# MATERIALS TESTING MANUAL
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MATERIALS TESTING MANUAL

INTRODUCTION

The Materials Testing Manual has been prepared for the purpose of standardizing sampling and testing procedures in the various laboratories which test materials for conformance to Arizona Department of Transportation specifications. The procedures found in this manual fall into one of four categories.

1. Those procedures which follow an established standard. The American Association of State Highway and Transportation Officials (AASHTO) is the primary contributor. Some procedures are conducted in accordance with the American Society for Testing and Materials (ASTM). Commonly used procedures in this category are listed in the Table of Contents, but these listings are not intended to include all procedures referenced in the Arizona Department of Transportation specifications.

2. Those procedures which generally follow an established standard, but which have been modified in some degree. They have been designated as modifications in a subheading beneath the title.

3. Those procedures that are modifications of testing procedures used by other state transportation departments.

4. Those procedures which have been originated by the Arizona Department of Transportation and which may not be fully described in national publications. These have been subtitled "An Arizona Method".

Each procedure in category 1 listed above is assigned an alphanumeric designation either by AASHTO or ASTM. AASHTO publishes the “Standard Specifications for Transportation Materials and Methods of Sampling and Testing” and the “AASHTO Provisional Standards” annually. Unless otherwise specified, the effective date of AASHTO standards is the first day of the month following publication and distribution of the published standards. Additional information regarding AASHTO standards can be obtained on the AASHTO website (www.transportation.org). The “Annual Book of ASTM Standards” is published in separate volumes throughout the year. Additional information regarding ASTM standards can be obtained on the ASTM website (www.astm.org).

Each procedure in category 2, 3, or 4 listed above is given a number prefixed by "ARIZ". Revised test designations will have a letter suffix, for example, the first revision of test method
ARIZ 201 would be ARIZ 201a, the second revision will be 201b, etc. Each procedure, or revision to it, is also identified by a date on which it becomes effective. References made to procedures in the manual which either do not include a letter suffix or have a suffix which has been superseded shall be understood to refer to the latest revision of the respective numbered procedure. The Table of Contents lists all Arizona sampling and testing procedures included in the manual, with their respective necessary letter suffixes.

The manual is divided into 10 different series, and an Appendix section, concerned with sampling and testing procedures for specific materials, design methods, or certification procedures.

It is the intent of Construction & Materials Group to describe the procedures in such detail that a competent operator who is unfamiliar with the method can obtain accurate results when the procedure is followed in detail. We will welcome any suggestions for improvement in these procedures that will assist in accomplishing this objective.

All revisions to the Materials Testing Manual shall officially originate from Construction & Materials Group, Quality Assurance Section, under the direction of the ADOT Standards Committee.

Revisions will be issued following approval by the ADOT Standards Committee. Revisions issued will be effective for projects with a bid opening date on or after the effective date of the revisions.

The Construction & Materials Group will welcome any suggestions for the improvement of the Materials Testing Manual, as it is hoped and intended that manual users will participate its formulation and revision.
GLOSSARY OF TERMS

**Abrasion** - The wearing away of the surface of a solid by friction.

**Abscissa** - The distance measured in the "x" or horizontal direction on a graph.

**Absorption** - Process of a soil or aggregate holding a fluid mechanically, by capillary action. This applies both to water and asphalt.

**Adhesion** - (1) Shearing resistance between soil and another material under zero externally applied temperature.

(2) Molecular attraction exerted between the surfaces of bodies in contact.

**Admixtures for Portland Cement Concrete** - Materials other than water, aggregate, Portland cement, and fiber reinforcement that is used as an ingredient of Portland cement concrete to modify its freshly mixed, setting, or hardened properties and that is added to the batch before or during its mixing.

**Aggregate** - Any hard, inert mineral material used for mixing in graduated fragments. It includes sand, gravel, crushed stone, and blast-furnace slag.

(1) Coarse aggregate for Portland cement concrete - that retained on a No. 8 (2.36 mm) sieve.

(2) Fine aggregate for Portland cement concrete - that passing a 3/8" (9.5 mm) sieve and almost entirely passing a No. 4 (4.75 mm) sieve, and predominantly retained on a No. 200 (75μm) sieve.

**Air-Entraining Agent** - An admixture used in Portland cement concrete to increase the amount of entrained air in the mixture. Air is entrained in minute bubbles in the concrete or mortar during mixing.

**Aliquot** - A part of the whole that divides evenly into the whole, as 3 mL is an aliquot portion, 1/8 of 24 mL.

**Anhydrous** - A material which has an affinity for water due to the fact that the water in its crystalline structure has been removed.

**Asphalt** - A dark brown to black cementitious material, solid or semi-solid in consistency, in which the predominating constituents are bitumens which occur in nature as such or are obtained as residual in refining petroleum.
Asphalt Cement - Asphalt that is refined to meet specifications for paving, industrial, and special purposes.

Asphalt Emulsion Slurry Seal - A mixture of slow-setting emulsified asphalt, fine aggregate and mineral filler, with water added to produce a slurry consistency.

Asphalt Joint Fillers - Asphaltic products used for filling cracks and joints in pavement and other structures.

Asphaltic Concrete - A thoroughly controlled hot mixture of asphalt cement and well-graded aggregate, thoroughly compacted into a uniform dense mass.

Auger - A drill for test holes in unconsolidated material modeled after the conventional screw auger.

Bitumen - Hydrocarbon material of natural and/or pyrogenous origin, frequently accompanied by their non-metallic derivatives, which may be gaseous, liquid, semi solid or solid, and is completely soluble in carbon disulfide.

Bleeding - (1) Upward migration of bituminous material resulting in a film of asphalt on the surface.

(2) Escape of water from freshly placed concrete commonly observed as an accumulation on a horizontal surface.

Boulder - A rock fragment, usually rounded by weathering or abrasion, which will be retained on a 3 inch (75 mm) sieve.

Capillarity - The rise or movement of a liquid in the interstices of a soil due to capillary forces.

Chemical Change - Any change resulting in formation of a new substance.

Clay (Clay Soil) - Fine-grained soil or the fine-grained portion of a soil that can be made to exhibit plasticity within a range of water contents and that exhibits considerable strength when air-dry.

Coal Tar - Tar produced by destructive distillation of bituminous coal.

Cohesion - (1) The property of a soil that makes its particles stick together (clay is a cohesive soil).

(2) The property of a bituminous material that holds the mass together.
Colloidal - Pertaining to soil particles that are so small (less than 0.001 mm) that the surface activity has an appreciable influence on the properties of the aggregate.

Compaction - The densification of a soil by means of mechanical manipulation.

Compressibility - Property of a soil pertaining to its susceptibility to decrease in volume when subjected to load.

Consistency - (1) The relative ease with which a soil can be deformed.

(2) Relative mobility of freshly mixed concrete commonly defined as slump.

Corrugations - Regular transverse undulations on the surface of sheet metal, consisting of alternate valleys and crests.

Curing of Concrete - A period provided to prevent formation of surface cracks due to rapid loss of water while concrete is plastic, and to assure attainment of maximum strength.

Curing Compound (Liquid) - A liquid, membrane-forming material used in curing Portland cement concrete, which prevents moisture loss during the early hardening period. Papers and other impermeable sheet materials can be used for the same purpose.

Cutback (Liquid) Asphalt - Asphalt cement which has been liquified by blending with petroleum distillates. Upon exposure to atmospheric conditions the diluents evaporate, leaving the asphalt cement to perform its function.

(1) Rapid Curing (RC) - Composed of asphalt cement and a naphtha or gasoline-type diluent of high volatility.

(2) Medium Curing (MC) - Composed of asphalt cement and a kerosene-type medium volatility.

(3) Slow Curing (SC) - Composed of asphalt cement and oils of low volatility.

Deflocculant - An agent that prevents fine soil particles in suspension from coalescing to form flocs; for example the working solution employed in the Sand Equivalent test.

Density (Unit Weight) - Weight of a material per unit volume.

Disintegration - Deterioration or separation into smaller fragments from any cause.
Distillation - Process of driving off gas or vapor from liquids or solids, with heat, in a retort or still, and condensing products therefrom.

Ductility - Capability of being drawn out or hammered thin. A common test for bituminous materials, which determines, indirectly, the adhesive quality, and temperature susceptibility.

Elasticity - That property of a body which causes it to recover its original shape and size when it is deformed and the deforming forces are removed.

Emulsified Asphalt - A heterogeneous system containing two normally immiscible phases (asphalt cement and water) with a small amount of emulsifying agent. It exists in a liquid state at normal temperatures. Emulsified asphalts may be anionic, with electro-negatively charged asphalt globules; or cationic, with electropositively charged asphalt globules, depending upon the emulsifying agent.

Erosion - The loosening and transporting of rock debris and soil by moving agents operating on the earth's surface. The four main agents are: wind, waves and currents in bodies of water, glaciers, and running water.

Expansion Joint - Joint permitting pavement to expand in length.

Fabricating Plant - A plant where steel members are riveted or welded together to form steel beams, trusses, metal pipe, etc.

Fineness Modulus - An index of the fineness or coarseness of an aggregate; the summation of the cumulative percentages of the material retained on the specified series of sieves divided by 100. It is **not** an indication of grading. Sieve sizes used are No. 100 (150 μm), No. 50 (300 μm), No. 30 (600 μm), No. 16 (1.18 mm), No. 8 (2.36 mm), No. 4 (4.75 mm), 3/8 inch (9.5 mm), 3/4 inch (19.0 mm), 1-1/2 inch (37.5 mm), and larger, increasing in the ratio of 2 to 1.

Floc - Loose, open-structured mass formed in a suspension by aggregation of minute particles.

Flocculation - Process of forming flocs.

Free Water (Ground Water) - Water that is free to move through a soil mass under the influence of gravity.

Friable - Easily broken or crumbled.
Frost Action - Freezing and thawing of moisture in materials and the resultant effects on these materials and on structures of which they are a part or with which they are in contact.

Gradation - Proportion of material of each grain-size present in a given aggregate sample.

Granular - Material that does not contain more than 35 percent of soil particles which will pass a No. 200 (75 µm) sieve.

Gravel - Rounded or semi-rounded particles of rock that will pass a 3 inch (75 mm) sieve and be retained on a No. 10 (2.00 mm) sieve.

Heave - Upward movement of soil caused by expansion or displacement resulting from phenomena such as moisture absorption, removal of overburden, driving of piles, frost action, etc.

Honeycomb - A surface or interior defect in a concrete mass characterized by lack of mortar between the coarse aggregate particles.

High-Early Strength Cement - A finer ground cement with a different chemical composition that reaches a high strength in a shorter period of time than regular cement.

Hydrated Lime - A dry powder obtained by treating quicklime with water enough to satisfy its chemical affinity for water under the conditions of its hydration. It is predominantly calcium hydroxide.

Hydraulic Cement - A cement that sets and hardens by chemical interaction with water and that is capable of doing so under water.

Hydrophilic - Literally, water-loving. The property that defines a material as having an affinity (attractive force) for water.

Hydrophobic - Literally, water-averting. The property that defines a material as lacking an affinity (attractive force) for water.

Hygroscopic Water - Water retained in an air-dried soil. It is held by each soil grain in a very thin film and has both physical and chemical affinity for the grain. This moisture is in equilibrium with the air humidity.

Joint (Pavement) - A narrow space separating two slabs or sections of pavement.

Leaching - The removal of soluble salts or other soluble particles by percolating water.
**Liquid Limit** - (1) The moisture content which is the boundary between the liquid and plastic states for the minus No. 40 (425 μm) fraction of a soil.

(2) That moisture content at which a soil fraction will close a standard groove for a length of about 1/2 inch (13 mm) when subjected to 25 blows in a liquid limit device.

**Macadam** - An asphalt pavement construction using a coarse, open-graded aggregate that is usually produced by crushing and screening stone, slag or gravel.

**Maximum Size of Soil and Aggregate** - The smallest sieve opening through which the entire amount of material will pass.

*Note:* Another definition, which applies only to aggregates used in Superpave mixes (Section 417, "Asphaltic Concrete (End Product) SHRP Volumetric Mix", of the ADOT Specifications) is: Maximum Aggregate Size - One size larger than the nominal maximum aggregate size.

**Mesh** - The square opening of a sieve.

**Modulus of Rupture** - A measure of the flexural strength of concrete when it is broken by bending. The higher the values the greater the strength.

**Moisture Content** - (1) The proportion of moisture in soil or aggregate material, expressed as a percent of the dry weight.

(2) Optimum Moisture Content - The percentage of moisture at which the greatest density of a particular soil can be obtained through compaction by a specified method.

(3) The proportion of moisture in bituminous mixtures, expressed as a percent of the wet weight.

**Nominal Maximum Size of Soil and Aggregate** - The smallest sieve opening through which the entire amount of material, by specification, is permitted to pass.

*Note:* Another definition, which applies only to aggregates used in Superpave mixes (Section 417, "Asphaltic Concrete (End Product) SHRP Volumetric Mix", of the ADOT Specifications) is: Nominal Maximum Aggregate Size - One size larger than the first sieve that retains more than 10 percent aggregate.

**Ordinate** - The distance measured in the "y" or vertical direction on a graph.
Overlay - One or more courses of asphalt construction on an existing pavement, generally including a leveling course, followed by a uniform course or courses to provide needed thickness.

Penetration - The consistency of a bituminous material expressed as the distance that a standard needle vertically penetrates a sample of the material under known conditions of loading, time and temperature.

Permeability - That property of a material which permits a liquid to flow through its pores or interstices.

pH - An index of acidity or alkalinity, in terms of the logarithm of the reciprocal of the hydrogen ion concentration. A pH of seven, the value for pure distilled water, is regarded as neutral; pH values of less than seven indicate acidity; and, pH values of greater than seven indicate alkalinity.

Plasticity - The property of a soil which allows it to be deformed beyond the point of elastic recovery without cracking or appreciable volume change.

Plasticity Index (PI) - Numerical difference between the liquid limit and the plastic limit.

Plastic Limit - (1) The water content corresponding to an arbitrary limit between the plastic and semisolid states of consistency of a soil.

(2) Water content at which a soil will just begin to crumble when rolled into a thread about 1/8 inch (3 mm) in diameter.

Porosity - The ratio, in percent, of the volume of void space (pores) of a material to the total volume of its mass.

Portland Cement - A hydraulic cement made by pulverizing portland-cement clinker (a clinker, partially fused by pyroprocessing, consisting predominantly of crystalline hydraulic calcium silicates), and usually containing calcium sulfate.

Prime Coat - The first application of a bituminous material to an existing absorbent surface.

Pumping - The ejection of foundation soil wet or dry, through joints or cracks, or along edges of rigid slabs, due to vertical movements under traffic.

Quicklime - A calcined material, the major part of which is calcium oxide or calcium oxide in association with a lesser amount of magnesium oxide, capable of slaking with water.
Raveling - Progressive disintegration of a pavement surface by dislodgment of aggregate particles.

Relative Density - (1) Soils - The ratio of (1) the difference between the void ratio of a cohesionless soil in the loosest state and any given void ratio, to (2) the difference between the void ratios in the loosest and in the densest states. This definition is expressed as an equation below:

\[ Dr = \frac{E_{\text{max}} - En}{E_{\text{max}} - E_{\text{min}}} \]

Where: \( Dr \) = Relative density of cohesionless soils
\( E_{\text{max}} \) = Void ratio of soil in the loosest state
\( E_{\text{min}} \) = Void ratio of soil in the densest state
\( En \) = Any given void ratio

(2) Asphaltic Concrete – Often used for specimens prepared by means of the Superpave gyratory compactor; the ratio, expressed as a percentage, of the bulk specific gravity (at any number of gyrations) to the theoretical maximum specific gravity.

Riprap - Broken rock used for the protection of embankments, cut slopes, etc., against agents of erosion, primarily water.

Road Mix - A mixture of aggregate and asphalt prepared in place.

Rock - Natural solid mineral matter occurring in large masses or fragments.

Rutting - Formation of longitudinal depressions by the lateral displacement of soils or surfaces under traffic.

Sand - Particles of rock that will pass the No. 4 (4.75 mm) sieve and be retained on the No. 200 (75 μm) sieve.

Saturated Surface-Dry - Term used to describe the condition of an aggregate in which the pores of all the particles are completely filled with water, but their surfaces are free from moisture.

Screed - To strike off excess material to bring the top surface to proper contour and elevation.

Screen - An apparatus, in which the apertures are circular, for separating sizes of material.
Seal Coat - A thin asphaltic surface treatment used to improve the texture of and to waterproof an asphalt pavement surface.

Settlement - The reduction in elevation of short sections of pavement or structures, due to compressibility of underlying soils.

Shear Strength - The maximum resistance of a soil to shearing stresses.

Shear Stress - An action or stress, resulting from applied forces, which tends to cause two contiguous parts of a body to slide relatively to each other in a direction parallel to their plane of contact.

Sieve - An apparatus, in which the apertures are square, for separating sizes of material.

Silt - Material passing the No. 200 (75 μm) sieve that is nonplastic or very slightly plastic and that exhibits little or no strength when air-dried.

Slump - The measure of the consistency of concrete determined by placing the concrete in a standard slump cone, removing the cone, and allowing the concrete to settle under its own weight.

Soil - Sediments or other consolidated accumulations of solid particles produced by the chemical and physical disintegration of rocks, and which may or may not contain organic matter.

Soil Profile - Vertical section of a soil, showing the nature and sequence of the various layers, as developed by deposition or weathering or both.

Soundness - Resistance to both physical and chemical deterioration.

Spalling - Peeling away of a surface, particularly Portland cement concrete.

Specific Gravity (Absolute) - The ratio of the weight in air of a given volume of material at a stated temperature, to the weight in air of the volume of water which the material displaces.

Specific Gravity (Apparent) - Ratio of the weight in air of a given volume of impermeable material (that is, the solid matter including the impermeable pores) to the weight of an equal volume of water at a stated temperature.

Specific Gravity (Bulk) - Ratio of the weight of a given volume of permeable material (including permeable and impermeable voids) to the weight of an equal volume of water at a stated temperature.
Spelter - The zinc coating used for rustproofing iron or steel by galvanizing.

Stability - Property of a material which enables it to retain its other essential characteristics throughout the range of conditions expected in service.

Stone - Crushed or naturally angular particles of rock that will pass a 3 inch (75 mm) sieve and be retained on a No. 10 (2.00 mm) sieve.

Stripping - Separation of bituminous films from aggregate particles due to presence of moisture.

Swell - Increase in volume due to the absorption of water into the intergranular pore space.

Tack Coat - A very thin spraying of bituminous material on an existing surface to increase the adhesion, or bond, between a new course of construction and the old surface.

Titration - The volumetric addition of a solution, of known concentration, to another solution to determine an end point reaction.

Top Soil - Usually the upper 6 inches (150 mm) of native soil and that portion used in dressing and landscaping earth slopes.

Viscosity - The property of a fluid to resist internal flow.

Void - Space in a mass not occupied by solid mineral matter; it may be occupied by air, or other fluids.

Warping - Deviation of pavement surface from original shape caused by temperature and moisture differentials within the slab.

Yield - Ratio of the volume of a Portland cement concrete mixture produced to that which theoretically should be produced.
NOTE: Additional information on Terminology and Definitions of Terms can be found in various other sources, including the following:

AASHTO M 146: "Terms Relating to Subgrade, Soil-Aggregate, and Fill Materials".

AASHTO R 10: "Definition of Terms for Specifications and Procedures".

ASTM C 125: "Standard Terminology Relating to Concrete and Concrete Aggregates".


SERIES 100
SAMPLING
SAMPLING BITUMINOUS MATERIALS
(An Arizona Method)

1. SCOPE

1.1 This procedure covers best practices for sampling of Bituminous materials (paving grade asphalt, crumb rubber asphalt and emulsions) in the field.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 For the purpose of this test method Bituminous materials other than Emulsions will be referred to as “Asphalt Binder”, and Emulsified Bituminous material i.e. (RS-1, SS-1, CSS-1, etc...) will be referred to as “Emulsions”.

2. SIZE OF SAMPLES

2.1 A minimum of 1 gal. of Asphalt Binder.

2.2 A minimum of two ½ gal. containers per sample of Emulsions.

3. CONTAINERS

3.1 Containers for Asphalt Binder, shall be double friction top cans.

3.2 Containers for Emulsion samples shall be wide mouth containers made of plastic.

4. PROTECTION AND PRESERVATION OF SAMPLES

4.1 Containers shall be new and free of any moisture, contaminants, or residue from any manufacturing process. The top and container shall fit together tightly.
4.2 The container shall be tightly sealed immediately after obtaining the sample.

4.3 The filled sample container shall not be cleaned using a solvent. If cleaning is necessary use a clean dry cloth.

4.4 Samples of Emulsion shall be protected from freezing.

4.5 Transferring samples from one container to another shall be avoided if possible.

4.6 Sample containers shall be identified using sample tags that are securely fastened to the side of the container so they will not be lost in transit, and shall be clearly marked for identification with a suitable permanent marker on the side of the container itself.

5. PROCEDURE

5.1 Samples of Asphalt Binder shall be taken from the last possible point before the bituminous material is introduced into the hot plant. This is usually from a spigot or faucet on the circulation line.

5.2 Bituminous materials applied to pavement surfaces, i.e. Tack Coat, Fog Coat shall be sampled from the distributor truck at the project.

5.3 Clearly identify the side (not the lid) of a new clean container of appropriate size with the sample number, date, project number, type of material, and any other pertinent information.

5.4 To ensure the sample is representative, draw off and discard a minimum of 1 gal. of Bituminous material prior to obtaining the sample from the sampling valve.

5.5 From the sampling valve draw off the minimum amount of Bituminous material required for the type of material being sampled. Care should be taken to avoid spilling any material on the outside of the container or over filling the container. The container should be filled to no closer than one inch from the top.

5.6 Immediately after obtaining the sample, the clearly identified container shall be tightly and positively sealed.
SAMPLING ASPHALTIC CONCRETE MIXTURES

(An Arizona Method)

1. SCOPE

1.1 This procedure describes the methods which are to be used when sampling asphaltic concrete mixtures in order to best assure representative samples of the materials being placed.

1.2 Sampling asphaltic concrete mixtures by this procedure may involve hazardous material, operations, or equipment. This procedure does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 For the purpose of this method, asphaltic concrete mixtures are referred to as either Asphaltic Concrete (AC) or Asphaltic Concrete Friction Course (ACFC), regardless of the asphalt binder type.

2. APPARATUS

2.1 Sampling Plate - A 4 foot x 1 foot x 1/16 inch steel plate, which has been prepared with a 1/8 inch hole at each corner of one end and a sufficient length of wire tied through each hole to form a loop approximately 4 feet in length. The sampling plate may be used when sampling AC mixtures.

2.2 Cookie-Cutter Template - As an alternate to obtaining the sample from the sampling plate, a rectangular metal template ("cookie cutter") and metal plate of sufficient size may be used to sample the AC mixture. The metal template and plate shall be of sufficient size so that the desired amount of material is obtained by a single use of the template and plate at any one location. The metal plate shall be prepared with a wire(s) of sufficient length attached to each corner on one side of the metal plate (the short side when the plate is not square) so the metal plate may be located by raising the wire(s) after the laydown machine has passed.
2.3 Miscellaneous Brushes - May be helpful when obtaining the entire amount of AC mixture from a cookie-cutter template.

2.4 Flat Square Point Shovel - A flat square point shovel shall be used when sampling AC mixtures. A flat square point shovel may also facilitate sampling ACFC mixtures and when sampling by other procedures as described below.

2.5 5-Gallon Bucket, or other suitable container - Shall be used when sampling AC mixtures and ACFC mixtures.

