90 + 0.5°	90 + 0.5°	90 + 0.5°	90 + 0.5°

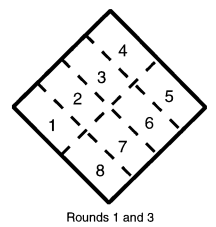
^{*}Measured at points slightly removed from the intersection. Measured separately for each compartment between all the interior faces and the adjacent face and between interior faces and top and bottom planes of the mold.

Permissible Variations of Specimen Molds Table 1

- 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.
- 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. Begin molding the specimens within an elapsed time of not more than $2\frac{1}{2}$ 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.



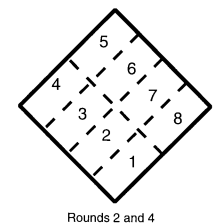


Figure 1

- 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, avoid freezing of cubes and record the time. Allow the sample to set undisturbed, away from vibration, for a minimum of four 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 Concrete 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 to 36 hours old.
- k. When the specimens are 24 to 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.

Field Method of Fabrication of 2 in (50 mm) Cube Specimens for Compressive Strength Testing of Grouts and Mortars WSDOT Test Method T 813

Part	ticipant Name Exam Date		
Pro	cedure 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.	Three cubes made for each time period of test?		
4.	All joints (mold halves, mold to base plate) shall be water tight?		
5.	Adequate coating of release agent applied to interior surfaces of the mold?		
6.	Molding began within 2½ minutes from completion of mixing?		
7.	Molding performed in two lifts? (Not necessary if mix is fluid.)		
8.	Lifts tamped 32 times, made up of 4 rounds of 8, each perpendicular to the other? (Not required if mix is fluid.)		
9.	For second layer, mortar forced out of the mold brought back in before each round? (Not required if mix is fluid.)		
10.	Mix extends slightly above the mold at the completion of tamping?		
11.	Mortar smoothed by drawing flat side of trowel across each cube at right angles?		
12.	Mortar leveled by drawing the flat side of trowel lightly along the length of mold?		
13.	Mortar cut off flush with mold with edge of trowel using sawing motion?		
14.	Time of molding recorded?		
15.	Cover plate placed on top of the mold and covered with wet burlap, towel or rag?		
16.	Covered sample sealed in a plastic sack in a level location out of sunlight?		
17.	Sample delivered to the laboratory in the mold within 24 hours?		
18.	Transmittal includes the time of molding?		
Firs	et Attempt: Pass Fail Second Attempt: Pass Fail		
Sign	nature of Examiner		

Comments:

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° ± 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° ± 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 $73.4^{\circ}F \pm 3.6^{\circ}F$ ($23^{\circ} \pm 2^{\circ}C$) 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 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 $(\pm 0.05 \text{ g})$ 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 ½ 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° ± 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.



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 the procedure:
 - a. Change item 3 to read: Fill the base completely in one lift.
 - *Note:* Filling the base with concrete by using multiple scoops or by pouring from a bucket or similar container has been found to be acceptable.
 - b. 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.
 - c. Delete items 5 through 11.

Air Content of Freshly Mixed Self Compacting Concrete by the Pressure Method WSDOT T 818

Parti	icipant Name Exam Date			
Pro	cedure 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.	Container filled in one layer, slightly overfilling?			
4.	Sides of the container lightly tapped with the mallet?			
5.	Concrete struck off level with top of container using the bar and rim cleaned off?			
6.	Inside of cover cleaned and moistened before clamping to base?			
Usir	ng a Type B Meter			
7.	Both petcocks open?			
8.	Air valve closed between air chamber and the bowl?			
9.	Water injected through petcock until it flows out the other petcock?			
10.	Water injection into the petcock continued while jarring and tapping the meter to insure all air is expelled?			
11.	Air pumped up to initial pressure line?			
12.	A few seconds allowed for the compressed air to stabilize?			
13.	Gauge adjusted to the initial pressure?			
14.	Both petcocks closed?			
15.	Air valve opened between chamber and bowl?			
16.	Sides of bowl tapped with the mallet?			
17.	Air percentage read after lightly tapping the gauge to stabilize the hand?			
18.	Air valve closed and then petcocks opened to release pressure before removing the cover?			
19.	Air content recorded to 0.1 percent?			
20.	All calculations performed correctly?			
First	t Attempt: Pass Fail Second Attempt: Pass Fail			
Sign	nature of Examiner			
This	checklist is derived, in part, from copyrighted material printed in ACI CP-1, published	by the		

American Concrete Institute.

Comments:



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:
 - 9. Molding Specimens
 - 9.2 Casting Cylinders is revised to read:

Place the concrete in the mold using a scoop, blunted trowel or shovel. Molds shall be filled in one layer by pouring material from a suitable container into the mold. Do not rod, vibrate, or tap the mold.

Note: Filling the mold with concrete by using multiple scoops or by pouring from a bucket or similar container has been found to be acceptable.

- 9.3 Consolidation is deleted
- 9.4 Finishing is revised to read:

Strike off the surface of the concrete level with the top of the mold using a float, trowel or steel strike off bar. Immediately after finishing place a plastic cylinder lid on the cylinder.

Making and Curing Self-Compacting Concrete Test Specimens in the Field WSDOT T 819

Participant Name	Exam I	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 be	gun within 15 minutes of sampling?	
4. Concrete poured into the	e mold using a suitable container?	
5. Mold filled in one lift?		
6. Excess concrete struck of	off?	
7. Specimens covered imm	nediately with plastic cylinder lid?	
First Attempt: Pass Fa	il Second Attempt:	Pass Fail
Signature of Examiner		
This checklist is derived, in p American Concrete Institute.	art, from copyrighted material printed in A	ACI CP-1, published by the
Comments:		

Bond Strength (Diagonal Shear) ASTM C 882

Participant Name

Pro	ocedure Element	Yes No				
1.	Confirmation that the concrete lab will be ready to mix mortar?					
2.	Half cylinders taken from moisture room?					
3.	Elliptical surface acid washed, rinsed with water and allowed to dry?					
4.	Half cylinders placed in cylinder molds awaiting the epoxy and mortar?					
5.	Epoxy mixed using manufacturer's mix ratio and directions?					
6.	With elliptical surface maintained horizontal, coat the surface with epoxy?					
7.	Epoxy layer thickness 10 to 15 mils (0.3 mm to 0.4 mm)?					
Aft	er the epoxy coating:					
8.	Fill up the remainder of the mold with fresh plastic mortar?					
9.	New mortar rodded and the top smoothed off?					
10.	Specimens (epoxy/mortar) cured for two (2) days or fourteen (14) days at 100% relative humidity?					
11.	Compressive load determined as per WSDOT Test Method T 22?					
12.	Compressive strength calculated based on elliptical surface area?					
Firs	st Attempt: Pass Fail Second Attempt: Pass Fail					
Sig	nature of Examiner					
Con	nments:					

Exam Date

Practice for Sampling of Geosynthetic Material for Testing

1. Scope

a. This practice covers the procedure for sampling Geosynthetic Material for testing.

2. Definitions

- a. Geogrid A regular network of integrally connected polymer tensile elements with an aperture geometry sufficient to permit mechanical interlock with the surrounding backfill.
- b. Geosynthetic Material general term which includes all geotextiles, geogrids, and prefabricated drainage mats.
- c. 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.
- d. Lot All geosynthetic material 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 geosynthetic material product.
- e. Lot Sample Sample(s) from one or more geosynthetic material rolls taken at random to represent an acceptance sampling lot and used as a source of laboratory samples.
- f. Production Unit As referred to in this practice, it shall be considered to be synonymous with the geosynthetic material roll as shipped by the manufacturer. Two or more geosynthetic material rolls joined together by sewn seams shall be considered as separate rolls.
- g. Minimum Average Roll Value The test results of any sampled roll in a lot shall meet or exceed the minimum values specified.

3. Significance and Use

- a. Sampling is an important part of testing and the sampler should make every effort to obtain samples that will show the nature and condition of the materials they represent.
- b. This sampling procedure will provide a representation of the lot which is adequate to establish minimum average roll values as defined by this practice.

4. Procedure

- a. Divide the shipment or consignment into lots as defined in 2.d.
- b. Determine the number of rolls in the shipment or consignment to be sampled using Table 1.

Number of Rolls in Lot	Number of Rolls to be Selected for Lot Sample
1 to 24	1
25 to 49	2
50 to 99	3
100 to 125	5
125 to 216	6
217 to 343	7
344 to 512	8
513 to 729	9
730 to 1,000	10

Number of Rolls to be Selected as Lot Sample Table 1

- c. Laboratory sample selection.
 - (1) Obtain a laboratory sample from each roll in the Lot Sample. The sample shall be a minimum of 6 feet long by the full width of the geosynthetic material roll with a total area greater than or equal to 6.0 yd².
 - (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 very ends of the roll).
 - (3) Protect the sample from exposure to Ultraviolet light.

5. Sample Submittal

- a. All geotextile samples submitted to the State Material Laboratory are to be prepared and shipped as follows:
 - 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.
- 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) Roll number(s).
 - (4) Proposed use(s).
 - (5) Certified test results from the manufacturer.
 - (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.

Practice for Sampling Geosynthetic Material for Testing WSDOT Test Method T 914

Participant Name		Exam D	ate		
Procedure Element				Yes N	٧o
1. The tester has a copy	y of the current proce	edure on hand?			
2. Sampling					
	nt/consignment into lolls to be sampled.	lot(s) and used Table 1 to o	determine	:	
b. Rolls to be samp	oled selected at rando	om.			
*		g by the full width of the g than or equal to 6.0 yd ²	geosynthe	etic	
d. Sample does not	include outer wrap	or inner wrap of the roll.			
3. Shipment Preparatio	on				
 a. Roll sample arou cardboard mailir 		ninimum, tube such as PC	V pipe or		
b. Wrap the sample	e to protect from ultra	a-violet light exposure.			
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 $70^{\circ} \pm 4^{\circ}F$ ($21^{\circ} \pm 2^{\circ}C$).
- 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 $70^{\circ} \pm 4^{\circ}$ F ($21 \pm 2^{\circ}$ 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 $70^{\circ} \pm 4^{\circ}$ F (21 ± 2°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

Summary and Use of Standard Practice

Abbreviations and Symbols

Definitions

Test Methods and Practices Used

Data Requirements for Initial Product Acceptance

- 1. General Product Information (required for all geosynthetic reinforcement products):
- 2. Installation Damage Data Requirements (RF_{ID}):
- 3. Creep Data Requirements (RF_{CR} and Creep Stiffness J):
- 4. Long-Term Durability Data Requirements (RF_D):7
- 5. Evaluation of Product Lines

Determination of Long-term Geosynthetic Strength for Initial Product Acceptance

- 1. Calculation of Long-Term Strength
- 2. Wall or Slope Class
- 3. Environment Aggressiveness
- 4. Requirements for Class 1 Walls and Slopes to Determine T_{al}
- 5. Requirements for Class 2 Walls and Slopes to Determine T_{al}
- 6. Minimum Polymer and Physical Property Requirements to Allow Use of Default Reduction Factors for RF and RF_D in Nonaggressive Environments

Quality Assurance Requirements for Products that have been Through Initial Acceptance

- 1. Data Verification Requirements
- 2. Quality Assurance (QA) Testing Approach
- 3. Quality Assurance (QA) Sampling
- 4. Quality Assurance (QA) Testing
 - A. Installation Damage Testing
 - B. Creep Testing
 - C. Durability Testing

- 5. Quality Assurance (QA) Criteria for Comparison to Initial Product Acceptance Test Results
 - A. Short-term Index Tensile Testing
 - B. Installation Damage Testing
 - C. Creep Rupture Testing for Prediction of Creep Limit
 - D. Creep Strain Testing for Prediction of Creep Limit
 - E. Assessment of the Creep Stiffness at Low Strain
 - F. Durability Testing

References

WSDOT Test Method No. 925, Appendix A

References

WSDOT Test Method No. 925, Appendix B

- B.1 Overview of Extrapolation Approach to Determine the Ultimate Limit State Creep Limit, T₁
- B.2 Step-By-Step Procedures for Extrapolating Creep Rupture Data –Conventional Method
- B.3 Procedures for Extrapolating Creep Rupture Data Stepped Isothermal Method (SIM)
- B.4 Determination of RF_{CR}
- B.5 Use of Creep Data from "Similar" Products and Evaluation of Product Lines

References

WSDOT Test Method No. 925, Appendix C

- C.1 Creep Strain Assessment Tools and Concepts
- C.2 Creep Strain Data Extrapolation
 - C.2.1 Step-By-Step Procedures for Extrapolating Creep Strain Data Conventional Method
 - C.2.2 Step-By-Step Procedures for Extrapolating Creep Strain Data Stepped Isothermal Method (SIM)
- C.3 Determination of RF_{CR}
- C.4 Estimation of Long-Term Creep Deformation
- C.5 Estimation of Creep Stiffness for Working Stress Design
- C.6 Evaluation of Product Lines

References

WSDOT Test Method No. 925, Appendix D

References



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 qualification or 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.

Abbreviations and Symbols

AASHTO = American Association of State Highway and Transportation Officials

 \mathbf{d}_{50} = The grain size at 50% passing by weight for the backfill.

HDPE = High Density Polyethylene

MARV = 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 = Mechanically Stabilized Earth

PET = Polyester

PP = Polypropylene

QPL = Qualified Products List

RF = Combined reduction factor to account for long-term degradation due

to installation damage, creep, and chemical/biological aging

 \mathbf{RF}_{CR} = Strength reduction factor to prevent long-term creep rupture of the

reinforcement

RF_D = Strength reduction factor to prevent rupture of the reinforcement due to long-

term chemical and biological degradation

 RF_{ID} = Strength reduction factor to account for installation damage to the reinforcement

 T_{al} = 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} = The ultimate tensile strength of the reinforcement determined from wide width

tensile tests

UV = Ultraviolet light

WSDOT = 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:

- 1. AASHTO Bridge Standard Specifications for Highway Bridges, 17th Edition, 2002
- 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
- 6. ASTM D4595 Standard Test Method for Tensile Properties of Geotextiles by the Wide-Width Strip Method
- 7. ASTM D 6637 Standard Test Method for Determining Tensile Properties of Geogrids by the Single or Multi-Rib Tensile Method.