3. SIZE OF SAMPLES

3.1 For AC mixtures designed with Marshall design procedures, minimum 75 pounds.

3.2 For AC mixtures designed with Gyratory design procedures, minimum 130 pounds.

3.3 For ACFC mixtures, minimum 50 pounds.

4. SAMPLING AC MIXTURES

4.1 The sampling plate shall be placed on the roadway just ahead of the laydown machine. Except for wider mats when a sample is being taken from the middle of the mat, the sampling plate is placed so that the end with the wire is approximately one foot in from the right or left edge of the mat being laid. The sampling should be alternated between the right and left edges when practical. The wire attached to the end of the plate shall be held to the ground to allow the laydown machine to pass over the plate and wire.

4.2 After the laydown machine has passed, locate the plate by raising the wire.

4.3 The sample shall be taken from the plate using a flat square point shovel. The sample shall consist of the full depth of material for one shovel width from the center portion of the plate over its entire length. Material covering the entire plate shall not be taken. A single pass of the shovel shall be made, moving along the surface of the plate until the shovel is full. Carefully deposit the AC mixture into a 5-gallon bucket, or other suitable container. Material which has sloughed into the resultant trench shall not be obtained. At the next undisturbed area of material on the plate, repeat shoveling and placing the material into the container. If necessary, additional material may be obtained by using an
additional plate(s) in the immediate vicinity and combining all material. The use of an additional plate(s) cannot be used in lieu of splitting.

**Note:** When sampling with the “cookie cutter”, the metal plate shall be placed on the roadway at the location where the sample is to be taken, just ahead of the laydown machine. If the metal template is not square, it shall be placed on the roadway so that the longest side is in a transverse direction across the roadway. The wire(s) shall be held to the ground to allow the laydown machine to pass over the plate and wire(s). After the laydown machine has passed, locate the plate by raising the wire(s). The template is pressed through the AC mixture until it rests squarely upon the plate. The entire amount of AC mixture is removed from the interior of the template and carefully placed into a 5-gallon bucket, or other suitable container. Obtaining multiple samples cannot be used in lieu of splitting.

5. **SAMPLING ACFC MIXTURES**

5.1 When sampling ACFC mixtures, an adequate amount of material shall be taken from the truck at the mixing plant and placed into a 5-gallon bucket, or other suitable container. The sample shall be taken from at least 3 random locations, approximately 12" below the surface, within five minutes from the time the loading of the truck is completed.

5.2 Material that is to be tested immediately after it has been sampled shall be protected to avoid heat loss while it is being transported to the laboratory.

6. **SAMPLING FINISHED AC PAVEMENT**

6.1 Samples of AC from finished pavement shall be taken through the complete thickness of the pavement or lift, in such a manner which causes minimum disturbance to the sample.

6.2 If a coring apparatus is used, the coring bit shall be subjected to enough vertical pressure to penetrate the pavement without causing damage to equipment or disturbance of the sample. The minimum core diameter shall be 4 inches.
6.3 If coring equipment is not available, the sample may be taken with the use of a saw, pick, jackhammer, or other suitable means if a suitable specimen can be obtained for the intended testing.

6.4 All samples shall be handled carefully so that they maintain their original form. The samples shall be transported on a relatively flat surface, and adequately protected to preserve their shape and to prevent damage.

6.5 The use of ice may be found helpful in obtaining and/or preserving the condition of the specimen.

6.6 Samples shall be delivered to the laboratory for testing as expeditiously as reasonably possible. Samples shall be transported carefully in a covered container out of extreme environmental conditions.

7. SAMPLING MISCELLANEOUS PLACEMENT OF AC MIXTURES

7.1 When required, samples of AC mixtures placed in miscellaneous areas shall be obtained from locations and by means to provide appropriate representation of the AC mixture being placed. Miscellaneous areas are locations where representative samples would be difficult to obtain in-place due to geometry, paving area size, limited access, or other factors. These areas could include paving slopes, median islands, utility trenches, tapers, radius paving and any other area designated by the Engineer.

8. SAMPLE IDENTIFICATION

8.1 Each sample shall be identified by an accompanying sample ticket. Sample tickets shall be filled out as required to provide necessary information. The remarks area of the sample ticket shall be used as necessary to provide additional information, including the phone number of an individual who can be contacted regarding the sample.

8.2 The source of the sample shall be the "original source" of the material, as indicated on the sample ticket.

8.3 An example of a completed sample ticket used by ADOT for construction projects is shown in Figure 1. Commonly used codes for filling out the sample ticket are shown on the back side of the sample ticket (see Figure 2).
8.4 The sample ticket consists of three copies. The center copy is kept by the person submitting the sample, the original copy is included inside the sample container, and the third copy is attached to the sample container. When filling out sample tickets, make certain information is clear and easily read on all three copies.
### FIGURE 1

**EXAMPLE**

Contact Phone No. - 999-999-9999

### FIGURE 2
1. SCOPE

1.1 This method describes the methods which are to be used when sampling soils and aggregates.

1.2 Sampling is equally as important as the testing, and the individual doing the sampling shall use every precaution to obtain samples that will be representative of the materials being sampled.

1.3 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.4 Table 1 shall be used to determine minimum sample weights based on the size of aggregate. The amount of material required may be greater depending on the tests that are to be performed on the material.

<table>
<thead>
<tr>
<th>Nominal Maximum Aggregate Size *</th>
<th>Sample Mass</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>lbs</td>
</tr>
<tr>
<td><strong>Fine Aggregate</strong></td>
<td></td>
</tr>
<tr>
<td>#8</td>
<td>22</td>
</tr>
<tr>
<td>#4</td>
<td>22</td>
</tr>
<tr>
<td><strong>Coarse Aggregate</strong></td>
<td></td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>22</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>35</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>55</td>
</tr>
<tr>
<td>1&quot;</td>
<td>110</td>
</tr>
<tr>
<td>1-1/2&quot;</td>
<td>165</td>
</tr>
<tr>
<td>2&quot;</td>
<td>220</td>
</tr>
<tr>
<td>2-1/2&quot;</td>
<td>275</td>
</tr>
<tr>
<td>3&quot;</td>
<td>330</td>
</tr>
</tbody>
</table>

* The smallest sieve opening through which the entire amount of material, by specification, is permitted to pass.
2. SAMPLING FROM STOCKPILES

2.1 In sampling materials from stockpiles it is difficult to ensure unbiased samples, due to the segregation which often occurs when the material is stockpiled with coarser particles rolling to the outside base of the pile. If power equipment is available then it would be advantageous to enlist the use of that equipment to develop a separate, small sampling pile composed of materials drawn from various levels and locations in the main stockpile. Once a small sampling pile has been established then a sample shall be taken from that pile by taking several increments and combining.

2.2 The stockpile may also be sampled by placing a wood or metal shield upslope from the point of sampling to prevent loose aggregate from sliding down into the sampling area. Remove approximately 3 to 6 inches of material from the sampling area. Utilizing a square point shovel, take a sample near the top, at the middle and near the bottom of the stockpile. The sample taken at each location shall be one shovelful of material. Repeat this operation at the sampling locations as shown in Figure 1, and combine all samples taken from the stockpile.

3. SAMPLING FROM BINS

3.1 A sample shall be taken by passing a sampling device through the entire cross-section of the flow of material as it is being discharged (see Figures 2 and 3). Sufficient material shall be allowed to pass at the beginning of discharge to ensure uniformity of material before the sample is taken. Repeat sampling procedure as necessary until the desired amount of material from each bin is obtained. Material from each bin shall be properly identified.

4. SAMPLING FROM A CONVEYOR BELT

4.1 Sampling from a conveyor belt may be performed either while the conveyor belt is running (by using a sampling device which diverts or intercepts the flow of material) or by taking a sample while the conveyor belt is stopped. The stopped belt method is also used when approving a sampling device used for sampling while the belt is running.
4.1.1 If the sample is obtained while the conveyor belt is running, samples of the aggregate shall be taken utilizing a sampling device to divert or intercept the entire flow of material in such a manner that all portions of the flow are diverted or intercepted for an equal amount of time.

4.1.2 Samples may be obtained by stopping the conveyor belt and sampling the full width of the belt utilizing a template which is shaped to the same contour of the belt. All material which is within the template area shall be removed, utilizing a brush to obtain all the fine aggregate material.

5. **SAMPLING FROM A WINDROW**

5.1 Figure 4 illustrates the method used to sample a windrow. At each point in the windrow where a sample is to be taken, remove sufficient material from the top of the windrow so that a representative sample can be obtained from the center of the freshly exposed top of the windrow using a square point shovel. The sample taken at each sampling location shall be one shovelful of material. Repeat the sampling as necessary, at the required number of locations in the windrow, to obtain the desired amount of material. The samples taken shall be combined.

6. **SAMPLING FROM THE ROADWAY**

6.1 In the case of sampling material in-place from the roadway, at least 3 samples shall be taken with a shovel at equally distributed locations across the width of the roadway. It may be necessary to use a hammer and chisel or similar tools to cut the hole in the compacted roadway. Care shall be taken to obtain all material from the hole which is dug. The samples taken shall be combined.

7. **REDUCING FIELD SAMPLES TO TESTING SIZE**

7.1 The reduction of samples to obtain the amount required for particular tests shall be performed in accordance with AASHTO R 76.
8. SAMPLE IDENTIFICATION

8.1 Each sample shall be identified by an accompanying sample ticket. Sample tickets shall be filled out as required to provide necessary information. The remarks area of the sample ticket shall be used as necessary to provide additional information, including the phone number of an individual who can be contacted regarding the sample.

8.2 The source of the sample shall be the “original source” of the material, as indicated on the sample ticket.

8.3 An example of a completed sample ticket used by ADOT for construction projects is shown in Figure 5. Commonly used codes for filling out the sample ticket are shown on the back side of the sample ticket (see Figure 6).

8.4 The sample ticket consists of three copies. The center copy in kept by the person submitting the sample, the original copy is included inside the sample container, and the third copy is attached to the sample container. When filling out sample tickets, make certain information is clear and easily read on all three copies.
Illustration of Sampling Locations for Different Stockpile Types

Figure 1
When aggregate is passed over a screen, the fines tend to drop through immediately and accumulate on one side of the hopper. A sample taken with a shovel or other small container will not be representative.

A sample taken by inserting the sampling device through the full flow of material will yield a representative sample. The restricted opening prevents the sampling device from filling all at once.

Illustration of Bin Sampling

FIGURE 2
Typical Bin Sampler

FIGURE 3

Undisturbed Windrow

Sampling Location (Center of freshly exposed top of windrow)

Sufficient Amount of Material Removed to Obtain a Representative Sample

Illustration of Sampling From a Windrow

FIGURE 4
SAMPLING HYDRATED LIME & LIME PRODUCTS

Scope

1. These methods are intended to outline procedures for sampling of hydrated lime and lime products in powdered form. In instances where a specific case is not covered, ASTM Designation C-50 shall be referred to.

Sampling

2. (a) Sampling from Bulk Shipment at Point of Discharge to Storage. — As the material is being dropped from the truck or car into the silo entrance at ground level, take samples with a scoop or small shovel along the side of the discharge pipe. Take enough increments of material to completely fill a one-gallon can or an equivalent container which will hold at least 5 pounds.

(b) In sampling material that is placed in silos by air pressure through a system of pipes, open the bleed-valve, which is at ground level, and divert the flow of material into a one-gallon can or other suitable container. After the can has been filled, or a 5-pound sample has been discharged, close the bleed-valve.

(c) In all Other Cases Not Covered. — If the above methods are not applicable, material shall be taken from the most convenient location, provided the sample obtained will be representative of the total quantity. Increments shall be taken with a scoop or small shovel, and from at least 3 random locations within the bulk.

Quantity Required

3. All samples shall be collected in quantity enough to fill a \(\frac{1}{2}\)-gallon, air-tight metal container. This is equivalent to approximately 5 pounds. For quicklime, the containers shall be made of glass.

Identification

4. (a) Each sample container shall have legibly marked thereon the name of the product, the name of the manufacturer, place of manufacture, and the brand name, if any.

(b) The sample card shall have the following information:

(1) Name of person sampling
(2) Name and designation of project
(3) Quantity represented
(4) Date sampled
(5) Manufacturer’s name, address
SAMPLING METALLIC MATERIALS

Scope

1. These methods are intended for use in sampling the respective metallic materials named.

Sampling Bronze Bearing Plates

2. Samples shall be not less than 6 inches in length and 1½ inch in diameter, cast as an integral part of the plate. At least one sample shall be taken, at the point of manufacture, for each melt.

Sampling Castings

3. (a) Cast Iron Castings. — Two samples, not less than 1.20 inch in diameter and 21 inches in length, shall be cast from each melt. These samples are ordinarily obtained at the time of manufacture. All castings as received shall be sounded by means of a hammer to detect possible flaws.

(b) Steel Castings. — Two samples not less than 1¼ inch in diameter and 10 inches in length shall be obtained from each melt at the time of manufacture. All castings as received shall be sounded with a hammer to detect possible flaws.

Sampling Fence Stays (Metal)

4. One random sample shall be cut that is representative of 10 bundles.

Sampling Galvanized Sheet Metal

5. One sample at least 6 inches x 6 inches shall be cut in such a manner as not to disturb or damage the coating. The area in which it is taken shall not be damaged.

Sampling Metal Pipe Products

6. (a) Bituminous Coated Corrugated Metal Pipe

   (1) Field Inspection. — Whenever the pipe is shipped from a source outside the Phoenix area, one sample at least 6 inches x 6 inches shall be taken and submitted for testing.

   (2) Central Laboratory Inspection. — One sample shall be obtained from each supplier each time a dipping tank is replenished, but no less than one sample per year shall be taken.

   (3) Samples shall be taken in an area where the coating has not been disturbed, and in such a manner as not to disturb the coating.

(b) Other Corrugated Metal Pipe

   (1) Field Inspection. — Field inspection shall be made before the pipe is installed. This shall consist of the examination of the pipe to determine that the workmanship and fabrication are in accordance with specifications and that the pipe and galvanized coating have not been damaged.

   (2) Central Laboratory Inspection. — The central laboratory shall obtain and test one sample of each size pipe at least once a year. The samples shall be sent by each supplier. Sample size shall be at least 6 inches x 6 inches.

(c) Cast Iron Pipe

   (1) One piece approximately 16 inches long shall be taken from each size pipe. It shall be taken at least 6 inches from the end of a section.
Sampling Metal Posts

7. Both steel guide posts and fence posts shall be represented by one post selected at random from each project.

Sampling Reinforcing Steel

8. (a) Specifications of the State Highway Department call for one sample at least 4 feet long, selected at random from each shipment.

Sampling Steel Strands, Wire, Anchorages

9. (a) Steel strand is composed of seven wires having a center wire enclosed tightly by 6 helically-wrapped outer wires, forming a single strand for pre-tensioning in prestressed concrete. Samples shall be cut directly from the reel. Discard the first 4 feet, and obtain one sample at least 4 feet long.

   (b) Steel wire is uncoated stress-relieved wire used for post-tensioning of prestressed concrete. The manner of obtaining samples shall be identical to 11(a).

   (c) Anchorages are wedge-type devices used to anchor post-tensioning wire assemblies. Samples shall be selected at random from a shipment.

Sampling Structural Steel

10. All required sampling of structural steel is performed by the Bridge Division.

Sampling Wire

11. (a) Barbed Wire. — Samples of barbed wire shall be taken directly from the spool. The first 3 feet of wire from the spool shall be discarded and one sample at least 3 feet long shall be obtained. The spool of wire shall be selected at random.

   (b) Wire Mesh. — Wire mesh samples shall be selected at random. Each sample shall have dimensions of at least 2 feet x 2 feet.
SAMPLING MISCELLANEOUS MATERIALS

Scope

1. These methods are intended as procedures for sampling those items not found in the preceding sections of sampling.

Sampling Brick

2. For the purpose of tests, full size brick shall be selected. Specimens shall be representative of the whole lot of brick from they are selected and shall include specimens representative of the complete range of colors and sizes of the brick in the shipment. If the brick shows variation in hardness or appearance, each type shall be selected in proportion to the quantity of each type in the lot.

Sampling Clay Pipe

3. One full-sized specimen or joint shall be selected at random from every 100 pipes, or fraction thereof, of each size.

Sampling Fence Stays (Wood)

4. These shall be inspected in the field and 1 copy of a written report stating that they comply with specifications shall be submitted to the Materials Division.

Sampling Glass Beads for Paints

5. When glass beads are furnished in cans or other types of individual containers, one container shall be selected at random, and a 5-pound sample shall be selected by the quartering or riffling methods used for sampling aggregate (ARIZ 105).

Sampling Paints and Solvents

6. (a) Aluminum pigment and vehicle for aluminum paint are shipped unmixed and, therefore, shall be sampled separately.

(b) A single package out of each batch shall be taken at random as a representative of the whole. In case the material is packaged in large containers, such as barrels or tanks, a number of small samples of not more than one quart shall be taken from the top, bottom, and intermediate points by means of a sampling tube. These small samples shall then be mixed to form a composite sample of not less than one quart. This sample shall be transferred to a clean glass or tin container and securely sealed.

Sampling Liquid Membrane Curing Compound

7. The compound to be sampled shall be shaken or stirred thoroughly in the container prior to sampling until both pigment and vehicle are blended to the extent that a representative sample can be obtained. The sample shall be placed in a ½-gallon container and sealed airtight.

Sampling Paper

8. A representative sample approximately 1 foot by 2 feet shall be cut from the roll or sheet. This sampling shall apply to Asbestos, White Top, Insulating, and Roofing Paper.

Sampling Treated Wood Pilings & Timbers

9. These materials will usually be inspected and sampled at the treating plant, in which case no sampling is required. If this is not the case, a piece 18 inches to 24 inches long shall be cut from one end of a piling or timber. If no laboratory reports have been received, it is advisable to determine from the Materials Division if sampling is necessary.
Sampling Wood Fence Posts

10. This includes guide, guard, and line fence posts. These shall be inspected in the field and reports sent to the Materials Division. Samples representative of any doubtful posts shall be submitted to the central laboratory for testing.

Sampling Reflector Buttons

11. One sample of 10 reflector buttons shall be selected at random from each lot.

Sampling Water for Concrete

12. (a) No sample of water is necessary if the water is potable and comes from a proven source; however, a memo to this effect must be submitted to the Materials Division.

(b) If possible, the sample shall be taken from a faucet connected to the pipeline used for the specified project, and water shall be pumped. Allow water to run at a high enough rate to discharge any sediment or other impurities from the pipe. Place a clean, 1-gallon jar beneath the running water and fill the jar completely.

Sampling Liquid Admixtures for Concrete

13. (a) Liquid material shall be submitted in a ¾-pint clean glass container. They shall be agitated thoroughly immediately prior to taking the sample. The liquid shall be drawn from one barrel selected at random out of the lot. The sample should be drawn from a depth sufficient enough to get proportionate quantities of components.

(b) The trade name of the agent must be noted on the sample card.

Sampling Creosote, Creosote-Coal Tar, and Creosote-Fuel Oil Solutions

14. (a) Because some of the creosote may crystallize below a temperature of 120°F, the creosote shall be heated to a temperature above 120°F before sampling. Because of separation of the lighter-weight water, the creosote must be thoroughly agitated, and three samples must be taken at different levels and combined to form a composite.

(b) At least ½ gallon of material shall be taken and sealed in an airtight container.

Sampling Preformed Expansion Joint Filler

15. Each sample shall be at least 6 inches by 18 inches, and enough samples shall be taken to represent each thickness in the lot. Samples shall be packed for transportation in such a manner that there will be no danger of distortion or breakage. No tests shall be made on samples deformed or broken in handling.

Sampling Concrete Pipe, Reinforced and Plain

16. The project for which the pipe is to be used shall be notified of the dates when the pipe has been approved, or the pipe may be stenciled “Inspected — AHD”. If neither of these has been accomplished, this division shall be notified immediately, and the pipe in question shall not be used until so approved.
SERIES 200
SOILS AND AGGREGATES
1. **SCOPE**

1.1 This procedure describes the method for sieving and determining the sieve analysis of fine and coarse graded soils and aggregates, including the determination of minus No. 200 material by elutriation.

1.2 The procedure for sample preparation, sieving, and calculating the sieve analysis which is given in Sections 3 through 10 applies in general for all sieving operations. Section 11 gives a brief outline of the procedure for performing the sieve analysis when the sample is dried to constant weight prior to sieving. Section 12 gives an outline and description of the procedure for performing the sieve analysis when the sample is not dried to constant weight prior to sieving. Additional methods are given in Arizona Test Method 248, “Alternate Procedures for Sieving of Coarse and Fine Graded Soils and Aggregates”.

1.3 A washed gradation, utilizing an appropriate alternate procedure as described in either Alternate #1, 3, 4, or 5 of ARIZ 248, is to be used for all soil and aggregate materials with specification requirements for gradation. The washing requirement may be waived if the Engineer determines that it is unnecessary. However, in cases of dispute, the referee method shall be a washed gradation. Composited samples of mineral aggregate for bituminous mix designs shall be tested in accordance with either Alternate #3, 4, or 5 of ARIZ 248. If desired, the washed gradation of samples from individual stockpiles or bins may be determined and used in calculating the composite gradation.

1.4 This test method may involve hazardous materials, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.
2. **APPARATUS**

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Sieves of sizes as required for screening, conforming to the requirements of ASTM E11.

2.3 Any mechanical shaker may be used which produces the required thoroughness of sieving, as specified in Subsection 4.5.

2.4 Balances or scales - Conforming to the requirements of AASHTO M 231, except when determining weights of 5000 grams or less, the readability and sensitivity shall be at least one gram; and when determining weights of greater than 5000 grams, the readability and sensitivity shall be at least 20 grams.

2.5 Sample splitters - Shall conform to the requirements of, and be used in accordance with the procedures given in, AASHTO R 76.

2.6 Heating/Drying Device - An oven or suitable heating device which is capable of drying samples without aggregate breakage or loss of material due to splattering. A microwave oven may be used to dry materials, provided proper attention is given to the use of apparatus and the intensity of heat generated.

2.7 Brass wire brush for cleaning fine sieves coarser than No. 100, and a soft bristle brush for sieves No. 100 and No. 200.

2.8 A suitable mechanical washing device with No. 200 mesh wire (Similar to Figure 1) for performing washing of fine soils and aggregates as described in Section 6, equipped with a metal tube(s) connected to a water and an air outlet; or a vessel of sufficient capacity and a nest of No. 16 and No. 200 sieves for use with the hand washing method. When utilizing mechanical washing equipment, the provision for air may be eliminated if the water pressure provides adequate agitation to wash sample as specified in Section 6.

2.9 Miscellaneous pans, scoops, spatulas, brushes, pulverizing equipment, etc. for preparing, washing, and drying samples.
3. SAMPLE PREPARATION

3.1 A representative sample of the amount indicated in Subsection 3.2 below shall be obtained. Dry the sample sufficiently to permit separation of particles on the No. 4 and larger sieves, and to develop a free-flowing condition in the portion passing the No. 4 sieve. The use of sunlight, ovens, fans, or warm air are the most common drying methods. Turn the sample frequently to prevent formation of hard clay lumps. If the sample contains hard clods or coated coarse aggregate particles, break up the clods by means which will not reduce the size of any rock. (The use of a rubber covered mallet or the raking of the material over an old screen have been found to be satisfactory methods.) When subsequent testing is to be performed on the material, it shall be assured that any temperature restrictions, as given in the appropriate test method(s), are not exceeded. A number of soil and aggregate tests, such as plasticity index, proctor maximum density, and sand equivalent, require that samples not be heated in excess of 140 °F.

3.2 If necessary, samples shall be reduced in size in accordance with the splitting or quartering methods in AASHTO R 76 to obtain the appropriate sample size as shown below.