- 8. ASTM D-1248 Standard Specification for Polyethylene Plastics Extrusion Materials for Wire and Cable
- 9. ASTM D-4101 Standard Specification for Polypropylene Injection and Extrusion Materials
- 10. WSDOT Test Method T 926 Geogrid Brittleness Test
- 11. ISO/DIS 10722-1 Procedure for simulating damage during installation. Part 1: Installation in granular materials
- 12. ASTM D5818 Standard Practice for Obtaining Samples of Geosynthetics from a Test Section for Assessment of Installation Damage
- 13. ASTM D2488 Standard Practice for Description and Identification of Soils (Visual-Manual Procedure)
- 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. AASHTO T96 Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine
- 16. ASTM D6992 Accelerated Tensile Creep and Creep-Rupture of Geosynthetic Materials Based on Time-Temperature Superposition Using the Stepped Isothermal Method
- 17. ASTM D5262 Standard Test Method for Evaluating Unconfined Tension Creep Behavior of Geosynthetics
- 18. *ISO/FDIS 9080:2001* Plastic piping and ducting systems Determination of long-term hydrostatic strength of thermoplastics materials in pipe form by extrapolation.
- 19. ASTM D2837 Standard Test Method for Obtaining Hydrostatic Design Basis for Thermoplastic Pipe Materials
- 20. ASTM D4355 Standard Test Method for Deterioration of Geotextiles from Exposure to Ultraviolet Light and Water (Xenon-Arc Type Apparatus)
- 21. ASTM D4603 Standard Test Method for Determining Inherent Viscosity of Poly(Ethylene Terephthalate) (PET) by Glass Capillary Viscometer
- 22. GRI-GG7 Carboxyl End Group Content of PET Yarns
- 23. GRI-GG8 Determination of the Number Average Molecular Weight of PET Yarns Based on a Relative Viscosity Value
- 24. ASTM D3045 Standard Practice for Heat Aging of Plastics Without Load
- 25. ASTM D 3417-99 Enthaplies of Fusion and Crystallinization of Polymers by DSC
- 26. ENV ISO 13438:1999 Geotextiles and Geotextile-Related Products Screening Test Method for Determining the Resistance to Oxidation
- 27. ASTM D 3895 Standard Test Method for Oxidative-Induction Time of Polyolefins by Differential Scanning Calorimetry
- 28. 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. The receiving laboratory should verify these dimensions upon receipt of the sample(s) using hand measurement techniques. This is especially critical for strength determination based on a single or limited number of ribs in the specimens tested.
 - 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).
 - 1. 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 (RF_{ID}):

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. Also include LA Wear test results for the backfill material used.
- 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 ${\rm RF}_{\rm ID}$ for each sample.

3. Creep Data Requirements (RF $_{CR}$ 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_{CR} 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 are desirable if available.
- h. If RF_{CR} 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.
- 1. 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. RFCR, 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% 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.
- q. For both creep rupture and low strain creep stiffness testing, if single rib, yarn, or narrow width specimens are used, 1,000 hour creep data in accordance with Appendices B and C that demonstrates the single rib, yarn, or narrow width test results are consistent with the results from multi-rib/wide width testing.
- 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:

$$X = \frac{\Delta H}{\Delta H^{\circ}}$$
 (times 100 for %)

where:

 ΔH is the latent heat under the DSC melt curve ΔH° is the latent heat for a 100% crystalline polymer

Sample Latent Heat. AH° **DSC Scan Speed** Homo-Melt Range (°C) **Polymer** Size (mg) (°C/min) (cal/gm) **HDPE** 5 100-145 68.4 10 PP 7.5 45 100-165 10 PET 10 200-245 30 10

Temperature scan should start 10° C below, continue through, and stop 10° C above the melt range. Recommended test parameters are as follows:

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. RFD, 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 reductfion 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_{ID} 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_{ID} 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. If it is determined that the RF_{ID} values obtained for a product line are not correlated with product weight per unit area, undamaged tensile strength, coating weight, or some other product parameter, and the variance of RF_{ID} between any two products in the product line is 0.1 or more, then each product in the product line shall be tested.

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 in accordance with AASHTO (2002, 2004). For ultimate limit state conditions:

$$T_{al} = \frac{T_{ult}}{RF}$$
 where:

$$RF = RF_{ID} \times RF_{CR} \times RF_{D}$$
 (2)

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_l by T_{lot} (see Appendix B). T_l only accounts for extrapolation uncertainty. Furthermore, T_{lot} is specific to the lot of material used for the creep testing. Normalizing by T_{lot} 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_l using T_{ulr} the MARV of the tensile strength for the product, nor is it correct to use T_l directly in the numerator to calculate T_{alr} .

$$RF_{CR} = \frac{T_{lot}}{T_l}
eq \frac{T_{ult}}{T_l}$$
 and $T_{al}
eq \frac{T_l}{RF_{ID} \times RF_D}$

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 Tl 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. The specific application of geosynthetic structure class shall be carried out in accordance with AASHTO (2002, 2004) and other requirements of the approval authority.

3. Environment Aggressiveness

A nonaggressive environment is defined based on soil gradation and particle characteristics, 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. Soil gradation and particle characteristics primarily affect potential high RF_{ID} values, chemical properties affect the potential for high RF_D values, and temperature affects potential for high RF_D and high RF_{CR} values. The aggressiveness of the soil gradation will depend on the distribution, the maximum size, the angularity, and the durability of the soil particles. In general, the more angular the soil, the more uniform its gradation, the greater the maximum particle size, and the more durable the particles, the more aggressive the soil is with regard to potential for installation damage. Installation damage for geosynthetic reinforcement has been approximately correlated to the d_{50} size of the soil, and the d_{50} size can be used as a basis to interpolate to a specific soil gradation using test results at other gradations (Elias, 2000). However, other gradation characteristics may need to be considered to more accurately interpolate to a specific soil gradation and angularity. While installation damage can be evaluated for the anticipated soil gradation and characteristics, it is generally undesirable to use soils and associated installation conditions that result in a RF_{ID} value that is greater than approximately 1.7 due to the likelihood of excessive variability in the results. The decision as to what gradation characteristics are to be considered too aggressive shall be made by the approval authority.

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 all but the most southerly tier of states in the USA, all wall and slope sites are anticipated to have an effective design temperature that is below 30° C.

For most soil conditions in the USA, the environment will likely be chemically 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. However, there are specific geological regions in the USA that are more likely to have chemically aggressive conditions as described in Elias (2000). Examples include salt affected soils in the arid western (especially southwest) regions of the USA, acid-sulphate soils that are commonly found in the Appalachian region of the USA, and calcareous soils commonly found in Florida, Texas, New Mexico, and many western states.

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, and the contract specifications provide soil chemical criteria that are consistent with nonaggressive conditions as described herein, the environment should be considered to be nonaggressive to determine the longterm strength. However, the backfill should be tested prior to use to verify that it is nonaggressive.

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_{a1}
 - The strength reduction factors RF_{ID} , 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.

Polymer Type	Property	Test Method	Criteria to Allow Use of Default RF*
PP and HDPE	UV Oxidation Resistance	ASTM D4355	Min. 70% strength retained after 500 hrs in weatherometer
PET	UV Oxidation Resistance	ASTM D4355	Min. 50% strength retained after 500 hrs in weatherometer if geosynthetic will be buried within one week, 70% if left exposed for more than one week.
PP and HDPE	Thermo- Oxidation Resistance	ENV ISO 13438:1999, Method A (PP) or B (HDPE)	Min. 50% strength retained after 28 days (PP) or 56 days (HDPE)
PET	Hydrolysis Resistance	Inherent Viscosity Method (ASTM D4603 and GRI Test Method GG8), or Determine Directly Using Gel Permeation Chromatography	Min. Number Average Molecular Weight of 25,000
PET	Hydrolysis Resistance	GRI Test Method GG7	Max. Carboxyl End Group Content of 30
All Polymers	Survivability	¹ Weight per Unit Area (ASTM D5261)	¹ Min. 270 g/m ²
All Polymers	% Post-Consumer Recycled Material by Weight	Certification of Materials Used	Maximum of 0%

^{*}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.

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.

 $^{^{1}}$ Alternatively, a default RF_D = 1.3 may be used if product specific installation damage testing is performed and it is determined that RF_{ID} is 1.7 or less, and if the other requirements in Table 1 are met.

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. The receiving laboratory should verify these dimensions upon receipt of the sample(s) using hand measurement techniques. This is especially critical for strength determination based on a single or limited number of ribs in the specimens tested.
- 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).
- 1. MARV for ultimate wide width tensile strength (ASTM D4595 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, a complete assessment of the product or product line in accordance with this Standard Practice instead of just a QA evaluation may be required by the approval authority 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_{ID} , RF_{CR} , and RF_{D} 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_{ID} , RF_{CR} , and RF_{D} 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 product in the product line with the highest RF_{ID} from the initial product acceptance testing using the soil with a d_{50} size which is equal to or larger than a d_{50} size of 4.75 mm, or other d_{50} size as determined by the approval authority, and the aggregate shall have a maximum LA Wear percent loss of 35 percent. The d_{50} size, angularity, and durability of the selected backfill should be consistent with the d_{50} size used for initial product acceptance (preferably, the same material should be used for both the acceptance testing and the quality assurance testing, if possible). 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 using the multi-rib procedure (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 (ASTM D6992) 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, if the product acceptance testing conducted indicates that the creep is log linear at the low strain levels tested, short-term (1,000 second) ramp and hold (R+H) tests as described in ASTM D6992 may be used and extrapolated to 1,000 hours in lieu of 1,000 hour creep tests. A minimum of two R+H tests shall be conducted for one product in the product line at the load level in which 2 percent strain at 1,000 hours was achieved in the product acceptance testing. If the product acceptance testing indicates that the creep is not log linear at the low strain level tested, then a minimum of two full 1,000 hour creep tests must be conducted at that load level. These tests shall be conducted on the same width specimens as was used for the product acceptance creep stiffness testing.

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 RF_D 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 RF_D 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:

$$t_{\alpha/2,n_1+n_2} = (3) \frac{(\overline{P_1} - \overline{P_2}) - \delta}{\sqrt{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}} \sqrt{\frac{n_1 n_2 (n_1 + n_2 - 2)}{(n_1 + n_2)}}$$

where,

 $\begin{array}{lll} \mathbf{t}_{\alpha/2,\mathrm{n}1+\mathrm{n}2\text{-}2} &=& \mathrm{value} \ \mathrm{of} \ \mathrm{the} \ \mathrm{t-distribution} \ \mathrm{for} \ \mathrm{the} \ \mathrm{installation} \ \mathrm{damage} \ \mathrm{samples} \\ P_I &=& \mathrm{the} \ \mathrm{mean} \ \mathrm{of} \ \mathrm{the} \ \mathrm{strength} \ \mathrm{retained} \ \mathrm{after} \ \mathrm{installation} \ \mathrm{damage} \\ && (\mathrm{i.e.,} \ \mathrm{T_{dam}}/\mathrm{T_{lot}}) \ \mathrm{obtained} \ \mathrm{for} \ \mathrm{initial} \ \mathrm{product} \ \mathrm{acceptance} \\ P_2 &=& \mathrm{the} \ \mathrm{mean} \ \mathrm{of} \ \mathrm{the} \ \mathrm{strength} \ \mathrm{retained} \ \mathrm{after} \ \mathrm{installation} \ \mathrm{damage} \\ && (\mathrm{i.e.,} \ \mathrm{T_{dam}}/\mathrm{T_{lot}}) \ \mathrm{obtained} \ \mathrm{for} \ \mathrm{QA} \ \mathrm{testing} \\ \delta &=& \mathrm{the} \ \mathrm{difference} \ \mathrm{in} \ \mathrm{the} \ \mathrm{means} \ \mathrm{for} \ \mathrm{the} \ \mathrm{populations} \ \mathrm{corresponding} \ \mathrm{to} \\ \mathrm{the} \ \mathrm{sample} \ \mathrm{means} \ P_I \ \mathrm{and} \ P_2 \ \mathrm{(assumed} \ \mathrm{equal} \ \mathrm{to} \ \mathrm{zero} \ \mathrm{for} \ \mathrm{this} \ \mathrm{test}) \\ \mathrm{s}_1 &=& \mathrm{the} \ \mathrm{standard} \ \mathrm{deviation} \ \mathrm{corresponding} \ \mathrm{to} \ P_1 \\ \mathrm{s}_2 &=& \mathrm{the} \ \mathrm{standard} \ \mathrm{deviation} \ \mathrm{corresponding} \ \mathrm{to} \ P_1 \\ \mathrm{s}_1 &=& \mathrm{the} \ \mathrm{number} \ \mathrm{of} \ \mathrm{data} \ \mathrm{points} \ \mathrm{corresponding} \ \mathrm{to} \ P_1 \\ \end{array}$

 $t_{\omega/2,n1+n2-2}$ calculated using Equation 3 shall be no greater than t determined from the applicable Student t table (or from the Microsoft EXCEL function TINV(α ,n-2)) at $\alpha=0.05$ and n1+n2-2 degrees of freedom. If this is not true, the difference between P_I and P_2 is determined to be statistically significant, and $P_I>P_2$, two additional samples from the same installation condition shall be tested and P_2 recalculated and statistically compared to P_I . 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} established.

= the number of data points corresponding to P_2

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):

$$(4 \log t_L = \log t_{reg} - \left[t_{\alpha/2, n-2} \sqrt{1 + \frac{1}{n} + \frac{\left(P - \overline{P}\right)^2}{\sum \left(P_i - \overline{P}\right)^2}} \right] \times \sigma$$

and

$$\sigma = \sqrt{\frac{\sum \left[\log t_i - \overline{\log t}\right]^2 - \frac{\left\{\sum \left[\left(P_i - \overline{P}\right)\left(\log t_i - \overline{\log t}\right)\right]\right\}^2}{\sum \left(P_i - \overline{P}\right)^2}}}{\sum \left(P_i - \overline{P}\right)^2}}$$
(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)

 $t_{\alpha/2,n-2}$ = value of the **t** distribution determined from applicable Student **t** table (or from the Microsoft EXCEL function TINV(α ,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

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

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.