<table>
<thead>
<tr>
<th>Nominal Maximum Size of Particle *</th>
<th>Minimum Weight of Sample, grams (lbs.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/8”</td>
<td>1000 (2.2)</td>
</tr>
<tr>
<td>1/2”</td>
<td>2000 (4.4)</td>
</tr>
<tr>
<td>3/4”</td>
<td>5000 (11)</td>
</tr>
<tr>
<td>1”</td>
<td>10000 (22)</td>
</tr>
<tr>
<td>1-1/2”</td>
<td>15000 (33)</td>
</tr>
<tr>
<td>2”</td>
<td>20000 (44)</td>
</tr>
<tr>
<td>2-1/2”</td>
<td>25000 (55)</td>
</tr>
<tr>
<td>3”</td>
<td>30000 (66)</td>
</tr>
<tr>
<td>3” Slot</td>
<td>35000 (77)</td>
</tr>
</tbody>
</table>

*The smallest sieve opening through which the entire amount of material, by specification, is permitted to pass.

3.3 If desired, the sample may be dried to constant weight. The sample may be considered to be at constant weight when, after an initial drying period of at least one hour, further drying causes or would cause less than 0.1% additional loss in weight within a five minute drying period. If desired, the material may be dried to constant weight utilizing a microwave oven in accordance with Subsection 2.6.
3.4 Allow the sample to cool, if necessary, and record the weight of the material.

4. **COARSE SIEVING**

4.1 Normally the coarse sieving of material is performed utilizing the No. 4 sieve as the smallest sieve.

4.2 When material being screened contains a large amount of passing No. 4 material, such that there may be overloading in the bottom pan during shaking operations, the sample shall be initially sieved over a No. 4 sieve to separate excess amounts of this material. This passing No. 4 material is saved and combined with the remaining portion of passing No. 4 material from subsequent coarse sieving. When material contains large rocks that are not to be sieved in the nest of coarse sieves during the actual shaking operation, the sample shall be pre-sieved to separate these particles. Unless all material will pass the largest sieve in the nested sieves, the material shall be pre-sieved over a sieve of the next larger size than that of the largest sieve size being utilized in the nested sieves. Large rocks separated in this manner are individually tested for passing the appropriate sieve, determining and recording the weight of any rock retained on these sieves.

4.3 Empty the sample into the nest of sieves that is to be used for screening material. If pre-sieving has not been performed, remove any particles which may be retained on the top sieve and test these for passing the appropriate sieve, determining and recording the weight of any rock retained on these sieves.

4.4 The material shall be subjected to sieving by hand or in a mechanical sieve shaker. The sieving action shall cause the particles to bounce and turn so as to present different orientations to the sieving surface. No particles shall be hand manipulated for passing any of the nested sieves. The sieving shall be of sufficient time to assure that the criteria for “thoroughness of sieving” described in Subsection 4.5 below is achieved.

4.5 The criteria for "thoroughness of sieving" is that after completion of sieving, not more than 0.5 percent by weight of the total sample passes any sieve during one minute of continuous hand sieving in an 8 inch or 12 inch sieve as appropriate. If the thoroughness of sieving is being determined for sieves larger than 12 inches, the material retained on the respective sieve, or portions of that material shall be placed in a 12 inch sieve of the same sieve size opening, so as to not overload the sieve. Hold the individual sieve, provided with a snug fitting pan and cover,
in a slightly inclined position in one hand. Strike 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, turn the sieve about one-sixth of a revolution at intervals of about 25 strokes. A frequent check should be performed to assure that the thoroughness of sieving specified is being maintained in sieving operations.

4.6 Overloading of sieves shall be avoided. The quantity of material on a given sieve at the completion of sieving shall not exceed the amount shown in the table below. As the weight retained on each individual sieve size is determined, it should be compared with the corresponding overloading criteria prior to combining the weighed material. If overloading occurs, proper corrective action to regulate the amount of material on a sieve shall be taken. This may be accomplished by splitting the material which is retained on the overloaded sieve, resieving each portion, and combining weights.

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Maximum Weight Allowed (grams/sq. in.)</th>
<th>Maximum Weight Allowed (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>12” Dia. Sieve</td>
</tr>
<tr>
<td>3”</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>2-1/2”</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>2”</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>1-1/2”</td>
<td>25</td>
<td>2827</td>
</tr>
<tr>
<td>1”</td>
<td>18</td>
<td>2036</td>
</tr>
<tr>
<td>3/4”</td>
<td>14</td>
<td>1583</td>
</tr>
<tr>
<td>1/2”</td>
<td>10</td>
<td>1131</td>
</tr>
<tr>
<td>3/8”</td>
<td>8</td>
<td>905</td>
</tr>
<tr>
<td>1/4”</td>
<td>6</td>
<td>679</td>
</tr>
<tr>
<td>No. 4</td>
<td>5</td>
<td>565</td>
</tr>
</tbody>
</table>

*Normally particles of material retained on these sieves are tested individually, so no maximum weight allowed is specified.

4.7 Starting with the largest of the nested sieves, the material retained on each sieve and in the bottom pan is weighed and recorded.

4.8 Do not discard any of the sieved material until the sum of the individual weights is compared to the weight of sample prior to sieving. If the difference between the two weights is less than or equal to 1.0% of the weight of sample prior to sieving, an adjustment in weight shall be made on the sieve which has the largest
weight retained, except no adjustments shall be made on the minus No. 4 material. If the difference is greater than 1.0%, the sample shall be recombined, resieved, and carefully reweighed.

5. SAMPLE FOR ELUTRIATION AND FINE SIEVING

5.1 A representative minimum 500 gram sample of the passing No. 4 material from the coarse sieving shall be obtained by the use of a splitter in accordance with AASHTO R 76. The selection of an exact predetermined weight shall not be attempted. The sample size may be reduced if the minimum of 500 grams is not obtained from coarse sieving, or if there is insufficient material for performing other desired tests utilizing the pass No. 4 material. When utilizing the mechanical washing device, the requirement for a minimum 500 gram sample may be reduced to a minimum of 200 grams for materials that tend to clog the No. 200 sieve (for example, fine soils such as blow sand, silty soils, or clay).

5.2 The weight of the sample for elutriation and fine sieving is recorded to the nearest gram as “Dry Wt. Pass No. 4 Split”.

5.3 Subject the sample to elutriation by either mechanical washing (Section 6) or hand washing (Section 7).

6. MECHANICAL WASHING

6.1 This method is generally utilized for passing No. 4 material, although it may also be used with the alternate procedure described in Section 2 of Arizona Test Method 248 for testing materials containing small amounts of Plus No. 4 material (100% passing 3/8"). Generally, a maximum of 600 grams may be tested utilizing the mechanical washing device.

6.2 Fill the washing device with water to the bottom of the windows. Transfer the sample to the washing device, and wash all material clinging to the sample container into the washing device. Utilizing the water tube, and air tube if necessary, agitate the sample vigorously enough so that it causes the material to go into suspension, but not enough to cause splattering to reach the top of the cylinder. Continue this washing action, taking care that there is no loss of sample by splattering or overflowing. (If the No. 200 sieve tends to plug excessively, this may be overcome to some extent by tapping the washing device with the palm of the hand or by washing down the inside of the sieve with a low stream of water.) Washing shall continue until the wash water becomes clear.
6.3 Turn off the water and compressed air and remove the metal tube(s), rinsing clinging material back into the sample. Wash the sample into a container of sufficient capacity to hold the water and sample; allow the particles to settle and decant the excess water.

6.4 Dry the sample to constant weight at a temperature that will not cause the material to be lost due to splattering.

6.5 Allow the sample to cool, reweigh and record to the nearest gram as the fine sieve “Total Dry Weight”.

6.6 Subtract the “Total Dry Weight” from the “Dry Wt. of Pass No. 4 Split”, and record as the “Elutriation”.

7. HAND WASHING (REFEREE METHOD)

7.1 Place the sample in a pan of sufficient size and capacity to allow washing without spillage. Cover with adequate water to thoroughly wash aggregate. Agitate the contents of the pan vigorously in order to completely separate all particles finer than the No. 200 sieve from the coarser particles, and to bring the finer material into suspension so that it will be removed by decantation of the water.

7.2 Decant the wash water through a nest of No. 16 and No. 200 sieves.

7.3 Repeat the washing and decanting cycle until the water becomes clear.

7.4 Thoroughly wash all material remaining on the No. 16 and the No. 200 sieves and return to the sample.

7.5 After the particles have settled in the pan, carefully decant any excess water, assuring that no particles are lost.

7.6 Dry the sample to constant weight at a temperature that will not cause material to be lost due to splattering.

7.7 Allow the sample to cool, reweigh and record to the nearest gram as the fine sieve “Total Dry Weight”.

7.8 Subtract the “Total Dry Weight” from the “Dry Wt. of Pass No. 4 Split”, and record as the “Elutriation”.
8. SIEVING OF FINE AGGREGATE

8.1 Place the washed and dried fine aggregate sample into the top of the nested sieves, close the nest of sieves with lid.

8.2 The material shall be subjected to sieving by hand or in a mechanical sieve shaker. The sieving action shall cause the particles to bounce and turn so as to present different orientations to the sieving surface. No particles shall be hand manipulated for passing any of the nested sieves. The sieving shall be of sufficient time to assure that the criteria for "thoroughness of sieving" described in Subsection 4.5 is achieved.

8.3 Overloading of sieves shall be avoided. The quantity of material on a given sieve at the completion of sieving shall not exceed 4 grams per square inch of sieving area (201 grams for an 8 inch diameter sieve and 452 grams for a 12 inch sieve). As the weight retained on each individual sieve size is determined, it should be compared with the corresponding overloading criteria prior to combining the weighed material. If overloading occurs, proper corrective action to regulate the amount of material on a sieve shall be taken. This may be accomplished by splitting the material which is retained on the overloaded sieve, resieving each portion, and combining weights.

8.4 Starting with the largest of the nested sieves, the material retained on the individual sieves and in the bottom pan shall be weighed and recorded.

8.5 Do not discard any of the sieved material until the sum of the individual weights is compared to the weight of sample prior to sieving. If the difference between the two weights is less than or equal to 1.0% of the weight of sample prior to sieving, an adjustment in weight shall be made on the sieve which has the largest weight retained, except no adjustments shall be made on the minus No. 200 material. If the difference is greater than 1.0% the sample shall be recombined, resieved, and carefully reweighed.

9. PRECAUTIONS

9.1 Check sieves at least daily for broken or distorted wire, and replace any sieves found to be damaged or excessively worn.

9.2 Sieves not conforming to AASHTO M 92 must be replaced.
9.3  Do not repair wire cloth.

9.4  Clean sieves carefully after each shaking, using the proper instrument to reduce chances of damaging the mesh.

9.5  All mechanical equipment shall be inspected frequently and maintained by greasing, cleaning, and repair of worn out parts.

10.  CALCULATIONS

10.1  The calculations for determining the sieve analysis are as follows. Examples of these calculations are given in Figures 2 and 3.

10.2  The calculations to determine the % passing values for the coarse sieve analysis are performed as described below:

10.2.1  For the largest sieve which has no material retained, record the percent passing as 100. Determine a factor for calculating the coarse sieve analysis by the following:

\[
\text{Coarse Sieve Factor} = \frac{100}{\text{Coarse Sieve Total}}
\]

10.2.2  The percent passing for each sieve in the coarse sieve analysis is determined by multiplying the weight retained on that sieve times the coarse sieve factor, and subtracting the result from the unrounded % passing for the next larger sieve, as shown below. Values for “weight retained times the coarse sieve factor” and “percent passing each sieve” shall be determined and used in the calculations to at least six decimal places. The percent passing value for each sieve is recorded in the sieve analysis to the nearest whole percent.

\[
\text{Percent Passing} = \left( \frac{\text{Unrounded} \% \text{Pass for next larger sieve}}{\text{Coarse Sieve Factor}} \right) - \left( \frac{\text{Weight Retained on Desired Sieve}}{\text{Sieve Factor}} \right)
\]
10.2.3 As a check on the coarse sieve analysis, multiply the weight of minus No. 4 material times the coarse sieve factor, as shown below. The result of this calculation, rounded to the nearest whole percent, should be the same as the value for percent passing the No. 4 sieve determined in the paragraph above.

\[
\text{Check for Percent Passing No. 4} = \left( \frac{\text{Wt. of Pass No. 4 sieve}}{\text{Coarse Sieve Factor}} \right)
\]

10.3 The calculations to determine the % passing values for the fine sieve analysis are performed as described below:

10.3.1 Determine a factor for the fine sieve analysis by dividing the percent passing the No. 4 sieve (which has been recorded to the nearest whole percent) by the "Dry Weight of Pass #4 Split", as shown below. Record the fine sieve factor to at least six decimal places. If all the pass No. 4 material from coarse sieving was subjected to elutriation and fine sieving, a fine sieve factor is not determined. Rather, the coarse sieve factor is utilized and the calculation of the percent passing each sieve is continuous through the entire sieve analysis.

\[
\text{Fine Sieve Factor} = \frac{\text{Rounded Percent Pass No. 4}}{\text{Dry Wt. of Pass No. 4 Split}}
\]

10.3.2 The percent passing for each sieve in the fine sieve analysis is determined by multiplying the weight retained on that sieve times the fine sieve factor, and subtracting the result from the unrounded % passing the next larger sieve, with the exception of the percent passing the No. 4 which has previously been recorded to the nearest whole percent. The equation for determining the percent passing each sieve is shown below. Values for "weight retained times the fine sieve factor" and "percent passing each sieve" shall be determined and used in the calculations to at least six decimal places. The percent passing value for each sieve is recorded in the sieve analysis to the nearest whole percent, except the percent passing the No. 200 sieve is recorded to the nearest 0.1 percent.

\[
\text{Percent Passing Desired Sieve} = \left( \frac{\text{Unrounded \% Pass for next Larger sieve}}{\text{Weight Retained on Desired Sieve \times Fine Sieve Factor}} \right) - \left( \frac{\text{Weight Retained on Desired Sieve}}{\text{Fine Sieve Factor}} \right)
\]
10.3.3 As a check on the fine sieve analysis, the weight of material passing the No. 200 from sieving is added to the elutriation weight, and this total is multiplied times the fine sieve factor, as shown below. The result of this calculation, rounded to the nearest 0.1 percent, should be the same as the value for the percent passing the No. 200 determined in the paragraph above.

\[
\text{Check for Percent Passing No. 200} = \left( \frac{\text{Weight of Passing No. 200} + \text{Elutriation Weight}}{\text{Weight}} \right) \times \left( \frac{\text{Fine Sieve Factor}}{} \right)
\]

10.4 If desired, obtain the percent retained on each sieve by subtracting the rounded % passing value for that sieve from the rounded % passing value for the next larger sieve, as shown below.

\[
\% \text{Retained for Desired Sieve} = \left( \frac{\text{Rounded} \%}{\text{Pass Next Larger Sieve}} \right) - \left( \frac{\text{Rounded} \%}{\text{Pass the Desired Sieve}} \right)
\]

10.5 Other methods may be used that differ from that specified in Subsections 10.2 and 10.3 above to determine % passing each sieve, so long as the method utilized has been proven to give equivalent results. However, any procedure which includes recording percent retained values prior to completing the calculation of all percent passing values is not allowed.

11. **PROCEDURE WHEN SAMPLE IS DRIED TO CONSTANT WEIGHT PRIOR TO SIEVING**

11.1 The following is a brief outline of the procedure to be used when the sample is dried to constant weight prior to sieving. An example of this procedure is shown in Figure 2.

11.2 Prepare sample, dry to constant weight, allow to cool, and record as “Coarse Sieve Total”, (3087 in the example).

11.3 Perform coarse sieving.

11.4 Record the weight of the material in the appropriate “Weights Retained” box for each sieve and the pan.
11.5 Sum the individual weights retained for each sieve, compare to the weight of sample prior to sieving (Coarse Sieve Total), and adjust or resieve as necessary.

11.6 Split the pass No. 4 material to at least 500 grams and record as "Dry Wt. of Pass No. 4 Split", (533).

11.7 Perform elutriation on the dry - #4 split, dry back to constant weight, allow to cool, and record as the "Total Dry Weight", (491).
### Example ECR Procedure When Sample Is Dried

TO CONSTANTWEIGHT PRIOR TO SIEVING - (SECTION 11)

**CONTACT PHONENO.** - (555)555-5555

<table>
<thead>
<tr>
<th>Test No.</th>
<th>Lot or Suffix</th>
<th>Sampled By</th>
<th>Date</th>
<th>Year</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>17</td>
<td>A. Jones</td>
<td>11/16/14</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cold Feed</th>
<th>Original Source</th>
<th>Project Engineer / Supervisor</th>
<th>Project Number</th>
<th>Trans Number</th>
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<tbody>
<tr>
<td>CM # 4444</td>
<td>B. Smith</td>
<td></td>
<td>F-099-9 (93)</td>
<td>H999901C</td>
</tr>
</tbody>
</table>

**Example Table:**

<table>
<thead>
<tr>
<th>Coarse Factor</th>
<th>Coarse Sieve Total</th>
<th>Cumulative</th>
<th>% Ret.</th>
<th>% Pass</th>
<th>Spec.</th>
<th>Fineness Modulus</th>
<th>Spec.</th>
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<tbody>
<tr>
<td>Coarse Factor</td>
<td></td>
<td></td>
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<td></td>
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</tr>
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<table>
<thead>
<tr>
<th>Wet Sample Preweight</th>
<th>Wet Wt. of #4 Split</th>
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<table>
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<tr>
<th>Weight Retained</th>
<th>% Ret.</th>
<th>% Pass</th>
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<tbody>
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<table>
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<th>Year</th>
<th>Time</th>
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<tr>
<td>17</td>
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<table>
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<th>Trans Number</th>
</tr>
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<tbody>
<tr>
<td>CM # 4444</td>
<td>B. Smith</td>
<td></td>
<td>F-099-9 (93)</td>
<td>H999901C</td>
</tr>
</tbody>
</table>

**Figure 2**
11.8 Determine and record "Elutriation" by subtracting the "Total Dry Weight" from the "Dry Wt. of Pass No. 4 Split", (533 - 491 = 42).

11.9 Perform fine sieving on the material left from elutriation using the No. 8 sieve down to the No. 200.

11.10 Record the weight of material in the appropriate “Weights Retained" box for each sieve.

11.11 Sum the individual weights retained for each sieve, compare to the weight of sample prior to sieving (fine sieve Total Dry Weight), and adjust or resieve as needed.

11.12 Determine the sieve analysis of the material as described in Section 10.

12. PROCEDURE WHEN SAMPLE IS NOT DRIED TO CONSTANT WEIGHT PRIOR TO SIEVING

12.1 The following is an outline and description of the procedure to be used when the sample is not dried to constant weight prior to sieving. An example of this procedure is shown in Figure 3.

12.2 Prepare sample, dry to free flowing condition, and record this weight as “Wet Sample Preweight”, (13010 in the example).

12.3 Perform coarse sieving.

12.4 Record the weight of the material in the appropriate "Weights Retained" box for each sieve, except record the weight of pass No. 4 material as the "Wet Wt. of - #4", (7365).

12.5 Sum the individual weights retained for each sieve, compare to weight of sample prior to sieving (Wet Sample Preweight), and adjust or resieve as needed.

12.6 The wet weight of pass No. 4 material is corrected for moisture using either a split of, or the entire amount of, the pass No. 4 material.
### Example for Procedure When Sample is Not Dried

TO CONSTANTWEIGHT PRIOR TO SIEVING - (SECTION 12)

#### CONTACT PHONE NO. - (555)555-5555

<table>
<thead>
<tr>
<th>REMARKS</th>
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<tr>
<td>Liquid Limit (LL) T-89</td>
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<tr>
<td>Plastic Limit (PL) T-100</td>
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<td>Plasticity Index (PI) = LL - PL T-90</td>
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<tr>
<td>@ 100 Revisions</td>
</tr>
<tr>
<td>@ 500 Revisions</td>
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#### Absorption, H/O

<table>
<thead>
<tr>
<th>ARIZ 248</th>
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</thead>
<tbody>
<tr>
<td>Specific Gravity, SSD ARIZ213</td>
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<tr>
<td>Specific Gravity, CD ARIZ213+</td>
</tr>
<tr>
<td>Specific Gravity, Apparent ARIZ213+</td>
</tr>
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</table>

#### Proctor Method

<table>
<thead>
<tr>
<th>ARIZ 248</th>
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<tbody>
<tr>
<td>Optimum Moisture +%</td>
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<tr>
<td>Maximum Density +PCF</td>
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</table>

#### Sand Equivalent

<table>
<thead>
<tr>
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</tr>
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<tbody>
<tr>
<td>At Least One Incrusted Face ARIZ 212</td>
</tr>
<tr>
<td>At Least Two Incrusted Faces ARIZ 212</td>
</tr>
</tbody>
</table>

#### Uncompacted Void Content

<table>
<thead>
<tr>
<th>ARIZ 248</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture Content +%</td>
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</tbody>
</table>

#### Flakiness Index

<table>
<thead>
<tr>
<th>ARIZ 233</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonates ARIZ 238</td>
</tr>
</tbody>
</table>

#### pH

| ARIZ 239 OR 237 +|

#### Resistivity (ohms-cm)

| ARIZ 236 |

#### Soluble Salts (PPM)

| ARIZ237 |

#### Unit Weight

| T-19 |

#### Voids

| T-19 |

#### Organic Impurities

| T-21 |

#### Chloride Content (PPM)

| ARIZ236 |

#### Sulfate Content (PPM)

| ARIZ233 |

#### Exchangeable Sodium (%)

| ARIZ279 |

#### Exchangeable Sodium (PPM)

| ARIZ279 |

#### TOTAL CUMULATIVE RET.

| 100 |

---

**FIGURE 3**
12.7 If a split of the pass No. 4 material is to be used to correct the weight of pass No. 4 material for moisture, immediately split the passing No. 4 material to at least 500 grams and record as the "#4 Split Wet Wt.", (506). Dry the sample to constant weight, allow to cool, and record the weight as "Dry Wt. of Pass #4 Split", (497). If an elutriation and fine sieve analysis is to be performed, this split is used for that testing. Determine and record the "weight of pass No. 4 material" (7234) by the following:

\[
\text{Weight of Pass No. 4 Material} = \left( \frac{\text{Wet Wt. of Pass No. 4}}{\text{Dry Wt. of Pass No. 4 Split}} \right) \times \left( \frac{\text{Pass No. 4 Split}}{\text{Wet Wt.}} \right)
\]

12.8 If the entire amount of the pass No. 4 material is to be used to correct the weight of pass No. 4 material for moisture, the material is dried to constant weight and allowed to cool. The dry weight is recorded as the "weight of pass No. 4 material". If an elutriation and fine sieve analysis is to be performed on this material, the dry weight is also recorded as the "Dry Wt. of Pass No. 4 Split".

12.9 Determine and record the sample “Coarse Sieve Total”, (12879), by the following:

\[
\text{Coarse Sieve Total} = \left( \frac{\text{Wet Sample Preweight}}{\text{Wet Wt. of Pass No. 4 Material}} \right) - \left( \frac{\text{Wet Wt. of Pass No. 4}}{\text{Pass No. 4 Split}} \right)
\]

12.10 If required, perform elutriation on the dry - #4 split, dry back to constant weight, allow to cool, and record as the "Total Dry Weight", (440).

12.11 Determine and record “Elutriation” by subtracting the “Total Dry Weight” from the “Dry Wt. of Pass No. 4 Split”, (497 - 440 = 57).

12.12 Perform fine sieving on the material left from elutriation using the No. 8 sieve down to the No. 200.

12.13 Record the weight of material in the appropriate “Weights Retained” box for each sieve.
12.14 Sum the individual weights retained for each sieve, compare to the weight of sample prior to sieving (fine sieve Total Dry Weight), and adjust or resieve as needed.

12.15 Determine the sieve analysis of the material as described in Section 10.

13. REPORT

13.1 The sieve analysis shall be reported either as shown in the example given in Figure 2 for a sample which has been dried to constant weight prior to sieving, or as shown in the example given in Figure 3 for a sample which has not been dried to constant weight prior to sieving.