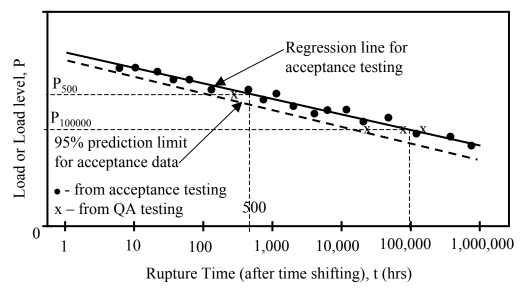
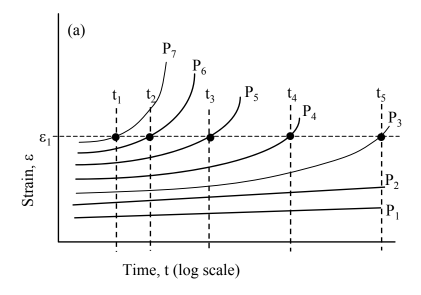


Figure 1. Conceptual illustration of the comparison of QA creep rupture test results to initial product acceptance creep rupture test results.

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.



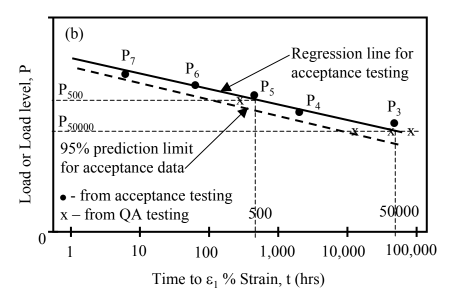


Figure 2. Conceptual illustration of the comparison of QA creep strain test results to initial product acceptance creep strain test results (a) creep strain curves, and (b) envelope of time to the specified strain.

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 curve. The estimated 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 strain 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- \mathbf{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.

 $t_{\alpha/2,n1+n2-2}$ calculated using Equation 3 shall be no greater than t determined from the applicable Student t table (or from the Microsoft EXCEL function TINV(α ,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 OA testing.

References

AASHTO, 2002, *Standard Specifications for Highway Bridges*, American Association of State Highway and Transportation Officials, Seventeenth Edition, Washington, D.C., USA.

AASHTO, 2004, *LRFD Bridge Design Specifications, with Interims*, American Association of State Highway and Transportation Officials, Third Edition, Washington, D.C., USA.

Allen, T.M., 1991, "Determination of Long-Term Strength of Geosynthetics: a State-of-the-Art Review", *Proceedings of Geosynthetics* '91, Atlanta, GA, USA, Vol. 1, pp. 351-379.

Elias, V., 2000, Corrosion/Degradation of Soil Reinforcements for Mechanically Stabilized Earth Walls and Reinforced Soil Slopes, FHWA-NHI-00-044, Federal Highway Administration, Washington, D.C.

Elias, V., DiMaggio, J., and DiMillio, A., 1997, "FHWA Technical Note on the Degradation-Reduction Factors for Geosynthetics," *Geotechnical Fabrics Report*, Vol. 15, No. 6, pp. 24-26.

Federal Highway Administration (FHWA), 1997, "Degradation Reduction Factors for Geosynthetics," *Federal Highway Administration Geotechnology Technical Note*.

Geosynthetic Research Institute, 1998, "Carboxyl End Group Content of Polyethylene Terephthalate (PET) Yarns," GRI Test Method GG7.

Geosynthetic Research Institute, 1998, "Determination of the Number Average Molecular Weight of Polyethylene Terephthalate (PET) Yarns based on a Relative Viscosity Value," GRI Test Method GG8.

Koerner, G. R., Hsuan, Y. G., and Hart, M., 2001, *Field Monitoring and Laboratory Study of Geosynthetics in Reinforcement Applications*, GRI Report #26, Geosynthetic Research Institute, Drexel University, Philadelphia, PA, USA, 116 pp.

Koerner, G. R., Hsuan, Y. G., and Koerner, R. M., 2002, "Field Measurements of Alkalinity (pH) Levels Behind Segmental Retaining Walls, or SRW's," 7th International Geosynthetics Conference, Nice, France, Delmas, Gourc, and Girard, ed's, Vol. 4, pp. 1443-1446.

Koerner, R. M. and Koerner, G. R., 1990, "A quantification and assessment of installation damage to geotextiles." *Proc. Fourth Intl. Conf. on Geotextiles, Geomembranes, and Related Products*, The Hague, pp. 597-602.

Risseeuw, P, and Schmidt, H. M., 1990, "Hydrolysis of HT Polyester Yarns at Moderate Temperatures," *Proceedings of the Fourth International Conference on Geotextiles, Geomembranes, and Related Products*, The Hague, pp. 691-696.

Sprague, C. J., Allen, S., and Thornton, S., 1999, "Installation Damage Testing, Sensitivity Assessment and Derivation of RF_{ID}," Geosynthetics '99, Boston, MA, USA, pp. 1123-1132.

Van Zanten, V., ed., 1986, Geotextiles and Geomembranes in Civil Engineering, Wiley-Halsted.

Wadsworth, H., 1998, *Handbook of Statistical Methods for Engineers and Scientists*, McGraw-Hill, 688 pp.

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 modified 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 shall be large enough to be used for wide width testing (ASTM D4595 for geotextiles or ASTM D6637, specimen preparation method B or C 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. The sample size shall be large enough to obtain the minimum number of specimens (i.e., 20) of the required dimensions.
- 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 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 to a minimum of 90 percent of Modified Proctor per ASTM D1557, or to another compaction standard typically used for geosynthetic reinforced structures if agreed upon in advance by the approval authority.
- 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 as described in Sprague and Allen (2003) 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

- 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_{ID} is then determined as follows:

$$RF_{ID} = \frac{T_{lot}}{T_{down}} \tag{A-1}$$

where, Tlot 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 RFID 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 (d50 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_{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_{ID}.

- If the installation damage test is conducted for a specific project, the actual backfill material planned for use in the geosynthetic structure should be used for the test. If the purpose of the installation damage testing is to generate RF_{ID} values for general use for future projects (i.e., the testing is not intended to be project specific), a range of soil backfill gradations/types should be used in the testing. The range of backfill materials selected should permit interpolation as needed to match the specific soil to be used as reinforced soil backfill. In general, the backfill materials tested should range from soil classified as a sand to coarse gravel (e.g., d₅₀ sizes ranging from 0.5 to 25 mm). The backfill materials selected should be angular to subangular and shall be durable. The coarse sand and gravel portions of the backfill material should have a Los Angeles Wear (LA Wear) percent loss after 500 revolutions (AASHTO T96) of no more than 35 percent. Additional installation damage tests may be conducted with a less durable backfill material, at the discretion of the manufacturer and the approval authority. If tests are conducted using a backfill material that does not meet the LA Wear requirement stated above, the condition of the backfill shall be evaluated for changes in angularity and gradation after each use. If changes in these two parameters are observed, the aggregate shall be immediately replaced with fresh material. This gradation/angularity evaluation should be conducted periodically even for more durable backfill material. Note that if the backfill materials available in the region for which the approval authority has jurisdiction consistently cannot meet the maximum LA Wear requirement of 35 percent loss, a less durable aggregate may be used for all the backfill materials tested, at the discretion of the approval authority.
- 11. Values of RF_{ID} may be estimated for a specified soil gradation using interpolation as illustrated in Figure A-1. The d₅₀ 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 and durability may be considered for this correlation and interpolation procedure per mutual agreement between the geosynthetic manufacturer and the approval authority. The range of backfill gradations, angularity, and durability will affect the range of applicability of the RF_{ID} values obtained from the testing. RF_{ID} values should not be extrapolated beyond the coarsest backfill soil tested.

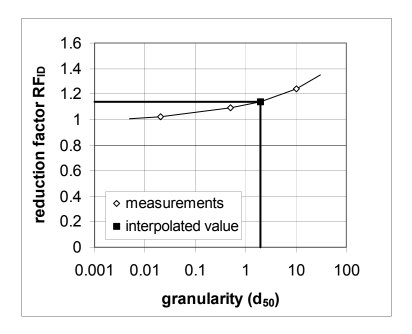
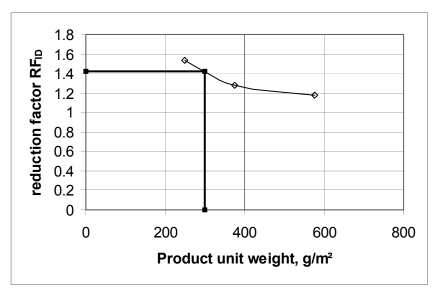


Figure A-1. Interpolation of RF_{ID} for a soil with $d_{50} = 2$ mm from installation damage data obtained for soils with $d_{50} = 0.02$, 0.5 and 10 mm (Note: 1in. = 0.0254 mm).

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.



Figure, A-2. Example interpolation of RF_{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).

If it is determined that the RF_{ID} values obtained for a product line are not correlated with product weight per unit area, undamaged tensile strength, coating weight, or some other product parameter, and the variance of RF_{ID} between any two products in the product line is 0.1 or more, then each product in the product line shall be tested. If in this case (i.e., no correlation could be found), the variance of RF_{ID} between any two products in the product line is less than 0.1 and the upper bound value of RF_{ID} is selected for all products in the product line, all products in the product line do not need to be tested.

References

Sprague, C. J., Allen, S. A., 2003, "Testing Installation Damage of Geosynthetic Reinforcement," Geotechnical Fabrics Report, Vol. 21, No. 6, pp. 24-27.

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 ASTM D6992 (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.

Single ribs for geogrids, or yarns or narrow width specimens for woven geotextiles, may be used for creep testing for the determination of RF_{CR} provided that it can be shown through a limited creep testing program conducted as described in Section B.5 later in this appendix that the rupture behavior and envelope for the single ribs, yarns, or narrow width specimens are the same as that for the full width product, with product width as defined in ASTM D5262. This comparison must demonstrate that there is no statistical difference between the full width product creep rupture regression line and the single rib, yarn, or narrow width specimen 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).

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.

B.1 Overview of Extrapolation Approach to Determine the Ultimate Limit State Creep Limit, T₁

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 stressstrain 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; 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. 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 for a given product, 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).

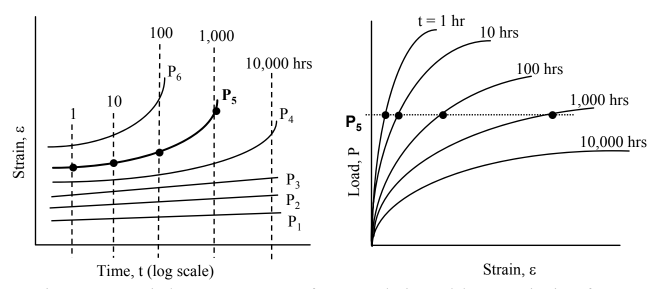


Figure B.2-1. Typical Stress Rupture Data for Geosynthetics, and the Determination of Shift Factors for Time-Temperature Superposition.

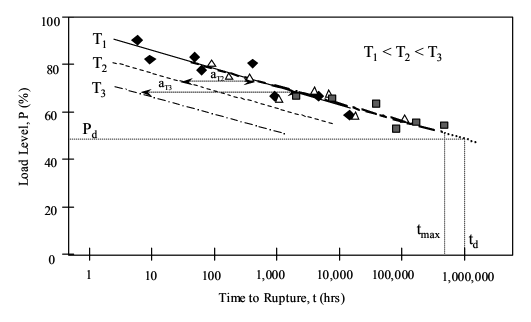


Figure B.2-2. Extrapolation of Stress Rupture Data and the Determination of the Creep Limit Load.

The time to rupture for the elevated temperature rupture data is shifted in accordance with the following equation:

$$t_{amb} = (t_{elev})(a_T) \tag{B.2-1}$$

where, t_{amb} is the predicted time at the ambient or temperature to reach rupture under the specified load, t_{elev} is the measured time at elevated temperature to reach a rupture under the specified load, and $a_{\scriptscriptstyle T}$ is the time shift factor. $a_{\scriptscriptstyle T}$ 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_{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. 1998a; 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° C increase in temperature (i.e., a 10° C increase would result in a time shift factor of 12 to 15) and 0.05 to 0.12 decades per 1° 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 0.05decades per 1° 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° 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 T₁ should be reduced by an extrapolation uncertainty factor as follows:

$$T_1 = P_{cl}/(1.2)^{x-1}$$
 (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 $\log t_d - \log t_{max}$ 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 for the product. 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 ASTM D 6992. 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 (ASTM D6992). At least six SIM rupture tests and six conventional rupture tests and shall be conducted one of the products in 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):

$$\log t_L = \log t_{reg} \pm \left[t_{\alpha, n-2} \sqrt{\frac{1}{n} + \frac{\left(P - \overline{P}\right)^2}{\sum \left(P_i - \overline{P}\right)^2}} \right] \times \sigma$$
and
(B.3-1)

$$\sigma = \sqrt{\frac{\sum \left[\log t_i - \overline{\log t}\right]^2 - \frac{\left[\sum \left[\left(P_i - \overline{P}\right)\left(\log t_i - \overline{\log t}\right)\right]\right]^2}{\sum \left(P_i - \overline{P}\right)^2}}}{\sum \left(P_i - \overline{P}\right)^2}}$$
(B.3-2)

where:

 $\log t_L = \text{lower and upper bound confidence limit. The + or - term in Equation B.2-1 results in the lower and upper bound confidence limits, respectively.}$

 t_{reg} = 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)

 $t_{\alpha,n-2}$ = value of the **t** distribution determined from applicable Student **t** table (or from the Microsoft EXCEL function TINV(α ,n-2)) at α = 0.10 and n-2 degrees of freedom (this corresponds to the 90% two-sided prediction limit).

n = the number of rupture or allowable run-out points in the original test sample (i.e., the conventional creep rupture data)

P = load level obtained at t_{reg} from the regression line developed from the conventional creep rupture testing

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 conventional creep rupture envelope)

 P_i = the rupture load level of the i'th point for the rupture points used in the regression for establishing the conventional creep rupture envelope

 $\log t$ = 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 = the rupture time of the i'th point for the rupture points used in the regression for establishing the conventional creep rupture envelope

Once $\log t_L$, both upper and lower bound, has been determined at the specified load level, compare these values to the log rupture time (i.e., $\log t_{SIM}$) obtained for the SIM creep rupture envelope test at the specified load level (e.g., 1,000 and 50,000 hours). The value of $\log t_{SIM}$ at the two specified load levels must be between the upper and lower bound confidence limits ($\log t_L$). If this requirement is not met, perform two additional SIM tests at each load level P for the specified t_{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 T₁.

B.4 Determination of RF_{CR}

The creep reduction factor, RF_{CR} , 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_{lot}). 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:

$$T_{al} = \frac{P_{95}}{RF_{ID} \times RF_D} \tag{B.4-1}$$

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 RF_{CR} , along with RF_{ID} and RF_{D} , are applied to T_{ult} 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.

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 (i.e., the primary product tested, which is typically a product in the middle of the range of products 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_{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 (i.e., the "primary" product) 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 RF_{CR} as a function of product weight or ultimate tensile strength, so that the RF_{CR} for the other products within the product line can be accurately interpolated. Furthermore, T_{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_{cl} 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_{al} must be the lesser of the determination from Equation 1 and the following determination:

$$T_{al} = \frac{P_{95}}{RF_{ID} \times RF_{D}}$$

where,

P₉₅ = 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

AASHTO, 2002, *Standard Specifications for Highway Bridges*, American Association of State Highway and Transportation Officials, Seventeenth Edition, Washington, D.C., USA, 686 p.

Andrawes, K. Z., McGown, A., and Murray, R. T., 1986, "The Load-Strain-Time Behavior of Geotextiles and Geogrids," *Proceedings of the Third International. Conference on Geotextiles*, Vienna, Austria, pp. 707-712.

Bush, D.I., 1990, "Variation of Long-Term Design Strength of Geosynthetics in Temperatures up to 40°C", *Proceedings of the Fourth International. Conference on Geotextiles, Geomembranes, and Related Products*, The Hague, Netherlands, pp. 673-676.

Chow and Van Laeken 1991, "Viscoelastic Relaxation in oriented Semicrystalline Polymers," *Polymer*, Vol. 32, pp. 1798-1802.

den Hoedt, G., Voskamp, W., van den Heuvel, C. J. M., 1994, "Creep and Time-to-Rupture of Polyester Geogrids at Elevated Temperatures," *Proceedings of the Fifth International. Conference on Geotextiles, Geomembranes, and Related Products*, Singapore, pp. 1125-1130.

Ferry, J. D., 1980, *Viscoelastic Properties of Polymers*, 3rd Edition, John Wiley and Sons, NY.

Findley, W. N., Lai, J. S., Onaran, K., 1976, *Creep and Relaxation of Nonlinear Viscoelastic Materials*, Dover Publications, NY, 369 pp.

Helwany, B., Wu, J. T. H., 1992, "A Generalized Creep Model for Geosynthetics," *International Symposium on Earth Reinforcement Practice*, Japan, pp.

Jewell, R.A. and Greenwood, J.H., 1988, "Long-Term Strength and Safety in Steep Soil Slopes Reinforced by Polymer Materials", *Geotextiles and Geomembranes*, Vol. 7, Nos. 1 and 2, pp. 81-118.

Koerner, R. M., 1990, "Determination of the Long-Term Design Strength of Stiff Geogrids," GRI Standard Practice GG4(a).

Lothspeich, S., and Thornton, S., 2000, "Comparison of Long-term Design Reduction Factors Based on Creep Rupture and Creep Strength Criteria for Several Geosynthetic Reinforcing Materials," *EuroGeo 2*, pp. 341-346.

McGown, A., Paine, N., and DuBois, D.D., 1984, "Use of Geogrid Properties in Limit Equilibrium Analysis", *Proceedings of the Symposium on Polymer Grid Reinforcement in Civil Engineering*, Paper No. 1.4, pp. 1-5.

Murray, R. T., McGown, A., 1988, "Assessment of the Time Dependent Behavior of Geotextiles for Reinforced Soil Applications," *Durability of Geotextiles RILEM*, Chapman and Hall.

Popelar, et al., 1990, "Viscoelastic Material Characterization and Modeling for Polyethylene," *Polymer Engineering and Science*, Vol. 30, no. 10, pp. 577-586.

Popelar, C.H., Kenner, V.H., and Wooster, J.P., 1991, "An Accelerated Method for Establishing the Long Term Performance of Polyethylene Gas Pipe Materials", *Polymer Engineering and Science*, Vol. 31, pp. 1693-1700.

Takaku, A., 1981, "Effect of Drawing on Creep Fracture of Polypropylene Fibers", *Journal of Applied Polymer Science*, Vol. 26, pp. 3565-3573.

Takemura, 1959, Theoretical Consideration of the Effect of Crystallinity on the Viscoelastic Properties of Polyethylene," *Journal of Applied Polymer Science*, Vol. 38, pp. 471-478.

Thornton, J. S., Allen, S. R., and Thomas, R. W., (1997) "Approaches for the Prediction of Long Term Viscoelastic Properties of Geosynthetics from ShortTerm Tests," Geosynthetics '97, IFAI, Long Beach, CA, Vol. 1, pp. 277-291.

Thornton, J. S., Allen, S. R., Thomas, R. W., and Sandri, D., 1998, "The Stepped Isothermal Method for Time-Temperature Superposition and Its Application to Creep Data of Polyester Yarns," *Sixth International Conference on Geotextiles, Geomembranes, and Related Products*, Atlanta, GA, pp.699-706.

Thornton, J. S., and Baker, T. L., 2002, "Comparison of SIM and conventional Methods for Determining Creep-Rupture Behavior of a Polypropylene Geotextile," 7th International Geosynthetics Conference, Nice, France, Vol. 4, pp. 1545-1550.

Thornton, J. S., Paulson, J., and Sandri, D., 1998a, "Conventional and Stepped Isothermal Methods for Characterizing Long-Term Creep Strength of Polyester Geogrids," *Sixth International Conference on Geotextiles, Geomembranes, and Related Products,* Atlanta, GA, pp.691-698.

Wadsworth, H., 1998, *Handbook of Statistical Methods for Engineers and Scientists*, McGraw-Hill, 688 pp.

Wilding, M.A. and Ward, I.M., 1978, "Tensile and Creep Recovery in Ultra-High Modulus Polyethylenes," Polymer, Vol. 19, pp.

Wilding, M.A. and Ward, I.M., 1981, "Creep and Recovery of Ultra High Modulus Polyethylene", *Polymer*, Vol. 22, pp. 870-876.

Wrigley, N. E., Austin, R. A., and Harrison, P. E., 1999, "The Long-Term Strength of Geogrid Reinforcement," Geosynthetics '99, Boston, MA, USA, Vol. 2, pp. 711-721.

Zhurkov, S. N., 1965, "Kenetic Concept of the Strength of Solids," *International Journal of Fracture Mechanics*, Vol. 1, pp. 311-323.

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. Full width specimens should be tested, unless it can be shown through a limited testing program that single rib, yarn, or narrow width specimens can be used without affecting the creep rupture envelope (see beginning of Appendix B), though in the case of creep strain testing and extrapolation, the rupture envelope is defined as the time to reach the instability limit strain (described later in Appendix C) 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 T_l , 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 (T_{amb}) 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. When using creep strain data, creep rupture is assumed to occur, or is at least eminent, when the strain exceeds the instability strain limit (see Section C.1) 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.

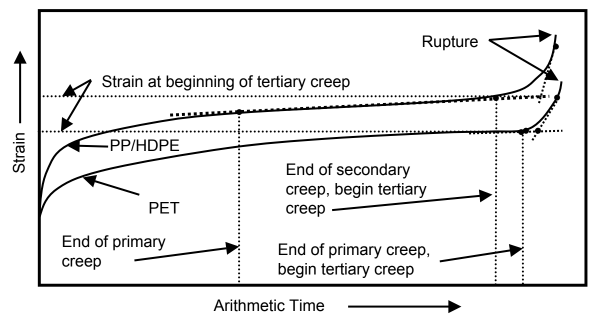
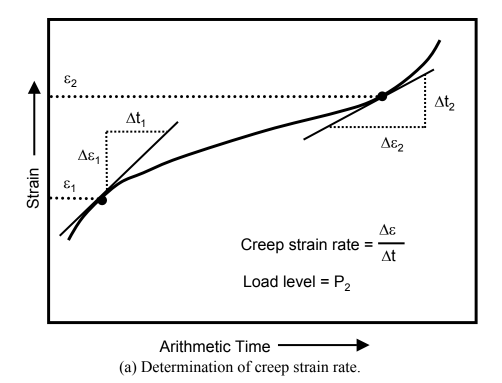


Figure C.1-1. Conceptual illustration of creep strain behavior, and the determination of the strain at the beginning of tertiary creep from creep strain data.

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.



O.001

O.0001

O.0001

O.00001

Figure C.1-2. Development and Use of Sherby-Dorn Plots to Analyze Creep Strain Data.

(b) Interpretation of Sherby-Dorn plots.

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.

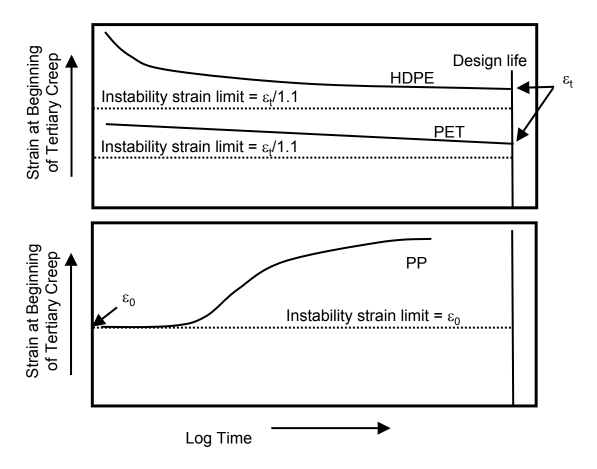


Figure C.1-3. Typical Near Rupture Behavior for Various Geosynthetic Polymers in Terms of Measured Strain.

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.

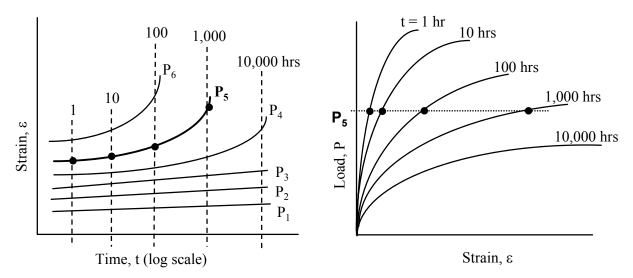
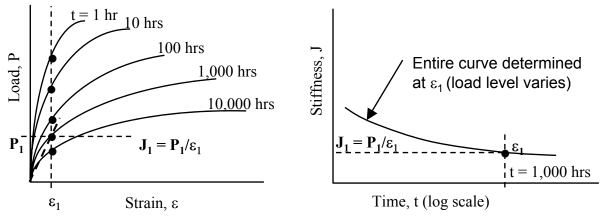


Figure C.1-4. Development of Isochronous Load-Strain Curves.

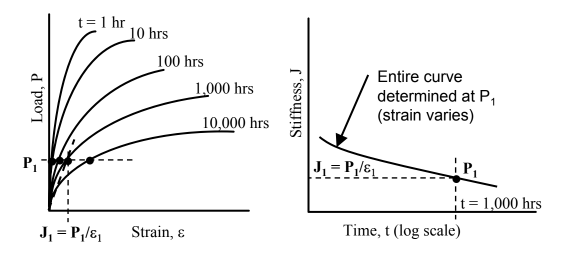
Creep stiffness curves as a function of time are also very useful to use for extrapolation purposes, or to simply estimate the long-term stiffness of the reinforcement for purposes of estimating deformation behavior. Select a constant strain or a constant load level from which to calculate the creep 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:

$$J = P/\varepsilon \tag{C.1-1}$$

where, "P" is the load, and " ϵ " 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_1 . 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.



(a) Creep stiffness determined at constant strain level.



(b) Creep stiffness determined at constant load level.

Figure C.1-5. Development of Creep Stiffness Curves.

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:

$$t_{amb} = (t_{elev})(a_T)$$
 (C.2.1-1)

where, t_{amb} is the predicted time at in-situ temperature to reach a specified creep stiffness or strain under the specified load, t_{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 R² 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.

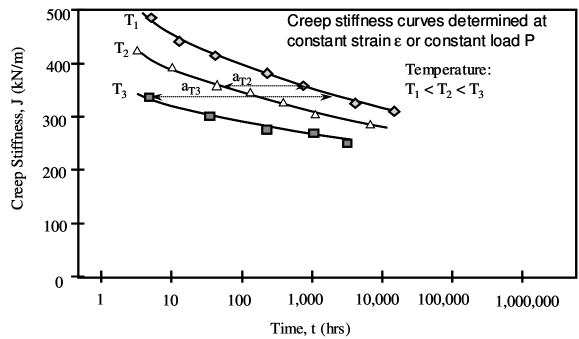
Step 6: Once the creep data has been extrapolated, determine the design, lot specific, creep limit load as follows:

$$T_1 = (J_{c1})(\varepsilon_i) \tag{C.2.1-2}$$

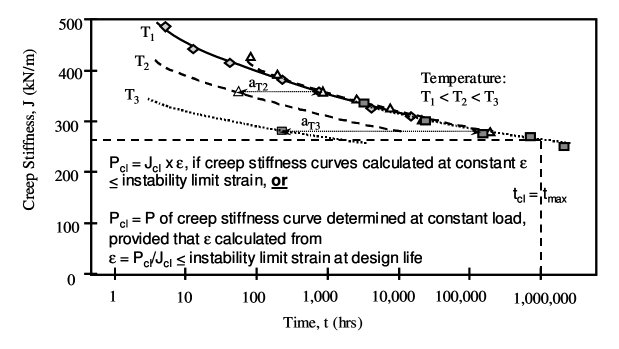
where, J_{cl} is the creep stiffness at the desired design life and temperature, and ϵ_i is the instability limit strain. If statistical extrapolation beyond the time shifted creep stiffness curves, or beyond the actual data if temperature accelerated creep data is not available, is necessary to reach the specified design life, the calculated creep load T_l should be reduced by an extrapolation uncertainty factor as follows:

$$T_1 = (J_{c1})(\varepsilon_i)/(1.2)^{x-1}$$
 (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)x 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.