13.2 A blank Soils and Aggregate Tabulation laboratory card is provided in Figure 4.
**Figure 4**

**Arizona Department of Transportation**

**Soil and Aggregate Tabulation**

<table>
<thead>
<tr>
<th>Lab Number</th>
<th>Org Number</th>
<th>Material</th>
<th>Type</th>
<th>Purpose</th>
<th>Test Lab</th>
<th>Size</th>
<th>Size %</th>
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<tbody>
<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

**Test No.**

<table>
<thead>
<tr>
<th>Lot or Suffix</th>
<th>Sampled In</th>
<th>Day</th>
<th>Month</th>
<th>Year</th>
<th>Time</th>
<th>Military Time</th>
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<td></td>
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</tbody>
</table>

**Sampled From**

<table>
<thead>
<tr>
<th>Site No.</th>
<th>Rdwy</th>
<th>Station</th>
<th>If Milepost, Input Decimal</th>
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</thead>
<tbody>
<tr>
<td></td>
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**Original Source**

<table>
<thead>
<tr>
<th>Project Engineer / Supervisor</th>
<th>Project Number</th>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Remarks**

**Contact Phone No.**

- **Liquid LIMIT (LL)**
  - T - 65
- **Plastic Limit (PL)**
  - T - 90
- **Plasticity Index (PI) = LL - PL**
  - T - 90

**Abrasion Method (A,B,C,D)**

- **100 Resolutions**
  - %
  - %
  - %
  - %
  - %

**Absorption, H2O**

- **ARIZ 201**
  - A - ARIZ 225
  - G - ARIZ 226
  - D - ARIZ 245
  - A1 - ARIZ 232
  - A2 - ARIZ 246

**Specific Gravity, SG**

- **ARIZ 201**

**Specific Gravity, OD**

- **ARIZ 201**

**Specific Gravity, Apparent**

- **ARIZ 201**

**Proctor Method**

- **Optimum Moisture**
  - %

**Max. Dry Density**

- **MDF**

**Sand Equivalent (ARIZ 248) RC**

- **ARIZ 248**

**Uncompacted void content**

- **ARIZ 247**

**Moisture Content**

- **ARIZ 252**

**At Least One Particular Face**

- **ARIZ 252**

**At Least Two Particular Faces**

- **ARIZ 252**

**Carbonates**

- **ARIZ 236**

**pH**

- **ARIZ 236 262 237**

**Resistivity (ohm-cm)**

- **ARIZ 236**

**Solute Salts (PAM)**

- **ARIZ 207**

**Unit Weight**

- **MDF**

**Voids**

- **MDF**

**Organic Impurities**

- **MDF**

**Clay Content (PPI)**

- **ARIZ 7.6**

**Sulfate Content (PPI)**

- **ARIZ 7.3**

**Exchangeable Sodium (%)**

- **ARIZ 7.9**

**Exchangeable Sodium (PPI)**

- **ARIZ 7.9**

**Total Cumulative % Ret. FINENESS MODULUS**

- **100**

**Weights Retained**

<table>
<thead>
<tr>
<th>% Ret.</th>
<th>% Pass</th>
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**Dry Wt. of #6 Split**

<table>
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<tr>
<th>% Ret.</th>
<th>% Pass #4</th>
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**Total**

<table>
<thead>
<tr>
<th>% Ret.</th>
<th>% Pass</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

**White**

**Yellow**

**Blue**

**Received Date**

**Test Operator & Date**

**Supervisor & Date**

**See Back Also**
COMPOSITE GRADING

(An Arizona Method)

Scope

1. This is a method of combining two or more samples of different gradations to produce a final product of a composited gradation.

Procedure

2. (a) For each sample; multiply the percent retained on each sieve by the "Decimal % of Composite" for the sample and record the results to the nearest 0.1%, except that the amount passing the No. 200 sieve shall be to the nearest 0.01%. Adjust the fractions of material for each aggregate sample as necessary to correspond to the total percent of composite for that sample, except that the percent passing the No. 200 sieve shall not be adjusted.

(b) Add the resultant percentages retained on each sieve for all samples and record as the composite % retained. Round the % retained for each sieve to the nearest 1%, except the amount passing the No. 200 sieve shall be rounded and recorded to the nearest 0.1%.

NOTE: Adjustments to the % retained and the % retained rounded may be necessary so that the total equals 100%, except that the % passing the No. 200 sieve shall not be adjusted.

(c) To obtain the composite percent passing each sieve, start with the Pass No. 200 fraction (rounded to the nearest whole percent), in the example 3%; and add the percent retained on the No. 200 sieve (3%), and record the sum as the % passing the No. 100 sieve (6%). Add the % retained on the No. 100 to this total and record as the % passing the No. 50 sieve (6 + 8 = 14). Repeat this procedure for all sieves, the final value should be 100%.

(d) If it is preferred, the % passing each sieve may be determined by beginning with the largest sieve which has material retained, in the example 13% was retained on the 1/2" sieve; subtract the % retained from 100 and record as % passing (100 - 13 = 87); subtract the % retained on the next smaller sieve and record as % passing (87 - 11 retained on the 3/8" = 76); continue this procedure for all sieves, the final value should be the percent passing the No. 200 sieve, rounded to the whole percent.
NOTE: Figure 1 shows an example of a sieve analysis as reported from ARIZ 201, for four individual samples. Figure 2 shows an example of the calculations described in paragraphs (a), (b), (c), and (d), and a completed composite grading of the four individual samples.

(e) If a calculated composite is desired using the percent passing each sieve, it may be obtained by multiplying the % pass each sieve by the "decimal % of composite" for each sample, and accumulate for the resultant composite percent pass each sieve in similar manner as in the method described above for composite % retained. After the composite % passing each sieve is calculated the composite % retained is determined by the following:

\[
\text{% Retained on Individual Sieve} = \text{% Passing next larger sieve size} - \text{% Passing desired sieve size}
\]

NOTE: Figure 3 shows an example of the calculations described in paragraph (e), and a completed composite grading of the four individual samples.
**Sample #1**

<table>
<thead>
<tr>
<th>% Oversize</th>
<th>Coarse Factor</th>
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<tbody>
<tr>
<td>G 3</td>
<td>015040 (no punch)</td>
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</table>

<table>
<thead>
<tr>
<th>Weights Retained</th>
<th>% Ret.</th>
<th>% Pass</th>
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<tbody>
<tr>
<td>3''</td>
<td>12</td>
<td>10</td>
</tr>
<tr>
<td>2 1/4''</td>
<td>17</td>
<td>21</td>
</tr>
<tr>
<td>2''</td>
<td>22</td>
<td>26</td>
</tr>
<tr>
<td>1 1/2''</td>
<td>27</td>
<td>31</td>
</tr>
<tr>
<td>1''</td>
<td>32</td>
<td>36</td>
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<tr>
<td>3/4''</td>
<td>37</td>
<td>41</td>
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<tr>
<td>3/8''</td>
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<td>46</td>
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<td>3/32''</td>
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<td>62</td>
<td>64</td>
</tr>
<tr>
<td>Total</td>
<td>67</td>
<td>72</td>
</tr>
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</table>

**Sample #2**

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<th>% Oversize</th>
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<td>G 3</td>
<td>039200 (no punch)</td>
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</table>

<table>
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<tr>
<th>Weights Retained</th>
<th>% Ret.</th>
<th>% Pass</th>
</tr>
</thead>
<tbody>
<tr>
<td>3''</td>
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<td>16</td>
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<td>64</td>
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<tr>
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<td>72</td>
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**Sample #3**

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<th>% Pass</th>
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</thead>
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<td>16</td>
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<td>64</td>
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<tr>
<td>Total</td>
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<td>72</td>
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</tbody>
</table>

**Sample #4**

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<tr>
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<th>Coarse Factor</th>
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<tbody>
<tr>
<td>G 3</td>
<td>067751 (no punch)</td>
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</table>

<table>
<thead>
<tr>
<th>Weights Retained</th>
<th>% Ret.</th>
<th>% Pass</th>
</tr>
</thead>
<tbody>
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<td>16</td>
</tr>
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<td>2''</td>
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<tr>
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<td>62</td>
<td>64</td>
</tr>
<tr>
<td>Total</td>
<td>67</td>
<td>72</td>
</tr>
</tbody>
</table>

---

**Figure 1**

- Table of weights and sieves for samples #1 to #4.
- Data includes % oversize, coarse factor, % retained, % pass.
- Additional columns for fine sieve factor and split.
- Total weight and percentage for each sample.
<table>
<thead>
<tr>
<th>SIEVE SIZE</th>
<th>SAMPLE #1</th>
<th>SAMPLE #2</th>
<th>SAMPLE #3</th>
<th>SAMPLE #4</th>
<th>COMPOSITE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>% OF COMPOSITE = 23</td>
<td>% OF COMPOSITE = 20</td>
<td>% OF COMPOSITE = 27</td>
<td>% OF COMPOSITE = 30</td>
<td>% Ret.</td>
</tr>
<tr>
<td>3/4</td>
<td>0</td>
<td>x.23</td>
<td>10</td>
<td>x.20</td>
<td>20</td>
</tr>
<tr>
<td>1/2</td>
<td>55</td>
<td>x.23</td>
<td>12.6</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>3/8</td>
<td>40</td>
<td>x.23</td>
<td>9.2</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>1/4</td>
<td>5</td>
<td>x.23</td>
<td>1.2</td>
<td>48</td>
<td>x.20</td>
</tr>
<tr>
<td>#4</td>
<td>0</td>
<td>x.23</td>
<td></td>
<td>27</td>
<td>x.20</td>
</tr>
<tr>
<td>#8</td>
<td>12</td>
<td>x.20</td>
<td>2.4</td>
<td>29</td>
<td>x.27</td>
</tr>
<tr>
<td>#10</td>
<td>1</td>
<td>x.20</td>
<td>0.2</td>
<td>10</td>
<td>x.27</td>
</tr>
<tr>
<td>#16</td>
<td>1</td>
<td>x.20</td>
<td>0.2</td>
<td>19</td>
<td>x.27</td>
</tr>
<tr>
<td>#30</td>
<td>0</td>
<td>x.20</td>
<td></td>
<td>12</td>
<td>x.27</td>
</tr>
<tr>
<td>#40</td>
<td>0</td>
<td>x.20</td>
<td></td>
<td>3</td>
<td>x.27</td>
</tr>
<tr>
<td>#50</td>
<td>0</td>
<td>x.20</td>
<td></td>
<td>3</td>
<td>x.27</td>
</tr>
<tr>
<td>#100</td>
<td>0</td>
<td>x.20</td>
<td></td>
<td>4</td>
<td>x.27</td>
</tr>
<tr>
<td>#200</td>
<td>0</td>
<td>x.20</td>
<td></td>
<td>3</td>
<td>x.27</td>
</tr>
<tr>
<td>Pass #200</td>
<td>0</td>
<td>x.23</td>
<td>0.8</td>
<td>x.20</td>
<td>0.16</td>
</tr>
<tr>
<td>TOTAL</td>
<td>100</td>
<td>23.0</td>
<td>99.8</td>
<td>19.96</td>
<td>99.9</td>
</tr>
</tbody>
</table>

FIGURE 2
<table>
<thead>
<tr>
<th>SIEVE SIZE</th>
<th>SAMPLE #1 % OF COMPOSITE = 23</th>
<th>SAMPLE #2 % OF COMPOSITE = 20</th>
<th>SAMPLE #4 % OF COMPOSITE = 27</th>
<th>SAMPLE #4 % OF COMPOSITE = 30</th>
<th>COMPOSITE % Pass</th>
<th>% Comp. % Pass</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4</td>
<td>100 x.23 23.0</td>
<td>100 x.20 20.0</td>
<td>100 x.27 27.0</td>
<td>100 x.30 30.0</td>
<td>100.0</td>
<td>100</td>
</tr>
<tr>
<td>1/2</td>
<td>45  x.23 10.4</td>
<td>100 x.20 20.0</td>
<td>100 x.27 27.0</td>
<td>100 x.30 30.0</td>
<td>87.4</td>
<td>13</td>
</tr>
<tr>
<td>3/8</td>
<td>5   x.23 1.2</td>
<td>90   x.20 18.0</td>
<td>100 x.27 27.0</td>
<td>100 x.30 30.0</td>
<td>76.2</td>
<td>11</td>
</tr>
<tr>
<td>1/4</td>
<td>0    x.23 0</td>
<td>42   x.20 8.4</td>
<td>99   x.27 26.7</td>
<td>100 x.30 30.0</td>
<td>65.1</td>
<td>11</td>
</tr>
<tr>
<td>#4</td>
<td>15   x.20 3.0</td>
<td>90   x.27 24.3</td>
<td>100 x.30 30.0</td>
<td>100 x.30 30.0</td>
<td>57.3</td>
<td>8</td>
</tr>
<tr>
<td>#8</td>
<td>3    x.20 0.6</td>
<td>61   x.27 16.5</td>
<td>96   x.30 28.6</td>
<td>100 x.30 30.0</td>
<td>45.9</td>
<td>11</td>
</tr>
<tr>
<td>#10</td>
<td>2    x.20 0.4</td>
<td>51   x.27 13.8</td>
<td>94   x.30 28.2</td>
<td>100 x.30 30.0</td>
<td>42.4</td>
<td>4</td>
</tr>
<tr>
<td>#16</td>
<td>1    x.20 0.2</td>
<td>32   x.27 8.6</td>
<td>88   x.30 26.4</td>
<td>100 x.30 30.0</td>
<td>35.2</td>
<td>7</td>
</tr>
<tr>
<td>#30</td>
<td>1    x.20 0.2</td>
<td>20   x.27 5.4</td>
<td>71   x.30 21.3</td>
<td>100 x.30 30.0</td>
<td>26.9</td>
<td>8</td>
</tr>
<tr>
<td>#40</td>
<td>1    x.20 0.2</td>
<td>17   x.27 4.6</td>
<td>53   x.30 15.9</td>
<td>100 x.30 30.0</td>
<td>20.7</td>
<td>6</td>
</tr>
<tr>
<td>#50</td>
<td>1    x.20 0.2</td>
<td>14   x.27 3.8</td>
<td>32   x.30 9.6</td>
<td>100 x.30 30.0</td>
<td>13.6</td>
<td>7</td>
</tr>
<tr>
<td>#100</td>
<td>1    x.20 0.2</td>
<td>10   x.27 2.7</td>
<td>9    x.30 2.7</td>
<td>100 x.30 30.0</td>
<td>5.6</td>
<td>8</td>
</tr>
<tr>
<td>#200</td>
<td>0.8   x.20 0.16</td>
<td>6.9   x.27 1.86</td>
<td>3.3   x.30 0.99</td>
<td>100 x.30 30.0</td>
<td>3.01</td>
<td>3</td>
</tr>
</tbody>
</table>

FIGURE 3
SPECIFIC GRAVITY AND ABSORPTION OF COARSE AGGREGATE

(A Modification of AASHTO Designation T 85)

1. SCOPE

1.1 This test method covers the determination of specific gravity and absorption of coarse aggregate. The specific gravity may be expressed as bulk oven dry (OD) specific gravity, bulk saturated surface dry (SSD) specific gravity, or apparent specific gravity. The bulk (SSD) specific gravity and absorption are based on aggregate after 15 to 19 hours soaking in water. This method is not intended to be used with lightweight aggregate.

1.2 The specific gravity and absorption of coarse aggregate is normally determined on Plus No. 4 Material. When corresponding fine aggregate specific gravity and absorption are determined on Minus No. 8 Material, in accordance with Arizona Test Method 211, the coarse aggregate specific gravity and absorption shall be performed on Plus No. 8 Material. "Coarse aggregate" as herein referenced will be for either Plus No. 4 or Plus No. 8 Material. Coarse aggregate specific gravity and absorption for mineral aggregates used in asphaltic concrete, other than asphaltic concrete friction course, shall be determined on Plus No. 4 Material.

1.3 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.4 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.5 Reference may be made to AASHTO T 85 for additional information, including: terminology, significance and use, precision and bias, development of equations, and interrelationships between specific gravities and absorption as defined in AASHTO T 85 and T 84.
2. **APPARATUS**

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Balance - A balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least one gram. The balance shall be equipped with a suitable apparatus for suspending the sample container in water from the center of the weighing platform or pan of the balance.

2.3 Sample Container - A wire basket, of adequate size for the material being tested, and so constructed to prevent trapping of air when submerged. For test samples of Plus No. 4 Material, the wire mesh shall be No. 6 or finer. For test samples of Plus No. 8 Material, the wire mesh shall be No. 10 or finer. The wire for suspending the container from the balance shall be of the smallest practical size to minimize any possible effects of a variable immersed length.

2.4 Water Tank - A watertight tank into which the sample and container are placed for complete immersion while suspended below the balance, equipped with an overflow for maintaining a constant water level.

2.5 Sieves - Conforming to ASTM E11, and consisting of the necessary sieve sizes as described in Subsection 3.2 through 3.3.

3. **SAMPLING**

3.1 Sample the aggregate in accordance with Arizona Test Method 105.

3.2 The sample shall be sieved over the appropriate No. 4 or No. 8 sieve and the specific gravity and absorption determined as given in Subsections 5.3 and 5.5; or, testing may be performed on separate coarse size fractions, as described in Subsection 3.4 and the average values for specific gravity and absorption determined as given in Subsections 5.4 and 5.6. When testing is performed on separate size fractions of the coarse aggregate, a sieve analysis of the material shall be performed in accordance with Arizona Test Method 201. If the specific gravity and absorption of the fine aggregate is to be determined, save the appropriate Pass No. 4 or Pass No. 8 Material for testing in accordance with Arizona Test Method 211, otherwise it can be discarded.
3.3 If testing is not to be performed using separate size fractions, a representative test sample of the Plus No. 4 or Plus No. 8 Material with a minimum mass as shown below shall be obtained.

<table>
<thead>
<tr>
<th>Nominal Maximum Sieve Size, inches</th>
<th>Minimum Mass of Test Sample, kilograms (pounds)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>125 (276)</td>
</tr>
<tr>
<td>5</td>
<td>75 (165)</td>
</tr>
<tr>
<td>4-1/2</td>
<td>50 (110)</td>
</tr>
<tr>
<td>4</td>
<td>40 (88)</td>
</tr>
<tr>
<td>3-1/2</td>
<td>25 (55)</td>
</tr>
<tr>
<td>3</td>
<td>18 (40)</td>
</tr>
<tr>
<td>2-1/2</td>
<td>12 (26)</td>
</tr>
<tr>
<td>2</td>
<td>8 (18)</td>
</tr>
<tr>
<td>1-1/2</td>
<td>5 (11)</td>
</tr>
<tr>
<td>1</td>
<td>4 (8.8)</td>
</tr>
<tr>
<td>3/4</td>
<td>3 (6.6)</td>
</tr>
<tr>
<td>1/2 or less</td>
<td>2 (4.4)</td>
</tr>
</tbody>
</table>

3.4 It may be desirable to test the material in separate size fractions. An example is given below which illustrates the procedure to be followed. [If the sample contains more than 15 percent retained on the 1-1/2" sieve, test the Plus 1-1/2" Material in one or more size fractions separately from the smaller size fraction(s).] When an aggregate is tested in separate size fractions, the minimum mass of test sample for each fraction shall be the difference between the masses specified above for the largest and smallest sieve sizes of the fraction. The smallest size fraction tested shall meet the minimum mass requirements listed above, corresponding to the largest sieve size (passing) of the fraction. Size fractions of more than one sieve shall be proportionate with the sieve analysis for the sizes included.

Example: (A sample with the following sieve analysis is tested in the selected separate size fractions shown below.)

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>% Retained</th>
<th>% Pass</th>
</tr>
</thead>
<tbody>
<tr>
<td>2&quot;</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1-1/2&quot;</td>
<td>16</td>
<td>84</td>
</tr>
<tr>
<td>1&quot;</td>
<td>10</td>
<td>74</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>8</td>
<td>66</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>6</td>
<td>60</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>4</td>
<td>56</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>10</td>
<td>46</td>
</tr>
<tr>
<td>#4</td>
<td>5</td>
<td>41</td>
</tr>
</tbody>
</table>
Size fraction of Pass 2” to Retained 1-1/2”: 8000 g - 5000 g = 3000 g

Size fraction of Pass 1-1/2” to Retained 3/4”: 5000 g - 3000 g = 2000 g

For each sieve in this size fraction, the proportionate amount is determined:

Pass 1-1/2” to Retained 1”: \[\frac{10}{(10 + 8)} \times 2000 \text{ g} = 1111 \text{ g}\]
Pass 1” to Retained 3/4”: \[\frac{8}{(10 + 8)} \times 2000 \text{ g} = 889 \text{ g}\]

Size fraction of Pass 3/4” to Retained No. 4: 3000 g

For each sieve in this size fraction, the proportionate amount is determined:

Pass 3/4” to Retained 1/2”: \[\frac{6}{(6 + 4 + 10 + 5)} \times 3000 \text{ g} = 720 \text{ g}\]
Pass 1/2” to Retained 3/8”: \[\frac{4}{(6 + 4 + 10 + 5)} \times 3000 \text{ g} = 480 \text{ g}\]
Pass 3/8” to Retained 1/4”: \[\frac{10}{(6 + 4 + 10 + 5)} \times 3000 \text{ g} = 1200 \text{ g}\]
Pass 1/4” to Retained No. 4: \[\frac{5}{(6 + 4 + 10 + 5)} \times 3000 \text{ g} = 600 \text{ g}\]

4. **PROCEDURE**

4.1 Thoroughly wash the test sample to remove dust or other coatings from the surface.

4.2 Dry the test sample in a suitable pan or vessel to constant mass (see Subsection 4.4 below) at a temperature of 230 ± 9 °F. Constant mass shall be determined as follows: Dry the sample for at least 1 hour at 230 ± 9 °F. Record the weight of the sample to at least the nearest 1.0 gram. Continue drying and weighing until the weight does not change more than 1.0 gram at drying intervals of a minimum of 30 minutes.

4.3 Cool in air at room temperature for 1 to 3 hours for test samples of 1-1/2" nominal maximum size, or longer for larger sizes until the aggregate has cooled to a temperature that is comfortable to handle (approximately 122 °F).

4.4 Subsequently immerse the aggregate in water at room temperature for a period of 15 to 19 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 testing, the 15 to 19 hour soaking may also be eliminated.

**Note 2:** Values for absorption and bulk (SSD) specific gravity may be significantly higher for aggregate not oven dried before soaking than for the same aggregate which has been dried and soaked as specified in Subsections 4.2 through 4.4. This is especially true of particles larger than 3 inches since the water may not be able to penetrate the pores to the center of the particle in the prescribed soaking period.

4.5 Remove the test sample from the water and roll it in a large absorbent cloth or towel until all visible films of water are removed from the surface of the aggregate, wiping the larger particles individually as necessary. Take care to avoid loss of water from aggregate pores during surface-drying by completing the operation as quickly as possible. Immediately determine the mass of the test sample in the saturated surface-dry condition. Record this and all subsequent masses to at least the nearest gram.

4.6 After determining the saturated-surface-dry mass place the test sample in the sample container and determine its mass in water at 73 ± 3 °F. Take care to remove all entrapped air before determining the mass by shaking the container while immersed. The container shall be immersed to a depth sufficient to cover it and the test sample during mass determination.

4.7 Dry the test sample to constant mass at a temperature of 230 ± 9 °F. Constant mass shall be determined as described in Subsection 4.2. Cool in air at room temperature for 1 to 3 hours, or until the aggregate has cooled to a temperature that is comfortable to handle (approximately 122 °F) and determine the mass. Record the mass of the sample to at least the nearest gram.
5. **CALCULATIONS**

5.1 Calculate the bulk (OD) specific gravity as shown below:

\[
\text{Bulk (OD) Specific Gravity} = \frac{A}{(B - C)}
\]

Where:
- \( A \) = Mass of oven-dry test sample in air, grams.
- \( B \) = Mass of saturated-surface-dry test sample in air, grams.
- \( C \) = Mass of saturated test sample in water, grams.

5.2 Calculate the bulk (SSD) specific gravity as shown below:

\[
\text{Bulk (SSD) Specific Gravity} = \frac{B}{(B - C)}
\]

5.3 Calculate the apparent specific gravity as shown below:

\[
\text{Apparent Specific Gravity} = \frac{A}{(A - C)}
\]

5.4 When the sample is tested in separate size fractions, the average value for bulk (OD) specific gravity, bulk (SSD) specific gravity, or apparent specific gravity can be computed as the average of the respective specific gravities for the individual size fractions, weighted in proportion to the mass percentage of the size fractions in the original sample.

\[
G = \frac{1}{\sum \frac{P_n (G_n)}{100}}
\]

Where:
- \( G \) = Average specific gravity. All forms of expression of specific gravity can be averaged in this manner.
- \( P_1, P_2, \ldots, P_n \) = Mass percentage for each size fraction in original sample (total equals 100).
- \( G_1, G_2, \ldots, G_n \) = Respective specific gravity values for each size fraction for the type of specific gravity being averaged.
5.5 Calculate the percent absorption as shown below:

\[
\text{Percent Absorption} = \frac{(B - A)}{A} \times 100
\]

5.6 When the sample is tested in separate size fractions, the average value for percent absorption can be computed as the average of the individual percent absorption values for the respective size fractions, weighted in proportion to the mass percentage of the size fractions in the original sample.

\[
A = \left[ \frac{(P1)(A1)}{100} \right] + \left[ \frac{(P2)(A2)}{100} \right] + \ldots + \left[ \frac{(Pn)(An)}{100} \right]
\]

Where: 
- \( A \) = Average percent absorption.
- \( P1, P2, \ldots, Pn \) = Mass percentages for each size fraction in original sample (total equals 100).
- \( A1, A2, \ldots, An \) = Individual percent absorption values for the respective size fractions.

6. REPORT

6.1 Report specific gravity results to the nearest 0.001, and indicate the type of specific gravity, whether bulk (OD), bulk (SSD), or apparent.

6.2 Report the absorption result to the nearest 0.01%.

6.3 If the specific gravity and absorption values were determined without first drying the aggregate, as permitted in Subsection 4.4, it shall be noted in the report.

6.4 The size of the material tested shall be noted, i.e., Plus No. 4 or Plus No. 8.

7. PROCEDURE FOR DETERMINING COMBINED SPECIFIC GRAVITIES AND ABSORPTION FOR DIFFERENT SOURCES OF MATERIAL

7.1 Two methods are given below for determining the combined specific gravities and absorption for different sources of material, as described in Subsections 7.1.1 and 7.1.2.

7.1.1 The specific gravity and absorption may be determined for coarse aggregates from different sources which have been composited in the desired proportions and thoroughly blended.
7.1.2 The specific gravity and absorption of the coarse aggregate from each individual source may be determined and the combined specific gravity and absorption calculated as described in Subsections 7.1.3 through 7.1.7 below. (Refer to the example given in Subsection 7.1.7 for an illustration of the procedure and calculations.) The same size of material, either Plus No. 4 or Plus No. 8, shall be used to determine the individual specific gravities and absorption for each of the different sources.

7.1.3 For each individual material in the composite, its contribution to the total percent of coarse aggregate in the composite is determined and recorded to the nearest 0.01% as "IP":

\[
\text{IP} = \left( \frac{\text{Percent of Individual Material in Composite}}{\text{Percent of Coarse Aggregate in Individual Material}} \right) \times 100
\]

Where: \(\text{IP} = \) Contribution by each individual material to the total percent of coarse aggregate in the composite.

7.1.4 For each individual source, the percent of coarse aggregate in the composite is determined by summing the values for "IP" for that source, and recording the total as "P".

7.1.5 The combined specific gravity is calculated by the following:

\[
G = \frac{P_1 + P_2 + \ldots + P_n}{G_1 + G_2 + \ldots + G_n}
\]

Where: \(P_1, P_2, \ldots, P_n = \) For each individual source, the percent of coarse aggregate in the composite, "P".

\(G_1, G_2, \ldots, G_n = \) The coarse specific gravity for each individual source.

7.1.6 The combined absorption is calculated by the following:
7.1.7 The following is an example of the procedure and calculations described in Subsections 7.1.2 through 7.1.6 above. The example given is for a composite consisting of 26% coarse aggregate, 12% intermediate, and 47% fines from the primary source; with 15% blend from a secondary source. The coarse aggregate has 98% Plus No. 4, the intermediate has 94% Plus No. 4, the fines have 9% Plus No. 4, and the blend has 24% Plus No. 4.

Specific gravity and absorption for each of the different sources:

Primary Source (Coarse, Intermediate, Fine):

Bulk (OD) Specific Gravity = 2.549
Bulk (SSD) Specific Gravity = 2.586
Apparent Specific Gravity = 2.651
Absorption = 1.45%

Secondary Source (Blend):

Bulk (OD) Specific Gravity = 2.665
Bulk (SSD) Specific Gravity = 2.724
Apparent Specific Gravity = 2.781
Absorption = 2.21%

Determination of "IP" for individual materials, and "P" for individual sources, for the Primary Source (Coarse, Intermediate, Fine) and the Secondary Source (Blend):

Primary Source:

"IP" Coarse = \frac{(26 \times 98)}{100} = 25.48%

"IP" Intermediate = \frac{(12 \times 94)}{100} = 11.28%
"IP" Fine $= \frac{(47) \times (9)}{100} = 4.23\%$

"P" for Primary Source = 40.99\%

Secondary Source:

"IP" Blend $= \frac{(15) \times (24)}{100} = 3.60\%$

"P" for Secondary Source = 3.60\%

$[P (Primary)] + [P (Secondary)] = 44.59\%$

<table>
<thead>
<tr>
<th>Combined</th>
<th>Bulk (OD)</th>
<th>Specific</th>
<th>Gravity</th>
<th>= $\frac{44.59}{40.99 + 3.60} = 2.558$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Bulk (SSD)</td>
<td>Specific</td>
<td>Gravity</td>
<td>= $\frac{44.59}{40.99 + 3.60} = 2.597$</td>
</tr>
<tr>
<td></td>
<td>Apparent</td>
<td>Specific</td>
<td>Gravity</td>
<td>= $\frac{44.59}{40.99 + 3.60} = 2.661$</td>
</tr>
<tr>
<td></td>
<td>Absorption</td>
<td></td>
<td></td>
<td>= $\frac{2.597 - 2.558}{2.558} \times 100 = 1.52%$</td>
</tr>
</tbody>
</table>
EXAMPLE OF CALCULATIONS FOR COARSE SPECIFIC GRAVITY

Bulk Sp. Gr. (O.D. basis) = \[
\frac{A}{B - C} = \frac{4197}{4291 - 2610} = 2.497
\]

Where: 
- A = mass of oven-dry sample in air, g.
- B = mass of saturated-surface-dry sample in air, g.
- C = mass of saturated sample in water, g.

Bulk Sp. Gr. (SSD basis) = \[
\frac{B}{B - C} = \frac{4291}{4291 - 2610} = 2.553
\]

Apparent Sp. Gr. = \[
\frac{A}{A - C} = \frac{4197}{4197 - 2610} = 2.645
\]

Absorption, percent = \[
\frac{B - A}{A} \times 100 = \frac{4291 - 4197}{4197} \times 100 = 2.24 \%
\]
SPECIFIC GRAVITY AND ABSORPTION OF FINE AGGREGATE

(A Modification of AASHTO Designation T 84)

1. SCOPE

1.1 This method covers the determination of specific gravity and absorption of fine aggregate. The specific gravity may be expressed as bulk oven dry (OD) specific gravity, bulk saturated surface dry (SSD) specific gravity, or apparent specific gravity.

1.2 The specific gravity and absorption of fine aggregate is normally determined on Minus No. 4 Material. When the fine aggregate sample (Minus No. 4) contains a substantial amount of Minus No. 4 to Plus No. 8 Material, such as in mineral aggregate for asphaltic concrete friction course, the fine aggregate specific gravity and absorption shall be performed on Minus No. 8 Material. The fine aggregate specific gravity and absorption for mineral aggregates used in asphaltic concrete, other than asphaltic concrete friction course, shall be determined on Minus No. 4 Material. "Fine aggregate" as herein referenced will be either Minus No. 4 or Minus No. 8 Material. Corresponding coarse aggregate specific gravity and absorption testing, utilizing the appropriate Plus No. 4 or Plus No. 8 Material, shall be performed in accordance with Arizona Test Method 210.

1.3 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.4 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.5 When the specific gravity and absorption of the fine aggregate and the coarse aggregate are known, the combined aggregate specific gravity and absorption is determined, if necessary, utilizing Arizona Test Method 251.
2. **APPARATUS**

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Balance - A balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.1 gram.

2.3 Pycnometer - A 500 mL volumetric flask into which the fine aggregate test sample can be readily introduced and in which the volume content can be reproduced within 0.2 mL. Figure 1 is an illustration of the type of flask that must be utilized. The volume of the flask filled to the mark shall be at least 50 percent greater than the space required to accommodate the test sample. If this requirement is not met for a particular aggregate, the normal sample size of 500 ± 10 grams may be reduced only enough to satisfy the requirement.

2.4 Mold - A metal mold in the form of a frustum of a cone with dimensions as follows: 40 ± 3 mm inside diameter at the top, 90 ± 3 mm inside diameter at the bottom, and 75 ± 3 mm in height, with the metal having a minimum thickness of 0.8 mm (See Figure 2).

2.5 Tamper - A metal tamper having a mass of 340 ± 15 grams, and having a flat circular tamping face 25 ± 3 mm in diameter (See Figure 2).

2.6 Oven - Capable of maintaining a temperature of 230 ± 9 °F.

3. **SAMPLING**

3.1 Sample the aggregate in accordance with Arizona Test Method 105.

4. **PREPARATION OF TEST SAMPLE**

4.1 Obtain a representative approximate 1200 gram test sample of the fine aggregate.
4.2 Dry the test sample in a suitable pan or vessel to constant mass (see Subsection 4.3 below) at a temperature of 230 ± 9 °F. (Constant mass shall be determined as follows: Dry the sample for a minimum of 1 hour at 230 ± 9 °F. Record the weight of the sample to the nearest 0.1 gram. Continue drying and weighing until the weight does not change more than 0.1 gram at drying intervals of a minimum of 30 minutes.) Allow the sample to cool to comfortable handling temperature, cover with sufficient water to completely immerse it throughout the soaking period, and permit to stand for 15 to 19 hours.

4.3 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 testing, the 15 to 19 hour soaking may also be eliminated.

**Note:** Values for absorption and bulk (SSD) specific gravity may be significantly higher for aggregate not oven dried before soaking than for the same aggregate which has been dried and soaked as specified in Subsection 4.2.

4.4 Decant excess water with care to avoid loss of fines, spread the sample on a flat nonabsorbent surface exposed to a gently moving current of ambient or warm air, and stir frequently to secure homogeneous drying. Continue stirring and drying until the sample approaches a free-flowing condition. As the material begins to dry sufficiently, it may be necessary to work it with the hands in a rubbing motion to break up any conglomerations, lumps, or balls of material that develop.

4.5 Follow the procedure (cone test for surface moisture) given in Subsection 4.6 below to determine whether or not surface moisture is present on the fine aggregate particles. It is intended that the first trial will be made with some surface water in the sample. Continue drying with constant stirring, working the material with a hand-rubbing motion as necessary, and test at frequent intervals until the test indicates that the sample has reached a surface-dry condition. If the first trial of the surface moisture test indicates that moisture is not present on the surface, it has been dried past the saturated surface-dry condition. In this case, thoroughly mix a few milliliters of water with the fine aggregate and permit the specimen to stand in a covered container for 30 minutes. Then resume the process of drying and testing at frequent intervals for the onset of the surface-dry condition.
4.6 With one hand, hold the mold firmly on a smooth nonabsorbent surface with the large diameter down. The mold must be held firmly in place throughout the process of filling the mold, tamping, and removal of excess material from around the base of the mold. With the other hand, place a portion of the partially dried fine aggregate loosely in the mold, filling it until overflowing occurs, and heaping additional material above the top of the mold. Lightly tamp the fine aggregate into the mold with 25 light drops of the tamper. Each drop should start about 0.2 inch above the top surface of the fine aggregate. Permit the tamper to fall freely on each drop. Adjust the starting height to the new surface elevation after each drop and distribute the drops over the surface. Remove loose material from around the base of the mold, and lift the mold vertically. If surface moisture is still present, the fine aggregate will retain the molded shape. When the fine aggregate slumps slightly, it indicates that it has reached a surface-dry condition. Some angular fine aggregate or material with a high proportion of fines may not slump in the cone test upon reaching a surface-dry condition. This may be the case if fines become airborne upon dropping a handful of the sand from the cone test 4 to 6 inches onto a surface. For these materials, the saturated surface-dry condition should be considered as the point that one side of the fine aggregate slumps slightly upon removing the mold.

5. PROCEDURE

5.1 Only a pycnometer calibrated in accordance with the requirements of Appendix A3 of the Materials Testing Manual shall be used to perform this test. The mass of the pycnometer filled to its calibration capacity with water at 73 ± 3 °F is determined and recorded to the nearest 0.1 gram. Prior to determining the mass of the pycnometer filled with water to the calibration mark, the inside of the neck of the pycnometer just above calibration level shall be dried with a rolled up paper towel.

5.2 Obtain a representative 500 ± 10 gram sample of the saturated surface-dry fine aggregate prepared as described in Section 4. Immediately determine and record the weight of the sample to the nearest 0.1 gram.

5.3 Partially fill the calibrated pycnometer with water. Introduce the sample into the pycnometer and fill with additional water to approximately 90 percent of total capacity so that the water level after the elimination of all air bubbles is above the bottom of the neck but below the calibration line. Roll and agitate the pycnometer to eliminate all air bubbles.

Note: Adding a few drops of 99% grade isopropyl alcohol (ASTM D 770), after removal of air bubbles and just prior to bringing
the water level to its calibrated capacity, has been found useful in dispersing foam on the water surface.

5.4 Rinse the inside of neck of pycnometer with water and add sufficient water to bring the water level in the pycnometer to its calibrated capacity. Adjust its temperature to 73 ± 3 °F, if necessary by immersion in circulating water. With a rolled up paper towel, dry the inside neck of pycnometer just above the calibration level. Determine and record the mass of the pycnometer, sample, and water to the nearest 0.1 gram.

5.5. Remove the fine aggregate from the pycnometer and dry to constant mass at a temperature of 230 ± 9 °F. (Constant mass shall be determined as follows: Dry the sample for a minimum of 1 hour at 230 ± 9 °F. Record the weight of the sample to the nearest 0.1 gram. Continue drying and weighing until the weight does not change more than 0.1 gram at drying intervals of a minimum of 30 minutes.) Cool in air at room temperature for 1.0 ± 0.5 hours. Determine and record the mass to the nearest 0.1 gram.

**Note:** In lieu of drying and weighing the sample which has been removed from the pycnometer, a second portion of the saturated surface dry sample may be used to determine the oven dry weight. This sample must be obtained at the same time and shall weigh within ± 0.2 grams of the sample which is introduced into the pycnometer.
6. CALCULATIONS

6.1 Calculate the bulk (OD) specific gravity as shown below:

\[
\text{Bulk (OD) Specific Gravity} = \frac{A}{(B + S - C)}
\]

Where:
- \( A \) = mass of oven-dry test sample in air.
- \( B \) = mass of pycnometer filled with water to calibration mark.
- \( S \) = mass of saturated surface-dry test sample.
- \( C \) = mass of pycnometer with test sample and water to calibration mark.

6.2 Calculate the bulk (SSD) specific gravity as shown below:

\[
\text{Bulk (SSD) Specific Gravity} = \frac{S}{(B + S - C)}
\]

6.3 Calculate the apparent specific gravity as shown below:

\[
\text{Apparent Specific Gravity} = \frac{A}{(B + A - C)}
\]

6.4 Calculate the percent absorption as shown below:

\[
\text{Percent Absorption} = \left( \frac{S - A}{A} \right) \times 100
\]

7. REPORT

7.1 Report specific gravity results to the nearest 0.001, and indicate the type of specific gravity, whether bulk (OD), bulk (SSD), or apparent.

7.2 Report the absorption result to the nearest 0.01%.

7.3 If the specific gravity and absorption values were determined without first drying the aggregate, as permitted in Subsection 4.3, it shall be noted in the report. The source of the sample and the procedures used to prevent drying prior to testing shall also be reported.
7.4 The size of the material tested shall be noted, i.e., Minus No. 4 or Minus No. 8.

8. PROCEDURE FOR DETERMINING COMBINED SPECIFIC GRAVITIES AND ABSORPTION FOR DIFFERENT SOURCES OF MATERIAL

8.1 Two methods are given below for determining the combined specific gravities and absorption for different sources of material, as described in Subsections 8.2 and 8.3.

8.2 The specific gravity and absorption may be determined for fine aggregates from different sources which have been composited in the desired proportions and thoroughly blended.

8.3 The specific gravity and absorption of the fine aggregate from each individual source may be determined and the combined specific gravity and absorption calculated as described in Subsections 8.3.1 through 8.3.4 below. (Refer to the example given in Subsection 8.4 for an illustration of the procedure and calculations) The same size of material, either Minus No. 4 or pass Minus No. 8, shall be used to determine the individual specific gravities and absorption for each of the different sources.

8.3.1 For each individual material in the composite, its contribution to the total percent of fine aggregate in the composite is determined and recorded to the nearest 0.01% as "IP":

\[
IP = \left( \frac{\text{Percent of Individual Material in Composite}}{\text{Percent of Fine Aggregate in Individual Material}} \right) \times 100
\]

Where: IP = Contribution by each individual material to the total percent of fine aggregate in the composite.

8.3.2 For each individual source, the percent of fine aggregate in the composite is determined by summing the values for "IP" for that source, and recording the total as "P".
8.3.3 The combined specific gravity is calculated by the following:

\[
G = \frac{\sum P_i \cdot G_i}{\sum P_i}
\]

Where: \( P_1, P_2, \ldots, P_n \) = For each individual source, the percent of fine aggregate in the composite, "P".

\( G_1, G_2, \ldots, G_n \) = The fine specific gravity for each individual source.

8.3.4 The combined absorption is calculated by the following:

\[
\text{Combined Absorption} = \left( \frac{\text{Combined Bulk (SSD) Specific Gravity}}{\text{Combined Bulk (OD) Specific Gravity}} - 1 \right) \times \left( \frac{\text{Combined Bulk (OD) Specific Gravity}}{\text{Combined Bulk (OD) Specific Gravity}} \right)
\]

8.4 The following is an example of the procedure and calculations described in Subsections 8.3.1 through 8.3.4 above. The example given is for a composite consisting of 26% coarse aggregate, 12% intermediate aggregate, and 47% fine aggregate from the primary source; with 15% aggregate from a secondary source. The coarse aggregate has 2% pass the No. 4 sieve, the intermediate aggregate has 6% pass the No. 4 sieve, the fine aggregate has 91% pass the No. 4 sieve, and the aggregate from the secondary source has 76% pass the No. 4 sieve.

Fine aggregate specific gravity and absorption for each of the different sources:

**Primary Source (Coarse, Intermediate, Fine):**

- Bulk (OD) Specific Gravity = 2.576
- Bulk (SSD) Specific Gravity = 2.611
- Apparent Specific Gravity = 2.669
- Absorption = 1.36%
Secondary Source:

Bulk (OD) Specific Gravity = 2.641
Bulk (SSD) Specific Gravity = 2.686
Apparent Specific Gravity = 2.764
Absorption = 1.70%

Determination of "IP" for individual materials, and "P" for individual sources:

**Primary Source (Coarse, Intermediate, Fine):**

"IP" Coarse = \( \frac{(26) \times (2)}{100} = 0.52\% \)

"IP" Intermediate = \( \frac{(12) \times (6)}{100} = 0.72\% \)

"IP" Fine = \( \frac{(47) \times (91)}{100} = 42.77\% \)

"P" for the Primary Source = IP (Coarse) + IP (Intermediate) + IP (Fine)

\[ = (0.52\%) + (0.72\%) + (42.77\%) = 44.01\% \]

**Secondary Source:**

"IP" (Secondary Source) = \( \frac{(15) \times (76)}{100} = 11.40\% \)

"P" for Secondary Source = IP (Secondary Source) = 11.40\%
Total fine aggregate in the composite = \( P(\text{Primary Source}) + P(\text{Secondary Source}) \)

\[ = (44.01\%) + (11.40\%) = 55.41\% \]

Combined Bulk (OD) Specific Gravity = \( \frac{55.41}{44.01 + \frac{11.40}{2.576 + 2.641}} = 2.589 \)

Combined Bulk (SSD) Specific Gravity = \( \frac{55.41}{44.01 + \frac{11.40}{2.611 + 2.686}} = 2.626 \)

Combined Apparent Specific Gravity = \( \frac{55.41}{44.01 + \frac{11.40}{2.669 + 2.764}} = 2.688 \)

Combined Absorption = \( \frac{2.626 - 2.589}{2.589} \times 100 = 1.43\% \)
EXAMPLE OF CALCULATIONS FOR FINE SPECIFIC GRAVITY

Bulk Sp. Gr. (O.D. basis) = \[
\frac{A}{B + S - C} = \frac{499.9}{683.7 + 503.3 - 990.1} = \frac{2.539}{2.538852}
\]

Where:  
- \( A \) = mass of oven-dry sample in air, g.  
- \( B \) = mass of pycnometer filled with water, g.  
- \( C \) = mass of pycnometer with sample and water to calibration mark, g.  
- \( S \) = mass of saturated-surface-dry sample, g.

Bulk Sp. Gr. (SSD basis) = \[
\frac{S}{B + S - C} = \frac{503.3}{683.7 + 503.3 - 990.1} = \frac{2.556}{2.556120}
\]

Apparent Sp. (\( \frac{A}{B + A - C} \)) = \[
\frac{499.9}{683.7 + 499.9 - 990.1} = \frac{2.583}{2.583463}
\]

Absorption, percent = \[
\frac{S - A}{A} \times 100 = \frac{503.3}{499.9} = 0.68 \%
\]
PERCENTAGE OF FRACTURED COARSE AGGREGATE PARTICLES

(An Arizona Method)

1. SCOPE

1.1 This test method describes the procedure for determining the percentage of fractured particles in coarse aggregate. This test procedure is performed on plus No. 8 and larger material, unless otherwise specified.

1.2 When this test procedure is referenced for the determination of “crushed faces”, such as in specifications or other test procedures, it shall be understood to be the determination of “fractured coarse aggregate particles”.

1.3 A fractured face is defined as an angular, broken surface of an aggregate particle created by crushing, by other artificial means, or by nature. A face is only considered fractured if it has sharp and well defined edges and the projected area of the fractured face is at least 25 percent of the maximum projected area of the aggregate particle visible considering the particle’s outline at all possible orientations of the aggregate particle.

1.4 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.5 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.
2.2 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 1 gram.

2.3 A magnifying glass, preferably mounted on a stand and with a built-in light.

2.4 Sieves conforming to ASTM E11, of sizes necessary for the material being tested.

2.5 Miscellaneous pans, containers, spatulas, etc.

3. **SAMPLE PREPARATION**

3.1 A representative sample of the aggregate shall be obtained. The coarse aggregate test sample shall be proportionate for the gradation of the material being tested. The size of the test sample shall be such that it contains a minimum of 500 particles. It is not expected that an actual count of the number of particles be made to assure compliance with this requirement. It is required, however, that a reasonably accurate estimate be made to determine that the test sample contains at least 500 particles. Fractured face determination must be made on all particles in the test sample.

4. **PROCEDURE**

4.1 Weigh the test sample, and record the weight to the nearest gram as "Wa". To facilitate the inspection of the particles, the test sample may be washed and dried to constant weight prior to determining the test sample weight.