(a) Creep stiffness curves and determination of shift factors.



(b) Extrapolation of creep stiffness curves and determination of creep limit.

Use of Time-Temperature Superposition to Extrapolate Creep Stiffness Curves. Figure C.2.1-1

C.2.2Step-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 (ASTM D6992). At least six SIM creep tests and six conventional creep tests and shall be conducted on at least one of the products in 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_{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_1 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_{cl} 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_1 , 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:

$$RF_{CR} = \frac{T_{lot}}{T_1} \tag{C.3-1}$$

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 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 overpredict 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).

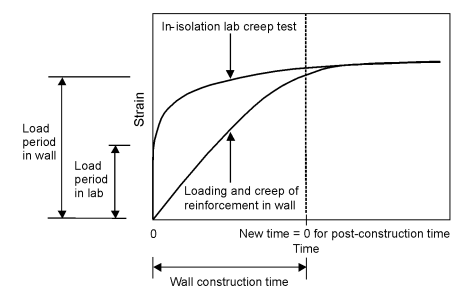


Figure C.4-1. Conceptual illustration showing adjustment of reinforcement creep strains measured in walls to in-isolation laboratory creep data (after Allen and Bathurst, 2002b).

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 at 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, at 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.

The load application rate during creep load ramp up should be consistent with the application rate used in the governing tensile test method (e.g., ASTM D4595 or D6637). If it is not possible to accurately apply the load at a specified rate (e.g., if dead weight is applied through the use of a jack), the actual application rate should be measured and recorded.

If it has been shown that single rib/narrow width specimens can be used in lieu of full width specimens for creep testing at high load levels (see beginning of this appendix), single rib/narrow width specimens may also be used for low strain creep stiffness testing if respective short-term tensile tests also indicate no significant specimen width effects. The maximum difference between the mean values of the load at 2 percent strain in the single rib/narrow width tensile tests and the full width tensile tests must be no greater than what is considered statistically insignificant based on a one sided student-t distribution at a level of

The slack tension, T_o, applied to the specimen based on the governing tensile test (e.g., ASTM D4595 or D6337) will likely be too large for creep stiffness testing due to the very low loads that are likely for this type of testing. A maximum slack tension of approximately 10 percent of the anticipated load at 2 percent strain or 9 N (2 lbf), whichever is less, is recommended for single rib or narrow width specimens. For full width specimens (i.e., per ASTM D5262), a maximum slack tension of approximately 10 percent of the anticipated load at 2 percent strain or 70 N (15 lbf), whichever is less, is recommended. Since these maximum slack tension values differ from what is specified in ASTM D4595 and ASTM D6637, a special set of tensile tests may need to be conducted for use with the low strain creep stiffness testing program. ASTM D4595 and ASTM D6637 allow both the slack tension and the slack displacement, d_o, to be set to zero for calculation purposes. For low strain creep stiffness testing, the slack displacement should be set to zero, but the slack tension should be left at its full value for stiffness calculation purposes.

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). Per agreement between the approval authority and the reinforcement manufacturer, load levels to produce additional 1,000 hour strain levels may be tested. A minimum of two specimens per product at each load level shall be tested.

To establish the load levels needed to produce the desired 1,000 hour strains for each product, conduct a series of 1,000 second ramp and hold (R+H) tests. An initial estimate of the load levels needed for the R+H tests can be obtained from the tensile tests used to establish T_{lot} for each product tested. Test a load level that is likely to produce a strain of approximately 2 percent at 1,000 hours, and then two other load levels to bracket the 2 percent load level (e.g., at load levels that would yield approximately 1 percent strain and 3 to 4 percent strain at 1,000 hours). Do three replicate R+H tests at each of those load levels, plotting the load level as a function of the estimated strain at 1,000 hours, assuming a log linear extrapolation is valid for the R+H test results. Perform a regression analysis of that data to obtain a more accurate estimation of the load level required to produce a strain of 2 percent at 1,000 hours, and run three replicate R+H tests at that load level. If one of the load levels used for the R + H tests does not produce an estimated strain of 2 percent at 1,000 hours, a fourth set of three replicate R+H tests may be needed, estimating the load level based on a regression of the R+H tests for the first three load levels. Then conduct two full 1,000 hour creep tests at the R+H load level that results in the closest estimate to 2 percent strain at 1,000 hours.

Note 1: It is desirable to conduct the full 1,000 hour creep tests at the R+H load levels tested so that a direct comparison can be made between the 1,000 hour creep tests and the R+H tests. If the creep observed in the 1,000 hour creep tests is in fact log linear, the R+H tests can then be used for quality assurance testing in the future.

Step 2: Use the 1,000 hour creep test results, 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, if the 1,000 hour creep stiffness tests do not achieve a strain of 2 percent (or other specified strain) at 1,000 hours..

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, if SIM has been determined to be applicable. In general, extrapolation should not be necessary if the "conventional" approach is used. 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 for determination of RF_{CR} . For creep stiffness assessment, a minimum of three products in the product line spanning the range of products in the line shall be tested as described in Section C.5. To interpolate to other products between the products tested, determine T_{lot} for each product tested, plotting the creep stiffness values obtained in Section C.5 as a function of T_{lot} . Creep stiffness values for other products in the product line not tested can be interpolated based on their tensile strength.

References

Allen, T.M., 1991, "Determination of Long-Term Strength of Geosynthetics: a State-of-the-Art Review", *Proceedings of Geosynthetics* '91, Atlanta, GA, USA, Vol. 1, pp. 351-379.

Allen, T. M., and Bathurst, R. J., 1996, "Combined Allowable Strength Reduction Factor for Geosynthetic Creep and Installation Damage", Geosynthetics International, Vol. 3, No. 3, pp. 407-439.

Allen, T.M., and Bathurst, R.J., 2002a, "Soil Reinforcement Loads in Geosynthetic Reinforced Walls at Working Stress Conditions", *Geosynthetics International*, *Geosynthetics International*, Vol. 9, Nos. 5-6, pp. 525-566.

Allen, T.M., and Bathurst, R.J., 2002b, "Observed Long-Term Performance of Geosynthetic Walls, and Implications for Design", in press with *Geosynthetics International*, *Geosynthetics International*, Vol. 9, Nos. 5-6, pp. 567-606.

Allen, T.M., Bathurst, R.J., Lee, W. F., Holtz, R. D., and Walters, D.L., 2003, "A New Working Stress Method for Prediction of Reinforcement Loads in Geosynthetic Walls", *Canadian Geotechnical Journal*, Vol. 40, No. 5, pp. 976-994.

Andrawes, K. Z., McGown, A., and Murray, R. T., 1986, "The Load-Strain-Time Behavior of Geotextiles and Geogrids," *Proceedings of the Third International. Conference on Geotextiles*, Vienna, Austria, pp. 707-712.

Bush, D.I., 1990, "Variation of Long-Term Design Strength of Geosynthetics in Temperatures up to 40°C", Proceedings of the Fourth International. Conference on Geotextiles, Geomembranes, and Related Products, The Hague, Netherlands, pp. 673-676.

den Hoedt, G., Voskamp, W., van den Heuvel, C. J. M., 1994, "Creep and Time-to-Rupture of Polyester Geogrids at Elevated Temperatures," *Proceedings of the Fifth International. Conference on Geotextiles, Geomembranes, and Related Products*, Singapore, pp. 1125-1130.

Ferry, J. D., 1980, Viscoelastic Properties of Polymers, 3rd Edition, John Wiley and Sons, NY

Ingold, T.S., Montanelli, F., and Rimoldi, P., 1994, "Extrapolation Techniques for Long Term Strengths of polymer Geogrids," *Proceedings of the Fifth International. Conference on Geotextiles, Geomembranes, and Related Products*, Singapore, pp. 1117-1120.

Jewell, R.A. and Greenwood, J.H., 1988, "Long-Term Strength and Safety in Steep Soil Slopes Reinforced by Polymer Materials", *Geotextiles and Geomembranes*, Vol. 7, Nos. 1 and 2, pp. 81-118.

Koerner, R. M., 1990, "Determination of the Long-Term Design Strength of Stiff Geogrids," GRI Standard Practice GG4(a).

Krumm, G., 1988, Der einfluss mechanischer schadigung auf das kriechverhalten von technischem filament - garn aus polyethylenterphthalat, Diplomarbeit TU - Aachen.

McGown, A., Andrawes, K. Z., Yeo, K. C. and DuBois, D.D., 1984, "The Load-Strain-Time Behavior of Tensar Geogrids", *Proceedings of the Symposium on Polymer Grid Reinforcement in Civil Engineering*, Paper No. 1.2, pp. 1-7.

McGown, A., Paine, N., and DuBois, D.D., 1984a, "Use of Geogrid Properties in Limit Equilibrium Analysis", *Proceedings of the Symposium on Polymer Grid Reinforcement in Civil Engineering*, Paper No. 1.4, pp. 1-5.

Popelar, C.H., Kenner, V.H., and Wooster, J.P., 1991, "An Accelerated Method for Establishing the Long Term Performance of Polyethylene Gas Pipe Materials", *Polymer Engineering and Science*, Vol. 31, pp. 1693-1700.

Takaku, A., 1981, "Effect of Drawing on Creep Fracture of Polypropylene Fibers", *Journal of Applied Polymer Science*, Vol. 26, pp. 3565-3573.

Thornton, J. S., Allen, S. R., and Thomas, R. W., 1997, "Approaches for the Prediction of Long Term Viscoelastic Properties of Geosynthetics from Short Term Tests," *Geosynthetics* '97, IFAI, Long Beach, CA, Vol. 1, pp. 277-291.

Thornton, J. S., Allen, S. R., Thomas, R. W., and Sandri, D., 1998, "The Stepped Isothermal Method for Time-Temperature Superposition and Its Application to Creep Data of Polyester Yarns," *Sixth International Conference on Geotextiles, Geomembranes, and Related Products*, Atlanta, GA, pp. 699-706.

Thornton, J. S., Paulson, J., and Sandri, D., 1998, "Conventional and Stepped Isothermal Methods for Characterizing Long-Term Creep Strength of Polyester Geogrids," *Sixth International Conference on Geotextiles, Geomembranes, and Related Products,* Atlanta, GA, pp. 691-698.

WSDOT Test Method No. 925, Appendix D

Product Specific Testing and Data Interpretation Requirements to Determine RFD for Geosynthetic Reinforcements

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 RF_D less than the default minimum of 1.3, or to determine RF_D 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.

The product specific durability studies for the determination of RF_D 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 RF_D at a temperature of 20° C (68° F) as a minimum. Determination of RF_D 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, 2000, and Elias, 2001). 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 (2001) suggests that 30 years of inservice 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 strongly recommended to keep the environment inside the oven as uniform as possible during the entire test duration and to keep oxidation products from building up inside the oven, 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% O₂), 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 Elias, et al. (1999) 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 and reaction with carbon dioxide during the long-term incubation (e.g., replace any air inside the reactor with nitrogen, use de-aired water, keep system well sealed, 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 75° C for polypropylene, high density polyethylene, and polyester, except as discussed above for polyolefin products which have microcracks 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 (2000) 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 tind 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:

$$RF_D = \frac{T_{lot}}{T_D} \tag{D-1}$$

where, T_{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 RF_D 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 RF^D. 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 (2000, 2001) for a more detailed description of the procedures required to obtain the needed information to determine RF_D using this approach.
- The polymer and physical characteristics of the exhumed material must meet the requirements for use in determining RF_D 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, 2001). In most cases, degradation has been minimal, if it is even significant enough be measurable. These laboratory studies appear to corroborate the lack of degradation that has been observed in the field (Elias, 2001). 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 (2001).

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

AASHTO, 2002, *Standard Specifications for Highway Bridges*, American Association of State Highway and Transportation Officials, Seventeenth Edition, Washington, D.C., USA, 686 p.

Allen, T. M., and Elias, V., 1996, *Durability of Geosynthetics for Highway Applications - Interim Report*, FHWA-RD-95-016, Washington, D.C.

Anderson, P. L., Jailloux, J.-M., and White, D. F., 1992, "Testing Durability of Polyester to Be Used in Earth-Reinforced Structures," Earth Reinforcement Practice, Balkema, Rotterdam, pp. 9-12.

American Society for Testing and Materials (ASTM), 1994, *Annual Book of Standards*, Vol. 8.04, Elastic Pipe and Building Materials, ASTM, Philadelphia, PA, USA.

American Society for Testing and Materials (ASTM), 1994, *Annual Book of Standards*, Vol. 4.08 and 4.09, Soil and Rock, ASTM, Philadelphia, PA, USA.

Bright, D. G., 1993, "The Environmental Stress Cracking of Polymers Used in Geosynthetic Products," Geosynthetics '93 Conference, Vancouver, B.C., pp. 925-934.

Elias, V., 1990, Durability/Corrosion of Soil Reinforced Structures, FHWA/RD-89/186, Federal Highway Administration, Washington, D.C.

Elias, V., 2000, *Corrosion/Degradation of Soil Reinforcements for Mechanically Stabilized Earth Walls and Reinforced Soil Slopes*, FHWA-NHI-00-044, Federal Highway Administration, Washington, D.C.

Elias, V., 2001, Long-Term Durability of Geosynthetics Based on Exhumed Samples from Construction Projects, FHWA Report FHWA RD-00-157, 53 pp.

Elias, V., Salman, A., Juran, I., Pearce, E., and Lu, S. 1999, *Testing Protocols for Oxidation and Hydrolysis of Geosynthetics*, FHWA Report FHWA RD-97-144, 200 pp.

Elias, V., DiMaggio, J., and DiMillio, A., 1997, "FHWA Technical Note on the Degradation-Reduction Factors for Geosynthetics," *Geotechnical Fabrics Report*, Vol. 15, No. 6, pp. 24-26.

Federal Highway Administration (FHWA), 1997, "Degradation Reduction Factors for Geosynthetics," *Federal Highway Administration Geotechnology Technical Note*.

Jailloux, J.-M., Anderson, P. L., and Thomas, R., 1992, "Chemical Compatibility Studies of Polyester Fibers and Yarns to Be Used in Geocomposites," Geotextiles and Geomembranes, Vol. 11, No. 3, pp. 277-290.