4.2 Spread the test sample onto a flat surface, and observe the aggregate particles under the magnifying glass with adequate light.

4.3 If not otherwise specified, separate aggregate particles having at least one fractured face from those that are unfractured. If specifications require that a percentage of the aggregate particles having at least two fractured faces be determined, separate aggregate particles having at least two fractured faces from those that have less than two fractured faces. A spatula or other blade-like tool may be helpful in separating particles.

4.4 Weigh the portion of fractured particles, and record the weight to the nearest gram. If aggregate particles with at least one fractured face have been
5. **CALCULATIONS**

5.1 If aggregate particles with at least one fractured face have been determined, calculate the percentage of fractured particles, \( FF \), and record to the nearest percent as shown below:

\[
FF = \frac{W_f}{W_a} \times 100
\]

Where:
- \( FF \) = Percentage of fractured particles with at least one fractured face.
- \( W_f \) = Weight of fractured particles with at least one fractured face.
- \( W_a \) = Weight of test sample.

5.2 If aggregate particles with at least two fractured faces have been determined, calculate the percentage of fractured particles, \( FF_2 \), and record to the nearest percent as shown below:

\[
FF_2 = \frac{W_2}{W_a} \times 100
\]

Where:
- \( FF_2 \) = Percentage of fractured particles with at least two fractured faces.
- \( W_2 \) = Weight of fractured particles with at least two fractured faces.
- \( W_a \) = Weight of test sample.
DETERMINATION OF CEMENTITIOUS MATERIAL CONTENT FOR CEMENTITIOUSLY TREATED MIXTURES

1. SCOPE

1.1 This procedure is used to determine the cementitious material content of a cement treated mixture.

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

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 4 inch proctor mold having a capacity of approximately 1/30 cubic foot, with an internal diameter of 4.000 plus 0.024 or minus 0.016 inches and a height of 4.584 plus 0.005 or minus 0.008 inches. The mold shall have a nominal wall thickness of 1/4 inch. It shall be equipped with an extension collar approximately 2-3/8 inches high.

2.3 A hand or mechanical rammer weighing 5.50 ± 0.02 pounds, having a flat face, and equipped with a suitable arrangement to control the height of drop to a free fall of 12 ± 0.06 (1/16) inches above the elevation of the soil. The rammer face shall be circular with a diameter of 2.000 plus 0.010 or minus 0.015 inches.

2.4 Hard steel straightedge, at least 10 inches in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.01 inch per 10 inches (0.1 percent) of length with the portion used for trimming the soil.

2.5 A circular sample follower with a diameter slightly less than the inner diameter of the proctor mold to aid in the removal of the mold collar.
2.6  Scale or balance capable of measuring the maximum weight to be determined, accurate to at least 0.1 g.

2.7  Oven capable of maintaining a temperature of 230 ± 9 °F

2.8  Sieves of sizes as required for screening conforming to the requirements of ASTM E11.

2.9  Miscellaneous mixing tools and pans.

2.10  Equipment as required for ARIZ 241, “Compressive Strength of Molded Cement Treated Base or Soil-Cement Specimens”.

2.11  Non-absorptive glass or plexiglass plates larger than the diameter of the 4 inch proctor mold.

3.  SAMPLE

3.1  Approximately 75 kg of material will be required. The amount required for the sieve analysis will be as determined in ARIZ 201, approximately 20000 g will be required for the optimum moisture determination, and approximately 40000 g will be required to develop the five separate 7500 g test samples.

3.2  If the sample is damp when received, it shall be dried until it becomes friable under a trowel. Drying may be accomplished by air drying or by the use of a drying apparatus such that the temperature of the sample does not exceed 140 °F. Prepare the sample for testing by separating the Plus No. 4 Material and breaking up the remaining soil aggregations to pass the No. 4 sieve in such a manner to avoid reducing the natural size of individual particles.

3.3  Select and prepare five separate test samples of approximately 7500 g and one 20000 g sample for optimum moisture content determination, corresponding to the sieve analysis of the total sample, in accordance with the following procedure.

Determine the gradation of the sample in accordance with ARIZ 201. For samples containing Plus 3/4 Inch Material, the material retained on the 1/2 inch, 3/8 inch, 1/4 inch, and No. 4 sieves shall be stored separately. Aggregate retained on the 3/4 inch sieve shall not be used, but its percentage shall be distributed proportionately among the aforementioned sieves.
NOTE: No change is made in the Percent Pass No.4 Material. See Subsection 3.3 of ARIZ 221 for further information regarding the proportional redistribution of the Plus 3/4 Inch Material.

3.4 Determine the optimum moisture content of the 20000 g sample in accordance with ARIZ 221 with the anticipated required cement content included in the material during the performance of the procedure.

3.5 The moisture content to be added to each of the five 7500 g test samples prior to compaction is the optimum moisture content obtained in Subsection 3.4.

3.6 Add the first of the cement contents to be used, to the first sample, and mix together thoroughly.

4. COMPACTION

4.1 Form a specimen by compacting the prepared mixture in the four inch mold (with the extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. Compact each layer with 25 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation.

4.2 Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the soil in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted soil even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Determine the wet density, “WD”, of the compacted soil by the following:

\[
WD = \frac{M1 - M2}{VM \times 453.6 \text{ (grams/lb)}}
\]

Where:  
WD = Wet density of compacted soil, lb./cu. ft.  
M1 = Weight of compacted specimen and mold, grams  
M2 = Weight of the mold, grams
4.3 Compact the other two specimens from the first sample, at the same cementitious material contents, in the same manner as Subsection 4.1.

4.4 Extrude all three specimens from their respective molds, place on glass or plexiglass plates, and store for curing in a moist condition.

4.5 Determine the moisture content of the specimens from the left over material from the first sample in accordance with AASHTO T 265, “Laboratory Determination of Moisture Content of Soils”. Calculate percent moisture and record to the nearest 0.1 percent by the following:

\[
\% \text{ Moisture} = \frac{WW - DW}{DW} \times 100
\]

Where: \( WW = \) Weight of wet soil, grams
\( DW = \) Weight of dry soil, grams

4.6 Calculate the dry density and record to the nearest 0.1 lb./cu. ft. by the following:

\[
DD = \frac{WD}{% \text{ Moisture} \times 100} \times 100
\]

Where: \( DD = \) Dry density of compacted soil, lb./cu. ft.
\( WD = \) Wet density of compacted soil, lb./cu. ft.

4.7 Repeat the procedures in Subsections 4.1 to 4.6 to the additional samples with increased cementitious material contents of 2% increments, until a complete bracketing of the specification requirements is met.

5. **PROCEDURE**

5.1 All specimens must be cured in a moist condition for six days and then immersed for a period of 24 hours in saturated lime water at 65 to 80 °F for 24 hours.
5.2 Specimens shall then be prepared and tested for compressive strength in accordance with ARIZ 241.

6. REPORT

6.1 Report the cylinder number, dry density, cementitious material content, moisture content, and 7-day compressive strength of all specimens tested, and the average 7-day compressive strength for the specimens tested at each of the cementitious material contents.

6.2 Following is an example of testing information obtained and how the information would be compared to the Special Provisions of a project for the purpose of determining the appropriate cementitious material content:

Assume the test information as shown in the following table was obtained:

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<thead>
<tr>
<th>Sample Number</th>
<th>Specimen Number</th>
<th>Density (lbs./cu. ft.)</th>
<th>Cementitious Material Content</th>
<th>Moisture Content</th>
<th>7-Day Strength (PSI)</th>
<th>Average 7-Day Strength (PSI)</th>
</tr>
</thead>
<tbody>
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<td>117.1</td>
<td>2.0</td>
<td>9.4</td>
<td>506</td>
<td>509</td>
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<td>2</td>
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<td>2.0</td>
<td>9.5</td>
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<td>2.0</td>
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<td>4.0</td>
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<td>6.0</td>
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<td>9.5</td>
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<td>1502</td>
</tr>
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<td>8.0</td>
<td>9.4</td>
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<td>9.4</td>
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<td>10.0</td>
<td>9.3</td>
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<td>10.0</td>
<td>9.4</td>
<td>1722</td>
<td>1722</td>
</tr>
</tbody>
</table>

A graphical presentation of the test information above would appear as follows:
If the Special Provisions required that a soil-cement mixture shall have a design compressive strength of at least 1250 pounds per square inch at seven days, this test information would indicate that in order to meet this requirement, 5.9 percent cementitious material by total weight should be used.
MOISTURE-DENSITY RELATIONS OF CEMENTITIOUSLY TREATED MIXTURES

1. SCOPE

1.1 This method covers procedures for determining the relationship between moisture content and the density of cement treated mixtures when compacted at varying hydration curing times dependent on and consistent with construction operation procedures.

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

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 4 inch proctor mold having a capacity of approximately 1/30 cubic foot, with an internal diameter of 4.000 plus 0.024 or minus 0.016 inches and a height of 4.584 plus 0.005 or minus 0.008 inches. The mold shall have a nominal wall thickness of 1/4 inch. It shall be equipped with an extension collar approximately 2-3/8 inches high.

2.3 A hand or mechanical rammer weighing 5.50 ± 0.02 pounds, having a flat face, and equipped with a suitable arrangement to control the height of drop to a free fall of 12 ± 0.06 (1/16) inches above the elevation of the soil. The rammer face shall be circular with a diameter of 2.000 plus 0.010 or minus 0.015 inches.

2.4 Hardened-steel straigtedge, at least 10 inches in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.01 inch per 10 inches (0.1 percent) of length with the portion used for trimming the soil.
A circular sample follower with a diameter slightly less than the inner diameter of the proctor mold to aid in the removal of the mold collar.

Scale or balance capable of measuring the maximum weight to be determined, accurate to at least 0.1 g.

Oven capable of maintaining a temperature of 230 ± 9 °F.

Sieves of sizes as required for screening conforming to the requirements of ASTM E11.

Miscellaneous mixing tools and pans.

3. SAMPLE PREPARATION

If the sample is damp when received, it shall be dried until it becomes friable under a trowel. Drying may be accomplished by air drying or by the use of a drying apparatus such that the temperature of the sample does not exceed 140 °F. Prepare the sample for testing by separating the Plus No. 4 Material and breaking up the remaining soil aggregations to pass the No. 4 sieve in such a manner as to avoid reducing the natural size of individual particles.

Select and prepare five separate 2500 g test samples, corresponding to the sieve analysis of the total sample, in accordance with the following procedure.

Determine the gradation of the sample in accordance with ARIZ 201. For samples containing Plus 3/4 Inch Material, the material retained on the 1/2 inch, 3/8 inch, 1/4 inch, and No. 4 sieves shall be stored separately. Aggregate retained on the 3/4 inch sieve shall not be used, but its percentage shall be distributed proportionately among the aforementioned sieves.

Determine a Percent Retained Factor for calculating the Adjusted Percent Retained by the following:

\[
\text{% Retained Factor} = \frac{\text{Total Retained on 3/4" Sieve}}{\text{Cumulative % Retained on 1/2", 3/8", 1/4", and No. 4 Sieves}}
\]

Record the Percent Retained Factor to at least four decimal places.
3.5 Calculate the Adjusted Percent Retained for the 1/2 inch, 3/8 inch, 1/4 inch and No. 4 sieves by the following:

\[
\text{Adjusted Percent Retained} = (\% \text{ Retained} \times \% \text{ Retained Factor}) + \% \text{ Retained}
\]

Record the adjusted percent retained for each sieve to the nearest hundredth.

3.6 The sum of the new Adjusted Percent Retained values should total 100% when rounded.

3.7 Multiply each of the newly Adjusted Percent Retained values by 2500, then divide by 100, to obtain the quantity of material for each sieve size that should be combined to create each sample.

**NOTE:** Due to rounding in the procedure, the sample may not equal exactly 2500 g.

3.8 An example of a coarse sieve analysis which requires adjusting to prepare test samples of 2500 g is shown in Table 1.

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>% Retained</th>
<th>Rounded Adjusted % Retained</th>
<th>Weight in g (2500 g sample)</th>
<th>Accumulative Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>2”</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2-1/2”</td>
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</tr>
<tr>
<td>1”</td>
<td>8</td>
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</tr>
<tr>
<td>3/4”</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
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<td>180</td>
<td>180</td>
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<tr>
<td>3/8”</td>
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<td>9.00</td>
<td>225</td>
<td>450</td>
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<td>#4</td>
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<td>225</td>
<td>675</td>
</tr>
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<tr>
<td>Total</td>
<td>100</td>
<td>100</td>
<td>2500</td>
<td></td>
</tr>
</tbody>
</table>

4. **COMPACTATION**

4.1 Add to the prepared samples the required amount of cementitious material as determined in ARIZ 220, “Cement Content Required for Cement Treated Mixtures.” If this has not yet been determined, an estimate shall be made and
this amount added. Mix thoroughly to a uniform color. Add sufficient water to dampen the mixture to approximately four to six percentage points below the estimated optimum moisture content, and mix thoroughly. Cover and allow to cure for 5 minutes to aid dispersion of the moisture.

4.2 Form a specimen by compacting the prepared mixture in the four inch mold (with the extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. Compact each layer with 25 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation.

4.3 Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the soil in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted specimen even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Determine the wet density, “WD”, of the compacted specimen by the following:

\[
WD = \frac{M_1 - M_2}{VM \times 453.6 \text{ (grams/lb)}}
\]

Where:
- WD = Wet density of compacted specimen, lb./cu. ft.
- M1 = Weight of compacted specimen and mold, grams
- M2 = Weight of the mold, grams
- VM = Volume of the mold, cu. ft., determined in accordance with ARIZ 225, Appendix A

4.4 Remove the material from the mold and slice vertically through the center. Take a representative minimum 300 gram sample from the full length and width of one of the cut faces. Weigh immediately, and dry to a constant weight in an oven at 230 ± 9 °F to determine the moisture content in accordance with AASHTO T 265. Record the weight of wet soil to the nearest 0.1 gram as "WW", and record the weight of oven dry soil to the nearest 0.1 gram as "DW". Calculate percent moisture and record to the nearest 0.1 percent by the following:
% Moisture = \frac{WW - DW}{DW} \times 100

Where: \( WW = \) Weight of wet soil, grams
\( DW = \) Weight of dry soil, grams

4.5 Calculate the dry density and record to the nearest 0.1 lb./cu. ft. by the following:

\[ DD = \frac{WD}{% \text{ Moisture} \times 100} \times 100 \]

Where: \( DD = \) Dry density of compacted soil, lb./cu. ft.
\( WD = \) Wet density of compacted soil, lb./cu. ft.

Repeat the procedure in Subsections 4.1 through 4.5 for the sample, increasing the moisture content for each specimen by approximately two percentage points, as necessary to establish a moisture-density curve which rises to a peak and then falls away.

4.6 Maximum density and optimum moisture content shall be determined as described in Subsections 7.1 through 7.4 of ARIZ 225.

Repeat the procedures in Subsections 4.1 through 4.6 on samples allowed to cure prior to compaction for the following curing time periods: 5 minutes, 30 minutes, and 60 minutes. (It will not be necessary to make a four-hour determination for all mixtures. This need should be established on an individual basis.) These separate determinations shall be used for developing a family of curves for use for compaction control with each curve designated according to its respective curing time. If it is determined that the delay in compaction time (i.e., the curing time between incorporation of the cement and compaction) will not vary throughout the project construction schedule, then the number of delayed compaction curves can be limited to only those applicable without running the complete family.
ROCK CORRECTION PROCEDURE
FOR MAXIMUM DENSITY DETERMINATION
OF CEMENT TREATED MIXTURES
(An Arizona Method)

Scope

1. This method is to be used to determine the maximum density of the total sample of cement treated mixture for which the percentage of material retained on the No. 4 sieve has been determined as well as the maximum dry density of the passing No. 4 fraction and the bulk specific gravity of the plus No. 4 fraction. This test procedure is basically the same as ARIZ 227, Rock Correction Procedure for Maximum Density Determination, except for the addition of the concept of delay compaction time.

Procedure

2. The test is made in the following manner:

(a) Determine the maximum dry density of the material passing the No. 4 sieve by means of ARIZ 221, consistent with the desired curing time.

(b) Determine the percentage of rock (larger than the No. 4 sieve) in the total sample.

(c) Determine the bulk specific gravity of the rock in accordance with AASHTO T 85, on an oven-dried basis.

NOTE: Once determined for a source, the specific gravity may usually be used for all rock corrections on material from that source. Any slight change in specific gravity should not change the maximum density to a large extent. (A change of ± 0.05 could result in a change of ± 0.6 lb. per cubic foot.) Similarly, the maximum dry density of the passing No. 4 material having been determined by Method A for a source may usually be used for that source providing the screen analysis of the passing No. 4 material and the Plasticity Index remain reasonably uniform.

(d) Determine the maximum dry density of the total sample by use of the formula (e) or by use of the nomograph (f).

(e) Determination of density by use of formula:

\[
D = \frac{(100 - R) \cdot d + (0.9 \cdot S \cdot x \cdot 62.4)}{100}
\]

Where:

- \( D \) = Dry density of total sample containing R percent rock, lbs/cu. ft. (Varies according to curing time required.)
- \( R \) = Percent of rock retained on No. 4 sieve.
- \( d \) = Dry density of material passing No. 4 sieve, lbs./cu. ft.
- \( S \) = Specific Gravity of rock.

Example:

- \( R = 29\% \) rock
- \( d = 110.0 \) lbs. per cu. ft. (2 hour curing time.)
- \( S = 2.50 \)
- \( D = \frac{(100 - 29) \cdot 110.0 + (56.2 \cdot 29 \cdot 2.50)}{100} \)
- \( D = \frac{(71 \cdot 110.0) + 4074}{100} \)
- \( D = 7810 + 4074 = 118.8 \) lbs. per cu. ft.

(f) Determination of density by use of nomograph: Use the nomograph shown in Fig. 1 following the instructions given. Data from the example above has been used to show methods for nomograph solutions.

Limitation of Method

3. The rock correction procedure is not accurate when the rock content (retained on No. 4 sieve) is in excess of 50% (or 60% in the case of uniformly graded rock which consists of 100% passing the 1 inch sieve), or when the material contains rock which will not pass a 3-inch sieve. If any of these conditions are present, the test should not be completed; however, the report should state that the test cannot be made due to excess rock, or large rock, and the sieve analysis reported to verify this. This limitation shall apply except for the case stated in Section 4.

Volcanic Cinders or Light Porous Material

6. The rock correction procedure is not accurate when the rock in the material consists of volcanic cinders or light weight porous material on which the specific gravity determination cannot be accurately made; for these materials, Methods C or D (ARIZ 226) shall be used.

Use of Results

5. The maximum density obtained by this method shall be used in comparison with the determination of the field density to determine the percentage of compaction.
ARIZONA DEPARTMENT OF TRANSPORTATION

MATERIALS SECTION

Rock Correction Procedure
for
Maximum Density Determination
of
Cement Treated Mixtures

\[ D = \frac{(100 - R)d + (56.2 RS)}{100} \]

Where:
- \( D \) = dry density of sample containing \( R \) percent rock, lb. per cu. ft.
- \( R \) = percent rock (plus No. 4 material)
- \( d \) = dry density of portion passing No. 4 sieve, lb. per cu. ft.
- \( S \) = bulk specific gravity of rock, oven dry basis.

Example for use of chart:
- Known - Dry density of cement treated material passing No. 4 sieve (2 hr. curing time) = 110.0 lb. per cu. ft. Plot at A.
- Note: Other curing time dry densities are plotted to indicate the anticipated pattern of results.
- Known - Specific gravity of rock = 2.5. Plot at B, draw line AB.
- Known - Percent of rock in total sample = 29. Plot at C, draw line CD and locate D at intersection with line AB.
- Draw line horizontally from point D to the left to locate point E. Dry density of total sample (2 hr. curing time) = 118.8 lb. per cu. ft.

Figure 1.
FIELD DENSITY OF CEMENT TREATED MIXTURES
BY SAND CONE METHOD OR BY
RUBBER BALLOON METHOD

Scope

1. These methods are used to determine the in-place density of cement treated mixtures by determining the weight and moisture content of material removed from a test hole and subsequent measure of the test hole volume by either the Rubber Balloon Method (ARIZ 231) or the Sand Cone Method (ARIZ 230). Tests ARIZ 230 or 231 should be followed except for the differences outlined under Procedures below.

Procedures

1. Determine the time elapsed between the incorporation of cement and water to the mixture and the completion of compaction. This information is needed to refer to the proper Rock Correction Delayed Curing line (ARIZ 222).

2. If the rock content is greater than 50% (or 60% in the case of materials which are to consist of 100% passing the 1-inch sieve) report the sieve analysis with a note stating that the density is not determinable due to excess rock.

3. If any rock is retained on the 3-inch sieve, verify this with a sieve analysis and call this the end point. This sieve analysis shall be reported with a note stating that the density is not determinable due to the presence of rock retained on the 3-inch sieve.
MAXIMUM DENSITY AND OPTIMUM MOISTURE OF SOILS
METHODS C & D
(A Modification of AASHO T 99, Methods C & D)

Scope

1. Method C shall be used only on volcanic cinders and light porous material containing less than 50% of material retained on the No. 4 sieve.

Method D shall be used only on volcanic cinders and light porous materials containing more than 50% and less than 80% of material retained on the No. 4 sieve. If the material contains more than 60% retained on the No. 4 sieve, report the sieve analysis with a note stating the density is not determinable due to excess material retained on the No. 4 sieve.

Apparatus

2. The apparatus shall consist of the following:

   (a) 4.0-in. proctor mold (for Method C) having a capacity of 1/30 (.0333) cu. ft. with internal diameter of 4.0 ±.005 in. and a height of 4.584 ±.005 in.

   (b) 6.0-in. proctor mold (for Method D) having a capacity of 1/13.33 (0.075) cu. ft., with internal diameter of 6.0 ±.005 in. and a height of 4.584 ±.005 in. (See Fig. 1.)

   (c) A hand rammer or mechanical compactor of 2-in. diameter, having a flat circular face and weighing 5.5 lb.

   (d) Steel straightedge 12" or longer, having one beveled edge.

   (e) Balance, capacity 20 kg. sensitive to 1 g.

   (f) Drying oven, operating range 110° ±5° C.

   (g) Miscellaneous mixing tools and pans.

Sample

3. Enough material should be on hand to make five compacted samples.

Approximately 13,000 g. for Method C.

Approximately 25,000 g. for Method D.

(These weights may have to be adjusted depending on the type of material.)

FIGURE 1
Cylindrical Mold, 6.0-In, for Soil Tests
(a) A sieve analysis, (ARIZ 201 and 203) must be run to determine the proportion of rock.

Samples which contain approximately 100% passing the % sieve shall be prepared in accordance with the following procedure:

A uniform sample of approximately 15,000 g. so as to obtain not less than five compaction samples or test charges, shall be used. The sample shall be separated by means of a mechanical sample splitter into equal portions of approximately the desired weight. With some practice the charges can be obtained by using only the splitter. The operator shall not add to or take away from any portion by scooping with the hand or other means, as this may cause segregation.

(b) Samples containing rock retained on the % sieve shall be processed in the following manner:

The material retained on the %, %, %, and No. 4 sieves shall be stored in separate containers during the sieve analysis procedure. Rock retained on the % sieve shall not be used, but its percentage shall be distributed proportionately among the sieves mentioned above. Table I shows a coarse sieve analysis which required adjusting, shows the adjusted percentages, and also shows the material to build up a 2200 g. charge. Essentially the same procedure shall be followed to build a 5000 g. charge in the case of Method D. These weights may require adjustment if the charge is too large.

(c) The adjusted % retained values are found by dividing each of the percentages retained on the %, %, %, and No. 4 (from Table I), by the total percentage of material coarser than No. 4, x 10, and adding these quotients to the respective sieves below the %, i.e., \(7 \div 36 = 1.9 = 2\); \(8 \div 36 = 2.2 = 2\); and \(12 \div 36 = 3.3 = 3\). The final correction should total 100%.