Jewell, R.A. and Greenwood, J.H., 1988, "Long-Term Strength and Safety in Steep Soil Slopes Reinforced by Polymer Materials", *Geotextiles and Geomembranes*, Vol. 7, Nos. 1 and 2, pp. 81-118.

Lustiger, A., 1983, "The Molecular Mechanism of Slow Crack Growth in Polyethylene," Proceedings, Eighth Plastic Fuel Gas Pipe Symposium, American Gas Association: Arlington, Virginia, pp. 54-56.

McMahon, W., Birdsall, H. A., Johnson, G. R., and Camilli, C. T., 1959, "Degradation Studies of Polyethylene Terephthalate," Journal Chemical Engineering Data, Vol. 4, No. 1, pp. 57-79.

Murray, R. T. and Farrar, D. M., 1988, "Temperature Distributions in Reinforced Soil Retaining Walls," Geotextiles and Geomembranes, No. 7, pp. 33-50.

Salman, A., Elias, V., Juran, I., Lu, S., and Pearce, E., 1997, "Durability of Geosynthetics Based on Accelerated Laboratory Testing," Geosynthetics '97, IFAI, San Diego, CA, pp. 217-234.

Salman, A., Elias, V., and DiMillio, A., 1998, "The Effect of Oxygen Pressure, Temperature, and manufacturing Processes on Laboratory Degradation of Polyolefin Based Geosynthetics," *Sixth International Conference on Geotextiles, Geomembranes, and Related Products*, Atlanta, GA (in press).

Schröder H. F., Bahr H., Lorenz E., Kneip E., Schmücking I., Zeylanov E. B., "Resistance of polyolefin geosynthetics to oxidation – a new accelerated test working at elevated oxygen pressure. Geosynthetics – State of the art – Recent developments," eds Delmas P., and Gourc J P., Proceedings of the Seventh International Conference on Geosynthetics, Nice, France. Balkema, Lisse, Netherlands, 2002, pp 1455-1458.

Shelton, W. S., and Bright, D. G., 1993, "Using the Arrhenius Equation and Rate Expressions, to Predict the Long-Term Behavior of Geosynthetic Polymers," Geosynthetics '93 Conference, Vol. 2, Vancouver, B.C., pp. 789-802.

Van Zanten, V., ed., 1986, Geotextiles and Geomembranes in Civil Engineering, Wiley-Halsted.

Wisse, J.D.M., Broos, C.J.M., and Boels, W.H., 1990, "Evaluation of the Life Expectancy of Polypropylene Geotextiles Used in Bottom Protection Structures Around the Ooster Schelde Storm Surge Barrier - A Case Study", *Proceedings of the Fourth International Conference on Geotextiles, Geomembranes, and Related Products*, The Hague, Netherlands, pp. 679-702.

Wrigley, N. E., 1987, "Durability and Long-Term Performance of Tensar Polymer Grids for Soil Reinforcement," Materials Science and Technology, Vol. 3, pp. 161-170.

Yanful, E. K., 1993, Oxygen Diffusion Through Soil Covers on Sulphidic Mine Tailings," ASCE Journal of Geotechnical Engineering, Vol. 119, No. 8, pp. 1207-1228.

Yanful, E. K., Riley, M. D., Woyshner, M. R., and Duncan, J., 1993, "Construction and Monitoring of a Composite Soil Cover on an Experimental Waste-Rock Pile near Newcastle, New Brunswick, Canada," Canadian Geotechnical Journal, No. 30, pp. 588-599.

WSDOT Test Method T 926

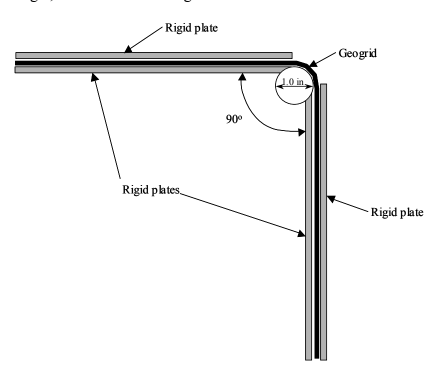
Geogrid Brittleness Test

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.



Geogrid bend test setup. Figure 1

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.

WSDOT Test Method for ASTM C 9391

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

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.

¹ This Test Method is based on ASTM C 939-97.

6. Apparatus

- 6.1 *Flow Cone*. The body and discharge tube can be stainless steel, cast aluminum, or other essentially non-corroding 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 $73.4 \pm 3^{\circ}F$ ($23 \pm 1.7^{\circ}C$), 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

WSDOT FOP for ASTM C 1611

Standard Test Method for Slump Flow of Self-Consolidating Concrete

1. Scope

- 1.1 This test method covers the determination of slump flow of self-consolidating concrete.
- 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 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. (Warning: Fresh hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.)
- 1.4 The text of this standard references notes and footnotes that provide explanatory material. These notes and footnotes (excluding those in tables and figures) shall not be considered as requirements of the standard.

2. Referenced Documents

2.1 ASTM Standards

C 143/C 143M

Test Method for Slump of Hydraulic-Cement Concrete

C 172 Practice for Sampling Freshly Mixed Concrete

C 173/C 173M

Test Method for Air Content of Freshly Mixed Concrete by the Volumetric Method

C 670 Practice for Preparing Precision and Bias Statements for Test Methods for Construction Materials

2.2 AASHTO Standards

T 119M/T 119

Standard Test Method for Slump of Hydraulic-Cement Concrete

T 347 Slump Flow of Self-Consolidating Concrete (SCC)

2.3 WAQTC Standards

TM 2 Sampling Freshly Mixed Concrete

3. Terminology

- 3.1 Definitions of terms specific to this standard:
 - 3.1.1 *halo,* n An observed cement paste or mortar ring that has clearly separated from the coarse aggregate, around the outside circumference of concrete after flowing from the slump cone.
 - 3.1.2 *spread,* n The distance of lateral flow of concrete during the slump-flow test.
 - 3.1.3 *stability,* n The ability of a concrete mixture to resist segregation of the paste from the aggregates.
 - 3.1.4 *viscosity,* n Resistance of a material to flow under an applied shearing stress.

4. Summary of Test Method

4.1 A sample of freshly mixed concrete is placed in a mold shaped as the frustum of a cone. The concrete is placed in one lift without tamping or vibration. The mold is raised, and the concrete allowed to spread. After spreading ceases, two diameters of the concrete mass are measured in approximately orthogonal directions, and slump flow is the average of the two diameters.

5. Significance and Use

- 5.1 This test method provides a procedure to determine the slump flow of self-consolidating concrete in the laboratory or the field.
- 5.2 This test method is used to monitor the consistency of fresh, unhardened self-consolidating concrete and its unconfined flow potential.
- 5.3 It is difficult to produce self-consolidating concrete that is both flowable and nonsegregating using coarse aggregates larger than 1 in (25 mm). Therefore, this test method is considered applicable to self-consolidating concrete having coarse aggregate up to 1 in (25 mm) in size.

6. Apparatus

- 6.1 Mold The mold used in this test method shall conform to that described in FOP for AASHTO T 119.
- Base Plate The base plate on which the mold rests shall be nonabsorbent, smooth, rigid, and have a minimum diameter of 36 in (915 mm).
 - **Note 1:** Field experience and results from the round robin test program have shown that base plates made from sealed/laminated plywood, acrylic plastic, or steel are suitable for performing this test.
- 6.3 Strike-off Bar As described in FOP for WAQTC T 152.

7. Sample

7.1 The sample of concrete from which test specimens are made shall be representative of the entire batch. Sample in accordance with FOP for WAQTC TM 2.

8. Procedure

- 8.1 The slump-flow test shall be performed on a flat, level, nonabsorbent base plate. Position and shim the base plate so it is fully supported, flat, and level.
- 8.2 Filling the Mold WSDOT requires the use of Procedure B.
 - 8.2.1 Filling Procedure B (Inverted Mold) Dampen and place the mold, with the smaller opening of the mold facing down, in the center of a flat, moistened base plate or concrete surface. Using a suitable container, fill the entire mold continuously (Note 2). The mold shall be held firmly in place during filling. Do not rod or tamp the SCC. Slightly overfill the mold.

Note 2: Filling the mold with concrete by using multiple scoops or by pouring from a bucket or similar container has been found to be acceptable.

- 8.3 Strike off the surface of the concrete level with the top of the mold by a sawing motion of the strike-off bar. Remove concrete from the area surrounding the base of the mold to preclude interference with the movement of the flowing concrete. Remove the mold from the concrete by raising it vertically. Raise the mold a distance of 9 ± 3 in $(225 \pm 75 \text{ mm})$ in 3 ± 1 seconds by a steady upward lift with no lateral or torsional motion. Complete the entire test from start of the filling through removal of the mold without interruption within an elapsed time of $2\frac{1}{2}$ minutes.
- 8.4 Wait for the concrete to stop flowing and then measure the largest diameter of the resulting circular spread of concrete to the nearest ¼ in (5 mm). When a halo is observed in the resulting circular spread of concrete, it shall be included as part of the diameter of the concrete. Measure a second diameter of the circular spread at an angle approximately perpendicular to the original measured diameter.
- 8.5 If the measurement of the two diameters differs by more than 2 in (50 mm), the test is invalid and shall be repeated.

9. Calculation

9.1 Calculate the slump flow using Eq 1:

Slump flow =
$$\frac{(d^1 + d^2)}{2}$$

where:

 d^1 = the largest diameter of the circular spread of the concrete, and

 d^2 = the circular spread of the concrete at an angle approximately perpendicular to d^1

9.2 Record the average of the two diameters to the nearest ½ in (5 mm).

10. Report

- 10.1 Report the slump flow to the nearest ¼ in (5 mm).
- 10.2 Report results on concrete delivery ticket (i.e., Certificate of Compliance).
- 10.3 The name of the tester who performed the field acceptance test is required on concrete delivery tickets containing test results.

11. Precision and Bias

See ASTM C1611/C 1611M for precision and bias.

Performance Exam Checklist WSDOT FOP for ASTM C 1611/C 1611M Standard Test Method for Slump Flow of Self-Consolidating Concrete

Participant Name		Exam D	ate			
	edure Element	y of the current procedure	on hand?		Yes	No
2. A	All equipment is fun urrent calibration/ve	actioning according to the erification tags present?	test procedure, and if	f required, has the	2	
6. N	Mold struck off level	l with top opening?				
	Excess material removed from base plate and mold raised 9 ± 3 inches, in 3 ± 1 seconds?					
8. A	After flow stabilized, measured largest diameter (including halo if necessary)?					
9. S	9. Second measurement taken approximately perpendicular to first measurement?					
10 First and second measurements agree within 2"?						
11. S	11. Slump flow was reported as an average of the two measurements?					
12. S	Slump flow reported	to the nearest $\frac{1}{4}$ "?				
	Attempt: Pass	Fail	Second Attempt:	Pass Fail		
Signat	ture of Examiner .					
Comm	ients:					

WSDOT FOP for ASTM C 1621/C 1621M1

Standard Test Method for Passing Ability of Self-Consolidating Concrete by J-Ring

1. Scope

- 1.1 This test method covers determination of the passing ability of self-consolidating concrete by using the J-Ring in combination with a slump cone mold. The test method is limited to concrete with maximum size of aggregate of 1 in (25 mm).
- 1.2 The values stated in either inch-pounds 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 this standard references notes that provide explanatory material. These notes (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 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. (*Warning:* Fresh hydraulic cementitious mixtures are caustic and may cause chemical burns to skin and tissue upon prolonged exposure.)

2. Referenced Documents

2.1 ASTM Standards

C 125 Terminology Relating to Concrete and Concrete Aggregates

C 143/C 143M

Test Method for Slump of Hydraulic-Cement Concrete

C 172 Practice for Sampling Freshly Mixed Concrete

C 173/C 173M

Test Method for Air Content of Freshly Mixed Concrete by the Volumetric Method

C 1611/C 1611M

Test Method for Slump Flow of Self-Consolidating Concrete

3. Terminology

- 3.1 Definitions
 - 3.1.1 For definitions of terms used in this test method, refer to Terminology C 125.
- 3.2 Definitions of terms specific to this standard:
 - 3.2.1 *Halo* An observed cement paste or mortar ring that has clearly separated from the coarse aggregate, around the outside circumference of concrete after flowing from the slump cone.

¹This Test Method is based on ASTM C 1621/C 1621M and has been modified per WSDOT standards. To view the redline modifications, contact the WSDOT Quality Systems Manager at 360-709-5412.

- 3.2.2 *J-ring* An apparatus consisting of a rigid ring supported on sixteen % in (16 mm) diameter rods equally spaced on a 12 in (300 mm) diameter circle 4 in (100 mm) above a flat surface as shown in Figure 1.
- 3.2.3 *J-ring flow* The distance of lateral flow of concrete using the J-Ring in combination with a slump cone.
- 3.2.4 *Passing ability* The ability of self-consolidating concrete to flow under its own weight (without vibration) and fill completely all spaces within intricate formwork, containing obstacles, such as reinforcement.

4. Summary of Test Method

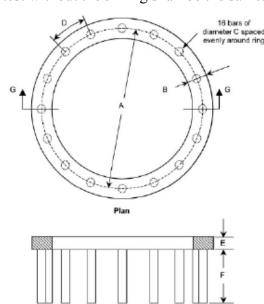
4.1 A sample of freshly mixed concrete is placed in a slump mold (inverted position) that is concentric with the J-Ring (Figure 2). The concrete is placed in one lift without tamping or vibration. The mold is raised, and the concrete is allowed to pass through J-Ring and subside (Figure 3).

The diameters of the concrete, in two directions approximately perpendicular to each other, are measured and averaged to obtain the J-Ring flow. The test is repeated without the J-Ring to obtain the slump flow.

The difference between the slump flow and J-Ring flow is an indicator of the passing ability of the concrete.

5. Significance and Use

5.1 This test method provides a procedure to determine the passing ability of self-consolidating concrete mixtures. The difference between the slump flow and J-Ring flow is an indication of the passing ability of the concrete. A difference less than 1 in (25 mm) indicates good passing ability and a difference greater than 2 in (50 mm) indicates poor passing ability. The orientation of the slump cone for the J-Ring test and for the slump flow test without the J-Ring shall be the same.



Section G-G

Dimension	in	mm	
Α	12.0 ± 0.13	300 ± 3.3	
В	1.5 ± 0.06	38 ± 1.5	
С	0.625 ± 0.13	16 ± 3.3	
D	2.36 ± 0.06	58.9 ± 1.5	
Е	1.0 ± 0.06	25 ± 1.5	
F	4.0 ± 0.06	200 ± 1.5	

Figure 1

5.2 This test method is applicable for laboratory use in comparing the passing ability of different concrete mixtures. It is also applicable in the field as a quality control test.

6. Apparatus

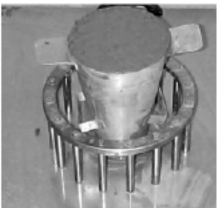
- 6.1 J-Ring The apparatus shall consist of a steel (or equivalent nonabsorbent, rigid material) ring measuring 12 in (300 mm) in diameter at the center of the ring and 1 in (25 mm) in thickness, and sixteen % in (16 mm) diameter smooth steel rods spaced evenly around the ring measuring 4 in (100 mm) in length (see Figure 1).
- 6.2 Mold The mold (slump cone) used in this test method is as described in FOP for AASHTO T 119.
- 6.3 Base Plate A nonabsorbent, rigid plate having a diameter of at least 36 in (915 mm). *Note 1:* Field experience has shown that base plates made from sealed or laminated plywood, rigid plastic, or steel are suitable for performing this test.
- 6.4 Strike Off Bar As described in FOP for WAQTC T 152.
- 6.5 Measuring Device A ruler, metal roll-up measuring tape, or similar rigid or semi-rigid length measuring instrument marked in increments of ¼ in (5 mm) or less.

7. Sample

7.1 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.

8. Procedure

8.1 Perform the test on a flat, level, and nonabsorbent base plate. Position and shim the base plate so that it is fully supported and level. Pre-moisten base-plate with a damp towel, rag, or sponge. Rest the J-Ring at the center of the base plate.



J-Ring Setup with Inverted Mold Filled with Concrete Figure 2

Ring Flow Figure 3

- 8.2 WSDOT uses only Procedure B.
 - 8.1.2 Filling Procedure B (Inverted Mold) Dampen the mold, and place it on the base plate with the smaller opening facing down and concentric with the J-Ring. Support the mold and fill the mold in one lift (Note 2). Heap the concrete above the top of the mold.
 - *Note 2:* Filling the mold with concrete by using multiple scoops or by pouring from a bucket or similar container has been found to be acceptable.

- 8.3 Strike off the surface of the concrete level with the top of the mold by a sawing motion of the strike off bar. Remove concrete from the area surrounding the mold to preclude interference with the movement of the flowing concrete. Raise the mold a distance of 9 ± 3 in $(230 \pm 75 \text{ mm})$ in 3 ± 1 s by a steady vertical lift with no lateral or torsional motion. Complete the entire procedure from start of the filling through removal of the mold without interruption within an elapsed time of $2\frac{1}{2}$ min.
- 8.4 Wait for the concrete to stop flowing and then measure the largest diameter (d1) of the resulting circular flow of concrete. When a halo is observed in the resulting circular flow of concrete, it shall be included as part of the diameter of the concrete. Measure a second diameter (d2) of the circular flow at approximately perpendicular to the first measured diameter (d1). Measure the diameters to the nearest $\frac{1}{4}$ in (5 mm). Determine the J-Ring flow in accordance with Section 9 of this test method.
- 8.5 Conduct a slump flow test without the J-Ring in accordance with Test Method C 1611/C 1611M. Use the same filling procedure as used with the J-Ring. Complete the tests with and without the J-Ring within 6 min.

9. Calculation

9.1 Calculate J-Ring flow according to the following equation:

J-Ring flow =
$$\frac{d^1 + d^2}{2}$$

9.2 Calculate the slump flow according to the following equation:

Slump flow =
$$\frac{d^1 + d^2}{2}$$

9.3 Calculate the difference between slump flow and J-Ring flow to the nearest ½ in (10 mm). This number represents the passing ability of the concrete.

10. Blocking Assessment

10.1 Identify blocking assessment according to Table 1.

Difference Between Slump Flow and J-Ring Flow	Blocking Assessment
0 to 1 in (0 to 25 mm)	No visible blocking
> 1 to 2 in (>25 to 50 mm)	Minimal to noticeable blocking
> 2 in (>50 mm)	Noticeable to extreme blocking

Blocking Assessment Table 1

11. Report

- 11.1 Report the filling procedure (A or B) that was used.
- 11.2 Report the J-Ring flow as the average of the two measured diameters to the nearest $\frac{1}{2}$ in (10 mm).
- Report the slump flow (without the J-Ring) as the average of the two measured diameters to the nearest ½ in (10 mm).
- Report the passing ability as the difference between the slump flow and J-Ring flow to the nearest ½ in (10 mm). Identify the blocking assessment.
- 11.5 Report results on concrete delivery ticket (i.e., Certificate of Compliance).
- 11.6 The name of the tester who performed the field acceptance test is required on concrete delivery tickets containing test results.

12. Precision and Bias

See ASTM C 1621/C 1621M for precision and bias.

Performance Exam Checklist WSDOT FOP for ASTM C 1621/C 1621M Standard Test Method for Passing Ability of Self-Consolidating Concrete by J-Ring

Participant Name	Exam Date				
Procedure Element	Yes N	0			
1. The tester has a copy of the current procedure	e on hand?				
2. All equipment is functioning according to the has the current calibration/verification tags pr	resent?				
3. Sample was taken per WSDOT FOP for WAG					
4. Molds and base plate dampened and base pla	te is flat, level and fully supported?				
5. Mold is centered in J-Ring and centered on b	ase plate?				
6. Mold filled completely in one lift (slightly ov	Mold filled completely in one lift (slightly overfilled)?				
7. Mold struck off level with top opening?					
8. Excess material removed from base plate and 3 ± 1 seconds?	I mold raised 9 ± 3 inches, in				
9. After flow has stabilized, measure largest dia	meter (including halo)?				
10. Second measurement taken approximately pe	Second measurement taken approximately perpendicular to first measurement?				
11. Measurements made to nearest 1/4"?					
12. Test performed within 6 minutes of FOP for A	Test performed within 6 minutes of FOP for ASTM C 1611?				
13. All calculations performed correctly?	All calculations performed correctly?				
14. Results reported to the nearest ½"?					
First Attempt: Pass Fail	Second Attempt: Pass Fail				
Signature of Examiner					
Comments:					

Performance Exam Checklist

Test for High and Low Temperature Recovery of Elastomeric Joint Seals for Concrete Pavements ASTM D 2628 AASHTO M220

Par	ticipant Nan	ne		Exam Date		
Pro	cedure Ele	ment			Yes	No
1.	Sample is	cut into 5	-inch pieces?			
2.	Specimens with talc?	for high	and low temp	erature recovery tests (2 each) lightly dusted		
3.			l between para e clamp asser	allel plates to 50% of the nominal width usin mbly?	g	
Hig	h Temperat	ture Reco	very			
1.	capable of	maintaini	$\sin 212 \pm 2^{\circ}F$	ompressed specimens placed in an oven $(100 \pm 1^{\circ}\text{C})$ and kept there for 70 hours?		
2.	Clamp asso	embly ren	noved from th	ne oven after 70 hours?		
3.	Unclampe	d the asse	mbly and care	efully removed the specimens?		
4.	Specimens	allowed	to rest on a w	ooden surface at room temperature for 1 hou	r?	
5.			-	s measured in the center of the 5-inch length using a dial caliper or other measuring device		
6.	Data entere		e computer da	ta base where % Recovery is automatically		
Lov	v Temperati		•			
1.				compressed specimens placed in a refrigerated $20 \pm 2^{\circ}F$ (-29 \pm 1°C) and kept there for 22 he		
2.	Clamp assembly removed from the chamber after 22 hours?					
3.	Unclampe	d the asse	mbly and care	efully removed the specimens?		
4.	Specimens recover for		ed to a woode	en surface in the chamber and allowed to		
5.	Specimens removed from the chamber and measured the recovered width in the center of the 5-inch length at the top of the longitudinal edge using a dial caliper or other measuring device?					
6.	Data entero		e computer da	ta base where % Recovery is automatically		
			Recovery, %	= recovered width × 100/nominal width		
Firs	st Attempt:	Pass	Fail	Second Attempt: Pass	Fail	

Signature of Examiner

Comments:

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 6931

Standard Test Method for Indirect Tensile (IDT) Strength of Bituminous Mixtures

ASTM D 6931 has been adopted by WSDOT with the following changes:

6. Specimens

- 6.1 Laboratory-Molded Specimens Prepare the 150 mm (5.9 in) laboratory-molded specimens in accordance with WSDOT FOP for AASHTO T 312, to a height of 62 ± 1.0 mm (2.44 \pm 0.04 in). A minimum of three replicates shall be prepared for each mixture.
 - 6.1.1 Air void (Va) of test specimen shall be $7.0 \pm 1.0 \%$.

7. Procedure

7.1 Section 7.1 shall be deleted in its entirety.

8. Calculation

8.1 Calculate the IDT strength as follows:

$$S_{T} = \frac{2F}{3.14 \text{ (hd)}}$$

Where:

 S_T = Indirect tensile strength (psi)

F = Total applied vertical load at failure (lbs)

h = Height of specimen (inches)

d = Diameter of specimen (inches)

Tester Qualification Practical Exam Checklist

Determining Indirect Tensile Strength of Compacted Bituminous Mixtures FOP for ASTM D 6931

Part	icipant Name Exam Date				
Pro -	cedure Element 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?				
3.	Specimen height is 62 ± 1.0 mm (2.44 ± 0.04 in) or 38.1 mm (1.5 in) minimum for cores?				
4.	Specimen meets air void tolerance of $7.0 + 1.0 \%$?				
5.	Specimen placed in water bath at $77 + 2^{\circ}F$ (25 + 1°C) for a minimum of 30 minutes but not longer than 120 minutes?				
6.	Press turned on and operating at a deformation rate of 2 in per minute?				
7.	Specimen placed on lower loading strip?				
8.	Upper loading strip lowered onto specimen with light contact?				
9.	Upper and lower loading strips parallel with each other?				
10.	Load applied at 2 in per minute?				
11.	Total applied vertical load recorded?				
12.	Indirect tensile strength in psi calculated and recorded correctly?				
Firs	t Attempt: Pass Fail Second Attempt: Pass Fail				
Sign	nature of Examiner				

Comments:

WSDOT FOP for ASTM D 7012

Standard Test Method for Unconfined Compressive Strength of Intact Rock Core Specimens

ASTM D 7012 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.

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 $1\frac{7}{8}$ inch (48 mm).

Performance Exam Checklist

Nondestructive Measurement of Thickness of Nonmagnetic Coatings on a Ferrous Base FOP for ASTM D 7091/SSPC Procedure 2 (Field)

Participant Name	Exam Date
Procedure Element	Yes No
1. The tester has a copy of the current procedure on ha	and?
2. All equipment is functioning according to the test p current calibration/verification tags present?	procedure, and if required, has the
3. Instrument calibrated in accordance with the manufuse employing a suitable thickness standard?	facturer's instructions before
4. Several readings taken and recorded taking into acc	count edge and curvature effects?
5. (Laboratory) The average thickness converted to oz conversion factor?	z . ft^2 (g/m ²) using appropriate
6. (Field) Average thickness reported in mils?	
First Attempt: Pass Fail Sec	cond Attempt: Pass Fail
Signature of Examiner	

Comments:

A	В
Abrasion resistance	Bark mulchT 123
Abrasion test:	Beams, concreteT 177, T 802, T 808
Los Angeles T 96	Bending beam rheometer T 313
Absorption test:	Bitumen content T 308, D 4
coarse aggregateT 85, T 96	Bituminous materials:
fine aggregate T 84	ash content T 111
Acidity, alkalinity, pHT 200	bending beam rheometerT 313
Aggregate: (see also coarse or fine aggregate)	distillation:
degradationT 113	cut-back asphalt T 78
effect on mortar strengthT 71	ductility T 51, T 113
extracted, mechanical analysisT 30	dynamic shear T 315
fine aggregate angularityT 304	emulsions
flat and elongated particlesD 4791	flash and fire points, open cup T 48
fracture T 335	flash point:
friable particles	tagliabueT 79
lightweight pieces T 113	flexural creepT 313
material finer than 0.075 mm sieve	float testT 50
moisture contentT 255	inorganic matter T 111
organic impuritiesT 21	loss on heating
preparation for HMA mix designT 724	melting joint and crack sealant
reducing to testing sizeT 248	penetrationT 49
sampling T 2, D 75	percentage in mixtures T 308
sieve analysis T 27	pressure agingR 28
splittingT 248	rheological propertiesT 313, T 315
unit weightT 19/T 19M	sampling T 40
Aggregate degradationT 96, T 113	sofening pointT 53
Aging, pressure R 28	soluable in organic solvents T 44
Air content of:	specific gravity T 228, T 229, D 1298
concrete, gravimetricT 121	supplier certificationQC 2
portland cement mortar T 137	viscosity:
pressure T 152	absolute
self-compacting concreteT 818	kinematicT 201
volumetric T 196	rotational
Air permeability fineness T 153	sayboltT 72
Alkali silica reaction	Bituminous roofing membranes T 53
Aqueous solutions T 200	Bituminous mixtures:
Ash, paper	agingR 30
Asphalt: (see bituminous materials)	air voids
Asphalt content	asphalt binder content T 308
Asphalt roofing materialsT 228, T 229	coating and strippingT 718
specific gravityT 228, T 229	compactionT 312
,	coresT 166, T 720, SOP 730