(d) Use the "adjusted % Ret." values to build up the compaction charges of 2200 g. each, i.e.,

\[ \frac{2200}{100} \times 9 = 198 \text{ g. of } \frac{1}{2}\]

rock needed (Table I). The accumulative total is for weighing the charge as follows:

1. Weigh out the first size (\(\frac{1}{2}\)) into the balance scoop (198 g. in Table I), then set the weights at 374 g. (198 + 176) and add the \(\frac{1}{2}\) rock until the instrument balances. Next set the weights at 638 g., (374 + 264) and add the \(\frac{1}{4}\) rock until the instrument again comes to balance. Continue in this manner until the charge weighing 2200 g. is built up. Weigh out 5 such charges and save them in separate pans.

Mixing

4. The amount of water that will be required for a sample is variable and requires some experience on the part of operator, but the following general rule may be of help. Add 4% water to the sample and mix thoroughly; when mixing is complete, pick up a handful of dirt and squeeze it, then open the hand, let the dirt fall, and immediately blow the breath upon the open hand. If the hand feels cool, but not cold, another 2% water shall be added, thoroughly mixed, and the squeeze and blow test repeated. Continue to add 2% portions of water, and check, until the feeling of coolness changes to cold. At this point, gather the material into a compact pile and cover it with a slightly damp cloth to inhibit drying. Record the total percentage of water added.

Compaction

5. Compact the material as follows:

(a) Place approximately \(\frac{1}{2}\) of the mixed material in the mold, moving the scoop around the top of the mold to insure even distribution of the material. Tamp the material lightly with the end of the hand rammer,
or with the knuckles if the mechanical rammer is used.

(b) Compact this first layer with the rammer falling through a distance of 12 inches, moving the rammer to a different place each time so the whole surface is covered.

Method "C" requires 25 blows per layer.

Method "D" requires 56 blows per layer.

(c) Add another \( \frac{1}{4} \) portion of the dampened material and compact it as before.

(d) Add and compact the third and final layer. This last layer, when compacted, should extend up into the extension collar a short distance.

(e) Place the mold in the mixing pan and remove the collar so as to not crack or crumble the compacted specimen, then strike off the top of the specimen even with the top of the mold, using the steel straightedge. If any holes are left as rocks are scraped out, these holes shall be filled with fine material and smoothed off.

Calculations

6. The samples are weighed and calculations are as follows:

(a) Remove the base, wipe away any dirt from the mold and weigh the sample and mold to the nearest gram. Record the weight on the work card.

(b) Subtract the weight of the mold to obtain the wet weight of the sample in grams.

(c) Calculate the wet Density (D\(_w\)) in pounds per cu. ft. as follows:

**Method C**

\[
D_w = \frac{\text{wet wt.} \times 0.06614}{\text{Vol. of 4" mold} \times 453.6 \, \text{g./lb.}}
\]

Where 0.06614 = \( \frac{1}{\text{Vol. of 4" mold} \times 453.6 \, \text{g./lb.}} \)

**Method D**

\[
D_w = \frac{\text{wet wt.} \times 0.02939}{\text{Vol. of 6" mold} \times 453.6 \, \text{g./lb.}}
\]

Where 0.02939 = \( \frac{1}{\text{Vol. of 6" mold} \times 453.6 \, \text{g./lb.}} \)

(d) The approximate dry density of the sample (D\(_d\)) may be calculated as follows:

\[
D_d = \frac{D_w}{100 + \% \text{water added}} \times 100
\]

Example:

Wet Density, \( D_w = 106.3 \, \text{lb./cu. ft.} \)

% Water added = 14

\[
D_d = \frac{106.3}{114} \times 100 = 93.2 \, \text{lb./cu. ft.}
\]

**NOTE:** These densities are approximate only and will be corrected when final moisture results are obtained.

The approximate dry density is useful in deciding how much water to add in later trial batches. For instance, if the 14% water in the example, shown on the previous page, gave a density of 93.2 lb. per cu. ft., and a later increase to 16% gave the same density of 93.2 lb. per cu. ft., then the next trial to be run should be dropped to 12% water. (This was tried and resulted in a compaction of 92.1 lb. per cu. ft., giving a curve which rose from 12% to 14% and fell away at 16%) One or two additional samples with either 10% or 18% water added should be compacted, to aid in establishing the curve. By carefully observing the approximate dry density of the compacted samples, the operator should be able to vary the moisture content as the test proceeds so that when the final results are plotted a curve will result that rises to a peak and then recedes.

**Moisture Determination**

7. The percentage of moisture in the compacted samples is determined as follows:

(a) Break the compacted sample into small pieces, being careful to lose none. Dry the sample in an oven at 110° ±5° C to constant weight, and record the weight.

**NOTE:** The "Speedy" Method is not applicable for moisture determinations.

(b) Calculate the moisture content as follows:

\[
\% \text{Moisture} = \frac{W_w - W_d}{W_d} \times 100
\]

Where:

\( W_w \) = weight of wet soil, g.

\( W_d \) = weight of dry soil, g.

Example:

\[
\% \text{Moisture} = \frac{322 \, \text{g.} - 286 \, \text{g.}}{286 \, \text{g.} - 22 \, \text{g.}} \times 100 = 13.6\%
\]
(c) The dry density in lb. per cu. ft. for each sample is calculated as follows:

\[ D = \frac{D_w}{100 + \% \text{ moisture}} \times 100 \]

Where:

\[ D = \text{dry density of soil, lb./cu. ft.} \]
\[ D_w = \text{wet density of soil, lb./cu. ft.} \]

Example:

\[ D = \frac{106.3 \text{ lb./cu. ft.}}{113.6} \times 100 = 93.6 \text{ lb./cu. ft.} \]

**Maximum Density and Optimum Moisture**

8. The optimum moisture is determined as follows:

(a) The results of all samples are plotted on graph paper, using the percentage of moisture as the horizontal axis and the dry weight per cubic foot as the vertical axis.

(b) Draw a curve through the resulting points. The peak of the curve represents the "Maximum Density", in pounds per cubic foot, and the percentage of water at the peak represents the "Optimum Moisture Content" necessary to produce maximum density when compacted under these conditions.

Figure 2 gives a typical example of the calculations involved.
### Example of Calculations for ARIZ-226

#### Method "C & D" Proctor

<table>
<thead>
<tr>
<th>Approx. % Water Added</th>
<th>Wet Weight of Mold &amp; Soil (gm.)</th>
<th>Weight of Mold (gm.)</th>
<th>Wet Weight of Soil (gm.)</th>
<th>Factor</th>
<th>Wet Density lb./cu. ft.</th>
<th>Approx. Dry Density lb/cuft</th>
<th>Actual % Water Added</th>
<th>Actual Dry Density lb./cu.f</th>
<th><img src="chart.png" alt="Per Cent Moisture" /></th>
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</thead>
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<tr>
<td>10</td>
<td>3335</td>
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(Factor for Method "D" = 0.02939)

#### Moisture Chart

<table>
<thead>
<tr>
<th>Approx. % Water Added</th>
<th>Wet Weight (gm.)</th>
<th>Dry Weight (gm.)</th>
<th>Weight Water (j-k)=l</th>
<th>Water Added (l x 100) = h</th>
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</tr>
<tr>
<td>16</td>
<td>300</td>
<td>258</td>
<td>42</td>
<td>16.2</td>
</tr>
<tr>
<td>18</td>
<td>300</td>
<td>254</td>
<td>46</td>
<td>18.1</td>
</tr>
</tbody>
</table>
1. SCOPE

1.1 This test method describes the procedure for determining the maximum dry density and optimum moisture content for a soil by the Proctor Method A. Some materials may be more appropriately tested by Arizona Test Method 245, "Maximum Dry Density and Optimum Moisture of Soils by Proctor Alternate Method D ".

1.2 Method A may be used for all maximum dry density and optimum moisture content determinations except for volcanic cinders or light porous material on which the specific gravity cannot be determined with consistency or when the moisture absorption for the coarse aggregate is greater than 4.0%.

1.3 Method A may be used except when greater than 50% (60% for Aggregate Base) of the material is retained on the No. 4 sieve.

1.4 An example is provided in Figure 2 for the calculations and determinations referenced herein.

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

1.6 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.
2.2 4 inch proctor mold having a capacity of approximately 1/30 cubic foot, with an internal diameter of 4.000 plus 0.024 or minus 0.016 inches and a height of 4.584 plus 0.005 or minus 0.008 inches. The mold shall have a nominal wall thickness of 1/4 inch. It shall be equipped with an extension collar approximately 2-3/8 inches high. A baseplate as shown in Figure 1 shall be provided.

2.3 A hand or mechanical rammer weighing 5.50 ± 0.02 pounds, having a flat face, and equipped with a suitable arrangement to control the height of drop to a free fall of 12 ± 0.06 (1/16) inches above the elevation of the soil. The rammer face shall be circular with a diameter of 2.000 plus 0.010 or minus 0.015 inches. If a mechanical apparatus is used, it must be monitored through the ADOT proficiency sample program and maintain a rating of 3 or better based on the results of testing ADOT and AMRL proficiency samples.

2.4 Hard steel straightedge, at least 10 inches in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.01 inch per 10 inches (0.1 percent) of length with the portion used for trimming the soil.

2.5 Scale or balance capable of measuring the maximum weight to be determined, accurate to at least one gram.

2.6 Scale or balance capable of measuring the maximum weight to be determined, accurate to at least 0.1 gram.

2.7 Oven capable of maintaining a temperature of 230 ± 9 °F.

2.8 No. 4 and 3 inch sieves conforming to the requirements of ASTM E11.

2.9 Miscellaneous mixing tools and pans.

2.10 Sample Extruder (optional) consisting of a jack, lever, frame, or other device for extruding the compacted sample from the mold.

3. CALIBRATION OF MOLD

3.1 Molds shall be calibrated in accordance with APPENDIX A of this procedure.
4. SAMPLE

4.1 Enough soil material shall be provided from the field to make five compacted specimens. A minimum sample size of 20,000 grams (approximately 45 lbs.) is normally required.

4.2 If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of a slow fan or other drying apparatus such that the temperature of the sample does not exceed 140 °F.

4.3 Thoroughly break up the aggregations in such a manner as to avoid reducing the natural size of individual particles.

4.4 Weigh out an approximate 20,000 gram sample of representative soil. Record the weight of the sample, and sieve the material over a No. 4 sieve. If the percentage of coarse aggregate or rock retained on the No. 4 sieve is not already known from gradation testing, save any material retained on the No. 4 sieve and weigh. Calculate the percent of coarse aggregate or rock particles retained on the No. 4 sieve according to the following equation:

\[
\text{PR4} = \frac{\text{WR4}}{\text{WT}} \times 100
\]

Where:

- PR4 = Percentage of coarse aggregate or rock particles retained on the No. 4 sieve
- WR4 = Weight of coarse aggregate or rock particles retained on the No. 4 sieve
- WT = Total weight of material sieved

4.5 If "PR4" is greater than 50% (60% for Aggregate Base), Alternate Method D, Arizona Test Method 245, shall be used to determine the maximum dry density. If "PR4" is less than or equal to 50% (60% for Aggregate Base), blend material passing the No. 4 sieve thoroughly and proceed to Section 5 of this test method. If a specific gravity and absorption determination, in accordance with AASHTO T 85, is to be made for the plus No. 4 material, save an adequate amount of this material, otherwise, discard it.
5. PROCEDURE

5.1 From the thoroughly blended passing No. 4 material from Subsection 4.5, split out 5 representative approximate 2500 gram samples.

5.2 Select one sample and thoroughly mix with sufficient water to dampen it to approximately three percentage points below optimum moisture content.

Note: If desired, an additional three samples may be mixed at this time with approximate moisture contents of 1% below optimum, 1% over optimum, and 3% over optimum. The moisture in each of these samples shall be retained by covering with a damp cloth or being sealed in air tight containers until they are compacted. One of the five samples should be retained for future use since it is necessary to have at least two points defined on each side of the moisture-density curve.

5.3 Heavy clay soils or materials which tend to break down, or those in which it is difficult to incorporate water, shall require approximately 12 hours for uniform moisture absorption to be achieved. This shall be accomplished by preparing separate samples for each increment of water to be added, and then placing and sealing these samples in air tight containers for the 12-hour period.

5.4 Form a specimen by compacting the prepared soil in the four inch mold (with extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. Compact each layer with 25 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation.

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

5.5 When compacting granular, free-draining materials, at moisture contents which are at or above optimum, the mold shall be prepared by first sealing the bottom of the mold with waterproofing grease. All excess grease shall be wiped from the mold and baseplate.
5.6 Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the soil in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted soil even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Determine the weight of compacted specimen and mold. Determine the wet density, "WD", of the compacted soil by the following:

\[
WD = \frac{M_1 - M_2}{VM \times 453.6 \text{ (grams/lb.)}}
\]

Where:
- \( WD \) = Wet density of compacted soil, lb./cu. ft.
- \( M_1 \) = Weight of compacted specimen and mold, grams
- \( M_2 \) = Weight of the mold, grams
- \( VM \) = Volume of the mold, cu. ft. (See Section 3 of this procedure.)

5.7 The estimated dry density, "ED", of the compacted soil may be calculated and recorded to the nearest 0.1 lb./cu. ft. as follows:

\[
ED = \frac{WD}{(\text{Approx. } \% \text{ of water added}) + 100} \times 100
\]

Where:
- \( ED \) = Estimated dry density of compacted soil, lb./cu. ft.
- \( WD \) = Wet density of compacted soil, lb./cu. ft.

**Note:** These estimated densities are approximate only and will be corrected when final moisture results are obtained.

5.8 The estimated dry density is useful in deciding how much water to add in later trial batches, if the procedure described in the note following Subsection 5.2, for initially preparing three additional samples with varying moisture contents is not utilized. By carefully observing the estimated dry density of the compacted samples, the operator should be able to vary the moisture content as the test proceeds so that when the final moisture-density results are plotted, a curve will result that rises to a peak and then falls away.
5.9 Remove the material from the mold and slice vertically through the center. Take a representative minimum 300 gram sample from the full length and width of one of the cut faces. Weigh immediately, and dry to a constant weight in an oven at $230 \pm 9 \, ^\circ F$ to determine the moisture content in accordance with AASHTO T 265. Record the weight of wet soil to the nearest 0.1 gram as "WW", and record the weight of oven dry soil to the nearest 0.1 gram as "DW".

5.10 For granular, free-draining materials, the moisture content shall be determined using the entire compacted proctor specimen. Determine the weight of wet soil, "WW", by subtracting the weight of the mold, "M2", from the weight of compacted specimen and mold, "M1". Record the weight of wet soil, "WW", and the weight of oven dry soil, "DW", to at least the nearest gram. All clinging material shall be removed from the inside of the mold and included with the specimen. To facilitate drying, the specimen may be broken up and spread out in a large pan, being careful to not lose any soil particles.

5.11 Select another of the samples which was split in Subsection 5.1, and if not already done, thoroughly mix with water in sufficient amount to increase the moisture content by approximately two percentage points.

5.12 Repeat the procedure in Subsections 5.3 through 5.10 for the sample at each moisture content, as necessary to establish a moisture-density curve which rises to a peak and then falls away.

6. **CALCULATIONS**

6.1 Calculate percent moisture and record to the nearest 0.1 percent by the following:

$$\text{% Moisture} = \frac{\text{WW} - \text{DW}}{\text{DW}} \times 100$$

Where:  
WW = weight of wet soil, grams  
DW = weight of oven dry soil, grams
6.2 Calculate dry density and record to the nearest 0.1 lb./cu. ft. by the following:

\[
DD = \frac{WD}{\% \text{ Moisture} + 100} \times 100
\]

Where:
- DD = Dry density of compacted soil, lb./cu. ft.
- WD = Wet density of compacted soil, lb./cu. ft.

7. **MOISTURE-DENSITY RELATIONSHIP**

7.1 The percent moisture and corresponding dry density for each of the compacted soil specimens shall be plotted on the graph provided on the proctor density test form shown in Figure 3. For a good plot, the majority of the graph is utilized. Normally, three increments on the horizontal axis shall equal one percent of moisture, and three increments on the vertical axis shall equal one lb./cu. ft. of dry density. If another number of increments other than three is utilized, the number of increments for one percent moisture and one lb./cu. ft. dry density shall always be the same.

7.2 On each side of the maximum density curve, at least two points should be utilized to form two straight lines. The intersection point of these two lines defines the peak point of the density-moisture content relationship, or the maximum density and optimum moisture content for the soil. In general it will be found that higher unit mass soils assume steeper slopes with high maximum dry densities at low optimum moisture contents, while the lower unit mass soils assume flatter, more gently sloped lines with high optimum moisture contents and low maximum dry densities. Figure 4 gives examples of moisture-density plots which show the different slopes associated with different maximum dry density ranges.

7.3 Optimum moisture content - The percent moisture content corresponding to the peak (intersection point of the two lines) of the moisture-density curve shall be termed the "optimum moisture content", and shall be reported as "OM" to the nearest 0.1 percent.
7.4 Maximum dry density - The dry density at optimum moisture content corresponding to the peak (intersection point of the two lines) of the moisture-density curve shall be termed the "maximum dry density", and shall be reported as "MD" to the nearest 0.1 lb./cu. ft.

Note: The optimum moisture and maximum dry density determinations above are for the material passing the No. 4 sieve. When testing field samples for comparison to proctor optimum moisture and maximum dry density, a correction to the proctor optimum moisture and maximum dry density must be made, in accordance with ARIZ 227, for the percent rock which the field sample contains.

8. REPORT

8.1 Record the moisture and density data on the laboratory test form along with the laboratory number, material source and type, and other information required. A blank laboratory test form is provided in Figure 3.
APPENDIX A

CALIBRATION OF PROCTOR MOLDS

1. CALIBRATION

1.1 Molds shall be calibrated at least once a year, or sooner if there is reason to question the accuracy of the calibration.

1.2 Lightly coat the bottom of the mold with a waterproofing grease. (Dow Corning High Vacuum Grease, or similar, has proven satisfactory for this application.) For split molds, waterproofing grease is also necessary on the edges of the split mold halves which join together.

1.3 Fit mold into baseplate and secure snugly into place.

1.4 Wipe excess grease from the mold and the baseplate.

1.5 Record weight of baseplate, empty mold, and glass plate to at least the nearest 0.1 gram.

Note: An example which illustrates the recording of calibration data and calculations is shown in Figure 5. Figure 6 is a blank calibration form.

1.6 With the mold and baseplate assembly on a flat and level surface fill the mold with distilled water at room temperature $77 \pm 9^\circ\text{F}$.

1.7 Determine and record the temperature of the water to the nearest one degree Fahrenheit.

1.8 With a small rod, remove any air bubbles that may be clinging to the sides or bottom of the mold. Add additional water to completely fill the mold, using a glass plate in such a way to ensure accurate filling of the mold, eliminating air bubbles and excess water. Check bottom of mold to assure there is no leakage.

1.9 Dry the base plate, glass and outside of mold with a dry, absorbent cloth. Care must be taken to not lose water from inside of mold during drying. Record weight of baseplate, mold filled with water, and glass plate to at least the nearest 0.1 gram.
1.10 Determine the weight of water to fill mold by subtracting the weight of baseplate, empty mold, and glass plate from the weight of the baseplate, mold filled with water, and glass plate.

1.11 For the temperature of the water, determine its corresponding unit weight from the table below.

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Unit Weight of Water</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temp °F</td>
<td>lbs/cu. Ft.</td>
</tr>
<tr>
<td>68</td>
<td>62.315</td>
</tr>
<tr>
<td>69</td>
<td>62.308</td>
</tr>
<tr>
<td>70</td>
<td>62.301</td>
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<td>71</td>
<td>62.293</td>
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<tr>
<td>72</td>
<td>62.285</td>
</tr>
<tr>
<td>73</td>
<td>62.277</td>
</tr>
<tr>
<td>74</td>
<td>62.269</td>
</tr>
<tr>
<td>75</td>
<td>62.261</td>
</tr>
<tr>
<td>76</td>
<td>62.252</td>
</tr>
<tr>
<td>77</td>
<td>62.243</td>
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<tr>
<td>78</td>
<td>62.234</td>
</tr>
<tr>
<td>79</td>
<td>62.225</td>
</tr>
<tr>
<td>80</td>
<td>62.216</td>
</tr>
<tr>
<td>81</td>
<td>62.206</td>
</tr>
<tr>
<td>82</td>
<td>62.196</td>
</tr>
<tr>
<td>83</td>
<td>62.186</td>
</tr>
<tr>
<td>84</td>
<td>62.176</td>
</tr>
<tr>
<td>85</td>
<td>62.166</td>
</tr>
<tr>
<td>86</td>
<td>62.155</td>
</tr>
</tbody>
</table>

1.12 Calculate the volume of the mold, in cu. ft., as shown on the calibration form in Figures 5 and 6, and record to the nearest four decimal places.
1.13 Thoroughly clean grease from the mold and base plate. On the mold, record the identification of the mold, the date of calibration, and the volume of the mold.

1.14 Documentation of the calibration data shall be kept on file.

2. REQUIRED DOCUMENTATION

2.1 Record of weights, temperatures, and calculations required in the calibration procedure.

2.2 Identification of mold.

2.3 Date of calibration.

2.4 Volume of the mold.

2.5 Operator performing calibration.

2.6 Supervisor check of calibration data.

2.7 Date of calibration expiration.
FIGURE 1

CYLINDRICAL MOLD, COLLAR, AND BASEPLATE
(4 INCH MOLD)
## Arizona Department of Transportation

**METHOD A or ALTERNATE METHOD D PROCTOR DENSITY**

(Arizona Test Method 225 or 245)

<table>
<thead>
<tr>
<th>b</th>
<th>M1</th>
<th>c</th>
<th>WD</th>
<th>ED</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>3884</td>
<td>1914</td>
<td>125.6</td>
<td>113.2</td>
</tr>
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<td>13</td>
<td>3455</td>
<td>1985</td>
<td>130.2</td>
<td>115.2</td>
</tr>
<tr>
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<td>2025</td>
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<td>115.6</td>
</tr>
<tr>
<td>17</td>
<td>3486</td>
<td>2016</td>
<td>132.3</td>
<td>113.1</td>
</tr>
</tbody>
</table>

### METHOD A

\[
\text{WT} = \frac{21556}{\text{WR4}} = 4462
\]

\[
\text{PR4} = \frac{\text{WR4} \times 100}{\text{WT}} = 21 \%
\]

### ALTERNATE METHOD D

\[
\text{WT} = \frac{\text{WR 3/4}}{\text{WR 3/4}} =
\]

\[
\text{PR 3/4} = \frac{\text{WR 3/4} \times 100}{\text{WT}} = \%
\]

Coarse Aggregate Bulk Oven Dry Specific Gravity: 2.476

Coarse Aggregate Absorption: 1.93 \%

OPTIMUM MOISTURE CONTENT = OM = 13.4 %

MAXIMUM DRY DENSITY (lb./cu. ft.) = MD = 117.1

**Remarks:**

---

**FIGURE 2**
**METHOD A or ALTERNATE METHOD D PROCTOR DENSITY**
(Arizona Test Method 225 or 245)

<table>
<thead>
<tr>
<th>Weight of Mold = M2</th>
<th>Volume of Mold = VM</th>
<th>a = VM x 453.6 =</th>
</tr>
</thead>
<tbody>
<tr>
<td>b</td>
<td>M1</td>
<td>c</td>
</tr>
<tr>
<td>Approx % of water</td>
<td>Wt. of Sample</td>
<td>Wet density</td>
</tr>
<tr>
<td>Added</td>
<td>and Mold</td>
<td>lb/cu ft</td>
</tr>
<tr>
<td>M1 - M2</td>
<td>M1 - M2</td>
<td>c x 100</td>
</tr>
<tr>
<td></td>
<td></td>
<td>a x 100</td>
</tr>
</tbody>
</table>

**Moisture Determination**

<table>
<thead>
<tr>
<th>WW</th>
<th>DW</th>
<th>d</th>
<th>e</th>
<th>DD</th>
</tr>
</thead>
<tbody>
<tr>
<td>WW x 100</td>
<td>DW x 100</td>
<td>d x 100</td>
<td>e x 100</td>
<td>WW - DW</td>
</tr>
<tr>
<td>b + 100</td>
<td>c</td>
<td>a</td>
<td>a</td>
<td>a</td>
</tr>
</tbody>
</table>

**METHOD A**

WT = _______ WR4 = _______ WR4 x 100 = _______%
PR4 = _______ WT

**ALTERNATE METHOD D**

WT = _______ WR 3/4 = _______ WR 3/4 x 100 = _______%
PR 3/4 = _______ WT

Coarse Aggregate
Bulk Oven Dry Specific Gravity:

Coarse Aggregate Absorption: _______%

OPTIMUM MOISTURE CONTENT = OM = _______%
MAXIMUM DRY DENSITY (lb./cu.ft.) = MD = _______

Remarks:

Percent Moisture

FIGURE 3
### Clayey Sand

<table>
<thead>
<tr>
<th>% Moisture</th>
<th>Dry Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>13.7</td>
<td>108.1</td>
</tr>
<tr>
<td>15.5</td>
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<tr>
<td>17.3</td>
<td>110.6</td>
</tr>
<tr>
<td>19.4</td>
<td>107.6</td>
</tr>
</tbody>
</table>

Maximum Density = 111.6 lb/cu ft  
Optimum Moisture = 16.7%

### Fine Clay

<table>
<thead>
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<th>% Moisture</th>
<th>Dry Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>21.2</td>
<td>94.5</td>
</tr>
<tr>
<td>22.3</td>
<td>95.8</td>
</tr>
<tr>
<td>25.2</td>
<td>95.1</td>
</tr>
<tr>
<td>27.8</td>
<td>93.1</td>
</tr>
</tbody>
</table>

Maximum Density = 96.7 lb/cu ft  
Optimum Moisture = 23.1%
ARIZONA DEPARTMENT OF TRANSPORTATION

CALIBRATION OF PROCTOR MOLD
ARIZ 225 Appendix A

✓ Four Inch Mold       Six Inch Mold       Mold I. D. #: 4A

Calibration Date: 08/15/15    Calibration Expiration Date: 08/15/16

Temperature of water used for Calibration: 73 ° F

Unit Weight of Water: 62.277 lb./cu. ft.