density:		fineness:	
bulk specific gravity-compacted	T 166	air permeability	T 153
cores	SOP 730	grout flow cone	
maximum theoretical	T 209	latex compatibility	T 313
nuclearTM	8, SOP 729	normal consistency	T 129
paraffin coated		soundness	T 107
ramdom testing	T 716	specific gravity	T 133
effect of water	T 718	supplier certification	QC 1
extraction	T 164	time of setting:	
ignition furnace	T 308	gilore needles	T 154
ignition furnace calbration	SOP 728	vicat needle	
gyratory compaction	T 312	Chemical analysis of hydraulic cement.	T 105
long-term aging	R 30	Chloride permeability of concrete	
maximum specific gravity	T 209	Chloride, water soluble	C 1218
mix design:		Classification (standards)T 59	e), D 2487
aggregate preparation	T 724	Clay gel absorption	D 2007
mixing	T 726	Clay lumps in aggregate	T 112
moisture content	T 329	Cleveland open cup	T 48
moisture induced damage		Coarse aggregate:	
nuclear density TM	8, SOP 729	clay lumps	T 112
preparation for HMA mix design		flat and elongated	
random Sampling		fracture	
recovery of asphalt	T 170	lightweight particles	T 113
reducing to testing size	T 712	Los Angeles abrasion test	T 96
sampling	T 168	material finer than 0.075 mm sieve	
short-term aging		sampling	
specific gravity- compacted HMA		sieve analysisT	
splitting		specific gravity and absorption	
stripping of		unit weight	
superpave mix design		Coating of bituminous aggregate mixtur	
superpave volumertics		paraffin for specific gravity test	
voids		Coatings	
Bond strength, epoxy		Coating, ferrous base	
Bond test joint sealants		Cohesive soil, compressive strength	
Burst test	D 3786	Cohesive soils	
•		Compacted hot mix asphaltTM	
U		Compaction and density of soilT	9, T 180
Calcium carbide gas pressure moistur	e tester	T 606	
T 217		Compaction of bituminous mixtures	
Calibration	SOP 728	Compost	T 420
Capillary rise of soils	T 610	Compressive strength:	
California profilograph		cohesive soil	T 208
Cement, hydraulic:		concrete:	
air content of mortar	T 137	molded specimens	
chemical analysis	T 105	specimens taken from hardened con-	
chloride ion		epoxy resins	
compressive strength		fine aggregate	T 71

hydraulic cementT 106, T 813	Cone penetration of lubricating grease T 217
unconfirmed cohesive:	Conflict monitor test
rock	Consistency T 49
soilT 208	Consistency, normal, of hydraulic cement T 59
Concrete	T 129
air content:	Consolidation T 297
gravimetricT 121	Consolidation properties of soils T 216
pressure T 152	Constant head testing T 215
self-compactingT 818	Core:
volumetricT 196	concreteT 810
alkali-silica reactionT 303	HMA T 166, T 720, SOP 730, SOP 734
beamsT 177, T 802, T 808	Cutback asphalts T 78
chloride ion, total T 260	Cyclic segregationSOP 733
chloride ion, water soluble	Cylinders, compressive strength of concrete T 22
chloride permeability T 277	Curing materials T 814
consistencyT 119	_
cores	D
curing materialsT 814	Deformation T 297
densityT 810	DegradationT 113, T 131
flexural strengths:	Delivery
center point loadingT 177	Demulsibility
speciment taken from hardened concrete T 24	Density. TM 8, T 228, T 229, T 310, SOP 615,
third point loadingT 97, T 177, T 802	SOP 72
joint sealers, hot poured D 5329, T 48, T 412	Density and compaction of soilT 99, T 180
joint sealers, two component rubber T 412	T 606
light weight C 495	Density:
molded specimens T 22	cores
preparation of specimens:	HMATM 8
cappingT 231	randon numbers T 716
in the field T 23	Density of paint
in the laboratory T 126	Density of paint Density of soil in-place:
self compacting T 819	nuclear soilsT 310, SOP 615
rebound hammer determination	Determination of bitumen
slump T 119	Directional reselectance T 314
sampling T 141, TM 2	Displacement method
self compacting:	Distillate fuels
air contentT 818	Distillation (general) T 55, T 78
cylinder T 819	Distillation:
splitting tensile strengthT 198	cut-back asphalt T 78
temperature T 310	emulsions T 59
time of setting T 197	
weight per cubic foot T 121	Disturbed soil samples, preparation:
yieldT 121	dry method
self consolidating concrete	Drainage T 297
J-Ring	Ductility of bituminous materials T 51, T 113
slump flow C 1611	Dynamic shear T 315

E	Float test, bituminous materials	T 50
Elastic recoveryCAL 331, CAL 332	Flow, hot melt joint sealant	D 5329
Electrical measurements	Fracture	T 335
Electrodes T 200	Freezing and thawing:	
Elongation T 51	bituminous mixture	T 718
Elutriation test of aggregates T 11	Friable particles	T 112
Embankment nuclear density T 310, SOP 615	Fuel oils	T 72
Emulsified asphalts, test of T 570, SOT 013	Furol viscosity	T 72
Environmental chamber test		
Epoxy resins:	G	
bond strength	Galvanizing:	
compressive strength	embrittlement	A 143
Extracted aggregates, mechanical analysis T 30	weight of coating	
Extracted aggregates, internament analysis 1 30	Geotextiles:	1 00
F	apparent opening size	D 4571
	burst test	
Falling weight deflections	conditioning	
Family of curves	deterioration from ultraviolet light	
Ferrous base, coating	geosynthestic reinforcement	
Fiber LengthT 126	grab breaking load	
Filler, mineralT 37	puncture	
Fine aggregate:	sampling	
angularity T 304	sewen seams	
absorptionT 84	tear strength	
clay lumps in	thickness	
elutriation	water permeability	
friable particles	wide width breaking load	
lightweight material T 113	Geosynthetic reinforcement	
mortar strength	sampling	
organic impuritiesT 21	Glass electrodes	
samplingT 2	Glass electrodes for pH determination	
sieve analysis T 27	Granular soilT	
specific gravity	Grab breaking load	
unit weightT19	Gravel (see coarse aggregate)	2 .002
Fineness of cement:	Grout cubesT	106. T 813
air permeability	Grout flow	
Fire point, bituminous materials T 48	Gyratory compactor	
Fire testing	Sylmoly Companies	
Flash point, bituminous materials:	Н	
cleveland open cup	High float	Т 50
tagliabueT79	Hydraulic cement (see cement)	1 30
Flat and elongated aggregate particles D 4791	Hydromulch	Т 127
Flexural creep	Hot mix (see bituminous mixture)	1 12/
Flexural strength concrete:	That this (See oftuninous mixture)	
center-point loading		
specimens taken from hardened concrete T 24		
third-point loading T 802		

l		M
Identification of soils	D 2488	Manufacturing processesT 40, T 168
Illumination:		Maximum dry densityT 99, T 180, T 606,
instrumental photometric	T 257	Mechanical analysis:
photovolt reflectance		aggregateT 27/T 11
retroreslective		extracted aggregateT 27/T 11
Impact sampling tests	T 244	mineral filterT 37
Ignition furnace:		soilT 88
calibration	SOP 728	Mechanical mixingT 162
fineness	D 1208	Mechanical properties of steel fasteners T 606
HMA mixture	T 308	Mechanical testing of steel
loss	D 1208	Melting joint and creack sealant D 5167
Inorganic matter in bituminous materials	T 111	Membrane, waterproofing T 413
_		Metallic materials:
J		impactT 244
J-Ring (see concrete)		Mineral filler:
Joint sealants:		mechanical analysisT 37
hot poured D 5329, T 4	8 T 412	MiscibilityT 59
two component rubber		Mixing, mechanical of hydraulic cement T 162
Joint sealants - bond test		Mixtures, bituminous (see bituminous mixtures)
Joint sealars for concrete		Modulus of soilsT 307
Joint sealant, flow test		Moisture analysis T 217
Joint Scalant, now test	D 3327	Moisture content HMA T 329
K		Moisture-densityT 310, SOP 615
V:	т 70	Moisture-density relations:
Kinematic viscosity (general)		soil:
Kinematic viscosity of asphalt	1 201	5.5 lb rammer
L		10 lb rammer T 180
		Mortar, cement:
Lane markers, type 2		length changeT 168
Latex - cement compatibility		mechanical mixingT 162
Latex nonvolatile content		water soluable chloride
Latex paints, pH		Mulch, barkT 123, T 125
Leachate Sample		
Length of concrete cores		N
Lightweight concrete		Normal consistency of hydraulic cement T 129
Lighweight particles in aggregate		Nuclear methods:
Liquid limit	T 89	densityT 310, TM 8, SOP 615, SOP 729
Longitudinal Joint Density		delisity 1 310, 110 8, 301 013, 301 729
Long-term aging		0
Loop Amplifying		
Los Angeles abrasion		Oils
Los Angeles test		Open cup:
Loss on heating of asphalt compounds	D 6	flash and fire points T 48
		Organic impurities in sand T 21
		Organic matter in soils T 267

P	Pressure testing	
Package stability T 200	Profilograph, California	
Paint:	PycnometerT 228,	T 209
condition in container FTMS 3011	D	
density	R	
directional reflectance E 1347	Random sampling	T 716
drying timeFTMS 4061, D 1640	Rebound hammer determination	
fineness of dispersion of pigmentD 1210	Recovery of asphalt	TP 2
hiding powerFTMS 4122	Reflective materials and devices T 257, E	4956
infrared identification	ResidueT 59	
loss on ignition	Resiliend modulus of soils	T 307
pigment content	Resistance (R valve) of soil	
viscosity	Resistivity of soilT 288,	T 417
volatiles	Retroreflectivity	T 429
volume of nonvolatiles	Reverse-flow viscometers	
Paraffin, coating of bituminous	Ring-and-ball apparatus	T 53
aggregate mixture	Ring-and-ball method, bituminous material	
pavement friction	Road oil	
Particle charge test	Rolling thin film oven	
Parting compound T 816	Rotational viscosity T 316, D 4402, D	
Paving materials T 228	•	
Penetration T 49	S	
Penetration, bituminous materials	Sampling:	
Permeability	aggregate	Т 2
Petroleum products:	bituminous materials	
water in	bituminous mixturesT 168, SC	
pH	concrete	
soils	fresh concrete	
Photovolt reflectance	gravel	
Pitches	geosynthetics	
Plastic fines in:	geotextiles, sampling	
aggregates and soils T 176	sand	
Plastic limit of soil	slag	
Plastic pavement marking tapeD 4505	stone	
Pore-water T 297	stone block	
Portland cement (see cement)	Sampling concrete/mortar/aggregates	
Pour pointT 187	Sand (see fine aggregate)	1 2
Pouring temperatureT 50	Sand equivalent test	T 176
Power interruption test T 424	Saybolt-Furol viscometers	
Preparation of concrete specimens:	Self compacting concrete:	, 1 /2
cappingT 231	air content	T 818
in field	cylinders	
in laboratory T 126	Semi-solid materials T 40, T 49, T 168,	
self compacting T 819	Settlement	
Preparation of disturbed soil samples for test:	Sewen seams	
dry method	Short-term aging	
Pressure aging R 28	choir wim uging	

Sieve analysis:	Solid phase materials	T 40, T 49
aggregateT 27	Solubility of bituminous materials.	
bark MulchT 123	Solvita test	
extracted aggregateT 30	Soundness test for:	
mineral fillerT 37	hydraulic cement	T 107
soilT 88, T 601	Portland cement	
Signal controlers T 421, T 422, T 423, T 424,	Specific gravity:	
T 425, T 427, T 428, SOP 429	bituminous materials	T 228, T 229
Signs	bituminous mixturesT 166	
Skid resistance T 242	coarse aggregate	T 85
Slag (see coarse aggregate)	fine aggregate	
Slump-self consolidating	hydraulic cement	T 133
Slump test of concrete	maximum, for bituminous paving	
Softening point of bituminous material T 53	T 209	
Soil permeabilityT 215	petroleum	D 1298
Soils:	soil	T 100
capillary riseT 610	tar	T 228
classification	water	
compaction and density correction for	Spike test	
coarse particlesT 99	Splitting:	
consolidation propertiesT 216	aggregate	T 248
density in placeT 310, SOP 615	hot mix asphalt	T 712
dry preparation for testT 87	Steel:	
family of curves	impact	T 244
identification	mechanical testing	
liquid limitT 89	testing	
mechanical analysis T 88	Stiffness of asphalt	
moisture:	Stone: (see also coarse aggregate)	
by calcium carbide methodT 217	sampling	T 2
by laboratory determinationT 265	Storage stability	
moisture-density relations: 5.5 lb rammer . T 99	Stormer viscosity	
moisture-density relations: 10-lb rammer T 180	Strength of mortar	
nuclearT 310, SOP 615	Stress	
organic matter T 267	Stripping of bituminous aggregate	
permeability T 215	mixtures	T 706, T 718
pHT 289	Superpave mix design	
plastic limitT 90	Superpave volumertics	
plasticity index T 90	1 1	
preparation for tests:	T	
dry method T 87	Tar:	
sand equivalent testT 176		т 220
specific gravityT 100	specific gravity	
unconfined compressive strength T 208	Tars	
resilient modulus of soils	Tear strength, geotextiles	
resistance (r-valve) T 611	Temperature tests	
resistivityT 288, T 417	Tension Tensile) properties/tests	
1 200, 1 41/	Thermoplastic insulation	D 2633

Time of setting:	Weight per cubic foot:	
cement, hydraulic:	aggregates	T 19
gillmore needles T 159	Wire, test for:	
vicat needleT 131	cross linked insulation	D 470
concreteT 197	Thermoplastic insulation	D 2633
Traffic signal controllersT 421, T 422, T 423, T 424, T 425, T 427, T 428, SOP 429	weight of coating	T 65
Triaxial compression	Z	
Type 2 lane markers T 426	Zinc coating:	
U	weight	T 65
Unconsigned compressure strength of soil		
rock		
Uncrushed solid material T 40 Unit weight of aggregate T 19		
V		
Vacuum capillary viscometer		
Vicuum testing T 297		
Viscometers		
Viscosity		
Viscosity tests: absolute		
kinematic T 201		
rotational T 316, D 4402, D 2196		
sayboltT 72		
stormer		
Volatile matter content		
Voltage spike test		
Volumertic properties SOP 731		
W		
Washing test for aggregates T 11		
Water contentT 55, T 59		
Water displacement test T 229		
Waterproofing membraneT 413		
Water penetration		
Water permeabilityD 4492		
Water soluable chloride in concrete C 1218		
Water, test for:		
emulsions T 59		
petroleum products		
specific gravity		