Test Operator: Joe Tester    Supervisor and Date: Joe Supervisor 08/17/15

<table>
<thead>
<tr>
<th>Weight of Baseplate, Empty Mold, and Glass Plate (grams)</th>
<th>Weight of Baseplate, Mold Filled with Water, and Glass Plate (grams)</th>
<th>Weight of Water to Fill Mold (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4458.7</td>
<td>5407.9</td>
<td>949.2</td>
</tr>
</tbody>
</table>

\[ V = \frac{\text{Weight of Water to Fill Mold (grams)}}{\left( \text{Unit Weight of Water (lb./cu. ft.)} \times 453.6 \text{ (grams/lb.)} \right)} \]

\[ V = \frac{949.2}{62.277 \times 453.6} = 0.0336 \text{ cu. ft.} \]

REMARKS:

Unit Weight of Water Table

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>68</td>
<td>62.315</td>
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<tr>
<td></td>
<td></td>
<td>86</td>
<td>62.155</td>
</tr>
</tbody>
</table>

FIGURE 5
ARIZONA DEPARTMENT OF TRANSPORTATION

CALIBRATION OF PROCTOR MOLD
ARIZ 225 Appendix A

Four Inch Mold       Six Inch Mold       Mold I. D. #: _______________________

Calibration Date: ___________________ Calibration Expiration Date: ___________________

Temperature of water used for Calibration: ___________ ° F

Unit Weight of Water: ___________ lb./cu. ft.

Test Operator: ________________________ Supervisor and Date: ________________________

<table>
<thead>
<tr>
<th>Weight of Baseplate, Empty Mold, and Glass Plate (grams)</th>
<th>Weight of Baseplate, Mold Filled with Water, and Glass Plate (grams)</th>
<th>Weight of Water to Fill Mold (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>V = [ \frac{\text{Volume of Mold (cu. ft.)}}{} ] \times \frac{\text{Weight of Water to Fill Mold (grams)}}{\text{Unit Weight of Water (lb./cu. ft.)} \times 453.6 \text{ (grams/lb.)}} ]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

V = \( \frac{()}{()} \times () = () \text{ cu. ft.} \)

REMARKS:

Unit Weight of Water Table

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
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</tr>
<tr>
<td>75</td>
<td>62.261</td>
<td>84</td>
<td>62.176</td>
</tr>
<tr>
<td>76</td>
<td>62.252</td>
<td>85</td>
<td>62.168</td>
</tr>
<tr>
<td>77</td>
<td>62.243</td>
<td>86</td>
<td>62.155</td>
</tr>
</tbody>
</table>

FIGURE 6
ROCK CORRECTION PROCEDURE
FOR MAXIMUM DRY DENSITY AND
OPTIMUM MOISTURE CONTENT DETERMINATION

(A Modification of AASHTO Designation T 99 - ANNEX)

1. SCOPE

1.1 This procedure is used to determine the corrected maximum dry density and optimum moisture content for a sample of material when coarse aggregate or rock particles are retained on the No. 4 sieve (for Method A proctor) or 3/4 inch sieve (for Alternate Method D proctor).

1.2 The rock correction procedure shall not be used when the material consists of volcanic cinders or light weight porous material on which the specific gravity cannot be determined with consistency or when the moisture absorption for the coarse aggregate is greater than 4.0%.

1.3 The rock correction procedure shall not be used when the percent rock retained on the No. 4 for Method A is less than 10% or greater than 50% (greater than 60% in the case of an aggregate base material); or when the percent rock retained on the 3/4 inch sieve for Alternate Method D is less than 10% or greater than 40%.

2. NECESSARY INFORMATION

2.1 Refer to, or determine the maximum dry density and optimum moisture content of the material by Method A (Arizona Test Method 225 or 232); or Alternate Method D (Arizona Test Method 245 or 246).

2.2 For Method A, the percentage of rock particles in the sample which are coarser than the No. 4 sieve is recorded as "PR4". For Alternate Method D, the percentage of rock particles in the sample which are coarser than the 3/4 inch sieve is recorded as "PR3/4".

2.3 If not known, determine the bulk oven-dry specific gravity of the coarse aggregate in accordance with AASHTO T 85.
Note: Once determined for a source, the specific gravity may usually be used for all rock corrections on material from that source. Any slight change in specific gravity should not change the maximum dry density to a large extent. (A change of ± 0.02 could result in a change of ± 0.6 lb. per cubic foot.) Similarly, the maximum dry density of the passing No. 4 or passing 3/4 inch material, having been determined by Method A or Alternate Method D for a source, may usually be used for that source providing the sieve analysis of the passing No. 4 or passing 3/4 inch material and the Plasticity Index remain reasonably uniform.

3. ROCK CORRECTED MAXIMUM DRY DENSITY

3.1 The corrected maximum dry density of the total sample for the amount of rock (plus No. 4 for Method A proctor or plus 3/4 inch for Alternate Method D proctor) shall be determined by the following equation:

\[
CMD = \frac{\left(100 - PR\right) \times (MD) + \left(56.2\right) \times (PR) \times (SG)}{100}
\]

Where: CMD = Corrected maximum dry density of the total sample containing "PR" percent coarse rock particles, lbs./cu. ft.

PR = "PR4", percent rock retained on the No. 4 sieve for Method A; or "PR3/4" percent rock retained on the 3/4 inch sieve for Alternate Method D.

MD = Maximum dry density (Method A for plus No. 4; or Alternate Method D for plus 3/4 inch), lbs./cu. ft.

SG = Bulk O.D. Specific Gravity of the coarse aggregate.
Example (For Method A):

PR = PR4 = 29% rock retained on the No. 4 sieve.
MD = 114.0 lbs./cu. ft.
SG = 2.499

\[
\text{CMD} = \left( \frac{100 - 29}{100} \right) \times 114.0 + \left( \frac{29}{2.499} \right)
\]

\[
\text{CMD} = \left( \frac{71}{100} \right) \times 114.0 + \left( \frac{4072.9}{100} \right) = \frac{8094.0 + 4072.9}{100}
\]

CMD = 121.7 lbs./cu. ft. [Corrected maximum dry density of the total sample containing 29% rock retained on the No. 4 sieve.]

Example (For Alternate Method D):

PR = PR3/4 = 32% rock retained on the 3/4 inch sieve.
MD = 112.6 lbs./cu. ft.
SG = 2.526

\[
\text{CMD} = \left( \frac{100 - 32}{100} \right) \times 112.6 + \left( \frac{32}{2.526} \right)
\]

\[
\text{CMD} = \left( \frac{68}{100} \right) \times 112.6 + \left( \frac{4542.8}{100} \right) = \frac{7656.8 + 4542.8}{100}
\]

CMD = 122.0 lbs./cu. ft. [Corrected maximum dry density of the total sample containing 32% rock retained on the 3/4 inch sieve.]

4. **ROCK CORRECTED OPTIMUM MOISTURE CONTENT**

4.1 The rock corrected optimum moisture content of the total sample shall be determined by the following equation:

\[
\text{COM} = \left( \frac{(OM) \times (100 - PR) + (PR)}{100} \right)
\]
Where: COM = Corrected optimum moisture content for the total sample.

OM = optimum moisture content for Pass No. 4 (Method A) or pass 3/4 inch material (Alternate Method D).

PR = "PR4", % rock retained on the No. 4 sieve for Method A; or "PR3/4", % rock retained on the 3/4 inch sieve for Alternate Method D.

Example (For Method A):

OM = 14.3%
PR = PR4 = 29% rock retained on the No. 4 sieve.

\[
\text{COM} = \frac{[14.3 \times (100 - 29)] + (29)}{100}
\]

\[
\text{COM} = \frac{[14.3 \times (71)] + (29)}{100} = \frac{1015.3 + (29)}{100}
\]

COM = 10.4% [Corrected optimum moisture content of the total sample containing 29% rock retained on the No. 4 sieve.]

Example (For Alternate Method D):

OM = 15.2%
PR = PR3/4 = 32% rock retained on the 3/4 inch sieve.

\[
\text{COM} = \frac{[15.2 \times (100 - 32)] + (32)}{100}
\]

\[
\text{COM} = \frac{[15.2 \times (68)] + (32)}{100} = \frac{1033.6 + (32)}{100}
\]

COM = 10.7% [Corrected optimum moisture content of the total sample containing 32% rock retained on the 3/4 inch sieve.]
5. **USE OF RESULTS**

5.1 The corrected maximum dry density obtained by this method is used, in comparison with the field density of the material, to determine the percentage of compaction.

5.2 The corrected optimum moisture content is used to determine the amount of moisture which should be added to the material to achieve maximum dry density.

6. **REPORT**

6.1 Report the corrected maximum dry density to the nearest 0.1 lb./cu. ft.

6.2 Report the corrected optimum moisture content to the nearest 0.1 percent.
CALIBRATION OF STANDARD SAND AND SAND CONE APPARATUS

(A Modification of AASHTO Designation T 191)

1. SCOPE

1.1 This method of test is to determine a weight per cubic foot of sand to be used in soil density determinations, and the volume of the sand to fill the funnel and baseplate on the sand cone apparatus.

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

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Sand Cone Density Apparatus with Base Plate - A 1 gallon jar and a detachable appliance consisting of a cylindrical valve with an orifice 1/2 inch in diameter and having a small funnel connecting to a standard G mason jar cap on one end and a large funnel on the other end. The valve shall have stops to prevent rotating the valve past the completely open or closed positions. The plate shall have a flanged center hole to receive the large funnel. The apparatus shall conform to the requirements of AASHTO T 191.

2.3 Sand to be standardized, clean, dry, free flowing and uncemented

2.4 Calibrated 1/13.33 (0.0750) cubic foot mold

2.5 Straightedge
2.6 Balance, 20 kg capacity, accurate to 1.0 g, or 35 lb capacity, accurate to 0.01 lb.

3. PROCEDURE

3.1 For determining the density of the sand, the weight of sand to fill the mold is determined as follows:

3.1.1 Fill the jar with the sand to be used and attach the funnel.

3.1.2 Weigh the empty 0.0750 cubic foot mold and baseplate.

3.1.3 Set the mold and baseplate in a flat square pan large enough to catch any excess sand.

3.1.4 Invert the apparatus and set the funnel directly over the mold.

3.1.5 Open the valve and let the sand flow freely into the mold being careful not to jar the apparatus, until the sand ceases to move in the jar. Ensure there are no vibrations in the immediate vicinity.

3.1.6 Close the valve and carefully re-move the apparatus from the mold.

3.1.7 Using the straightedge, strike off the excess sand, being careful not to jar the mold. Work the straightedge in the least number of strokes possible until sand is level with the mold. Tap the side of the mold. Using a brush, clean off excess sand from mold and baseplate, and weigh and record.

3.1.8 Repeat steps 3.1.1 through 3.1.7 twice more.

3.2 For determining the volume of funnel and base plate, the weight of sand to fill the funnel and base plate is determined as follows:

3.2.1 The jar shall now be refilled with the sand, the funnel attached, and the apparatus weighed and recorded.

3.2.2 Place the base plate in the bottom of a level, smooth pan. Ensure that there is not a gap between the baseplate and the bottom of the pan.
3.2.3 Invert the apparatus and place on the base plate in the same manner as the apparatus would be placed over a field density hole.

3.2.4 Open the valve all the way and allow the sand to flow freely, being careful not to jar the apparatus, until the sand ceases to move. Ensure there are no vibrations in the immediate vicinity.

3.2.5 Close the valve and carefully remove the jar and record the weight.

3.2.6 Repeat steps 3.2.1 through 3.2.5 twice more.

4. **CALCULATION**

4.1 The calculations are as follows. See Figures 2 and 3 for example forms.

4.1.1 Density of Standard Sand ($D_s$), pcf:

\[
D_s = \frac{\text{Average of 3 Weights (grams)}}{(453.6 \text{ g/lb}) \times \text{(volume of mold)}}
\]

Example:

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>Wt. of Baseplate and Mold Filled with Sand (grams)</th>
<th>Wt. of Baseplate and Empty Mold (grams)</th>
<th>Wt. of Sand to Fill Mold (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>9,436</td>
<td>6,649</td>
<td>2,787</td>
</tr>
<tr>
<td>2</td>
<td>9,430</td>
<td>6,649</td>
<td>2,781</td>
</tr>
<tr>
<td>3</td>
<td>9,429</td>
<td>6,649</td>
<td>2,780</td>
</tr>
<tr>
<td></td>
<td>Total 8,348</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The Average of 3 Weights, determined in accordance with Subsections 3.1.1 through 3.1.8, is:

\[
\text{Average of 3 Weights} = \frac{8,348}{3} = 2,783
\]

Using the equation for $D_s$ (shown above this example), the Density of Standard Sand in this example is:

\[
D_s = \frac{2,783}{(453.6) \times (0.0750)} = 81.8 \text{ pcf}
\]
4.1.2 Volume of Funnel and Base Plate, cu.ft.:

\[
\text{Volume of Funnel and Base Plate (cu.ft.)} = \frac{\text{Average of 3 Weights (grams)}}{(453.6) \times (D_s)}
\]

Example:

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>Initial Wt. of Apparatus (grams)</th>
<th>Final Wt. of Apparatus (grams)</th>
<th>Wt. of Sand to Fill Funnel and Baseplate (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6,348</td>
<td>4,870</td>
<td>1,478</td>
</tr>
<tr>
<td>2</td>
<td>6,347</td>
<td>4,871</td>
<td>1,476</td>
</tr>
<tr>
<td>3</td>
<td>6,345</td>
<td>4,874</td>
<td>1,471</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td>4,425</td>
</tr>
</tbody>
</table>

The Average of 3 Weights, determined in accordance with Subsections 3.2.1 through 3.2.6, is:

\[
\text{Average of 3 Weights} = \frac{4,425}{3} = 1,475
\]

Using the equation for the Volume of Funnel and Base Plate (shown above this example), the Volume of Funnel and Base Plate in this example is:

\[
\text{Volume of Funnel and Base Plate (cu.ft.)} = \frac{1,475}{(453.6) \times (81.8)} = 0.0398 \text{ cu.ft.}
\]
Jar shall be of proportions such that there will be no shoulder void when determining the bulk density of sand.
CALIBRATION OF DENSITY SAND AND SAND CONE APPARATUS
ARIZ 229
(A Modification of AASHTO T191)

Date of Calibration: _______________ Test Operator: _______________
I. D. No. of Mold used in calibration: _____________________________
Volume of Mold used in calibration: ________________________________
Identification of Sand: ___________________________________________
Identification of Sand Cone Apparatus: ____________________________

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>Wt. of Baseplate and Mold Filled with Sand (grams)</th>
<th>Wt. of Baseplate and Empty Mold (grams)</th>
<th>Wt. of Sand to Fill Mold (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Average Weight of Sand to Fill Mold = _______________ grams

Density of Sand, $D_s = \frac{\text{Average Weight of Sand to Fill Mold}}{\frac{453.6 \text{ grams}}{\text{lb.}}} \times \frac{\text{Volume of Mold}}{\text{lb./cu. ft.}}$

$D_s = \frac{\text{Average Weight of Sand to Fill Mold}}{\frac{453.6 \text{ grams}}{\text{lb.}}} \times \frac{\text{Volume of Mold}}{\text{lb./cu. ft.}}$

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>Initial Wt. of Apparatus (grams)</th>
<th>Final Wt. of Apparatus (grams)</th>
<th>Wt. of Sand to Fill Funnel and Baseplate (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Average Weight of Sand to Fill Funnel and Baseplate = _______________ grams

Volume of Funnel and Baseplate, $V_{fb} = \frac{\text{Average Weight of Sand to Fill Funnel and Baseplate}}{\frac{453.6 \text{ grams}}{\text{lb.}}} \times \frac{\text{Density of Sand}}{\text{cu. ft.}}$

$V_{fb} = \frac{\text{Average Weight of Sand to Fill Funnel and Baseplate}}{\frac{453.6 \text{ grams}}{\text{lb.}}} \times \frac{\text{Density of Sand}}{\text{cu. ft.}}$

Remarks: __________________________________________________________

Supervisor and Date: _____________________________________________
Calibration Expiration Date: ________________________________

FIGURE 2
## Calibration of Density Sand and Sand Cone Apparatus

**ARIZ 229**  
*(A Modification of AASHTO T191)*

Date of Calibration: ___________  
Test Operator: ___________

1. D. No. of Mold used in calibration: **26**
2. Volume of Mold used in calibration: **0.0751**

Identification of Sand: 
Identification of Sand Cone Apparatus: 

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>Wt. of Baseplate and Mold Filled with Sand (grams)</th>
<th>Wt. of Baseplate and Empty Mold (grams)</th>
<th>Wt. of Sand to Fill Mold (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>9436</td>
<td>6649</td>
<td>2787</td>
</tr>
<tr>
<td>2</td>
<td>9430</td>
<td>6649</td>
<td>2781</td>
</tr>
<tr>
<td>3</td>
<td>9429</td>
<td>6649</td>
<td>2780</td>
</tr>
</tbody>
</table>

Average Weight of Sand to Fill Mold = **2783** grams

Density of Sand, \( D_s \) = \[ \frac{\text{Average Weight of Sand to Fill Mold}}{(453.6 \text{ grams} / \text{lb.}) \times (\text{Volume of Mold})} \]

\[ D_s = \frac{2783}{(453.6) \times (0.0751)} = 81.7 \text{ lb./cu. ft.} \]

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>Initial Wt. of Apparatus (grams)</th>
<th>Final Wt. of Apparatus (grams)</th>
<th>Wt. of Sand to Fill Funnel and Baseplate (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6348</td>
<td>4870</td>
<td>1478</td>
</tr>
<tr>
<td>2</td>
<td>6347</td>
<td>4871</td>
<td>1476</td>
</tr>
<tr>
<td>3</td>
<td>6345</td>
<td>4874</td>
<td>1471</td>
</tr>
</tbody>
</table>

Average Weight of Sand to Fill Funnel and Baseplate = **1475** grams

Volume of Funnel and Baseplate, \( V_{fb} \) = \[ \frac{\text{Average Weight of Sand to Fill Funnel and Baseplate}}{(453.6 \text{ grams} / \text{lb.}) \times (\text{Density of Sand})} \]

\[ V_{fb} = \frac{1475}{(453.6) \times (81.7)} = 0.0398 \text{ cu. ft.} \]

Remarks: ___________

Supervisor and Date: ___________
Calibration Expiration Date: ___________

**FIGURE 3**
FIELD DENSITY BY THE SAND CONE METHOD

(A Modification of AASHTO Designation T 191)

SCOPE

1. (a) This method is used to determine the density of compacted soils or aggregates by determining the weight and moisture content material removed from a test hole and measuring the volume of the test hole.

   (b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

APPARATUS

2. Apparatus for this test procedure shall consist of the following:

   (a) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.01 lbs or at least the nearest gram.

   (a) Miscellaneous digging tools.

   (b) Sand cone apparatus consisting of base plate, cone and sand jar.

   (c) Standard sand. (Sand shall be kept dry and free flowing).

   (d) Containers with air tight covers (cylinder molds are satisfactory).

   (e) Oven, hot plate, stove or Speedy Moisture Tester.
NOTE: Calibration of the sand and sand cone apparatus shall be done in accordance with AZ 229.

PREPARATION OF TEST SITE

3. The surface of the area where the test is to be conducted shall be prepared as follows:

   (a) Clean away all loose soil and rock from an area of about 3 feet square at the spot where the test is to be made. In areas compacted by ‘Sheep’s foot’ rollers, it is necessary to get below the depth of the ‘foot’ imprints.

   (b) The top of the material at the chosen location shall be prepared to a plane and level surface for an area slightly larger than the size of the base plate. The base plate shall then be placed on this level surface.

PROCEDURE

4. (a) A hole shall be dug approximately the diameter of the hole in the base plate and to the desired depth. (Usually 6 inches to 8 inches). While digging, especially using a hammer and chisel, care must be taken to avoid prying as this may deform the hole, disturb the surrounding material and give a false reading. All of the material removed from the hole shall be carefully recovered and put into a suitable container and covered with a lid or damp cloth, also making sure to get the hole as clean as possible. This operation shall be done as quickly as possible to avoid any excessive drying of the sample.

Suggested test hole volumes and corresponding moisture sample weights are given in Table 1. There will be occasions where the values listed in Table 1 will be difficult to arrive at or follow, such as in the case where we are limited to a shallow depth of compacted material. This table is offered as a guide and should be followed in most cases, however, deviations from these values are allowable when conditions warrant. The ‘Speedy’ Moisture Method (AASHTO T-217) may be used to determine the moisture content. The ‘Speedy’ Method will give the percent moisture on the passing the No. 4 material. If the sample contains material retained on the No. 4 sieve the ‘Speedy’ results must be adjusted in accordance with the following formula to obtain the percent moisture of the total sample.
\[
W = \frac{w(100 - R) + R}{100}
\]

Where:

\[
W = \% \text{ moisture in total sample}
\]
\[
w = \% \text{ moisture in Pass No. 4 material}
\]
\[
R = \% \text{ rock (Plus No. 4 sieve)}
\]

An example of this formula is shown under Calculations in this procedure.

<table>
<thead>
<tr>
<th>Maximum Particle Size Retained</th>
<th>Minimum Test Hole Volume</th>
<th>Minimum Moisture Content Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 4 sieve</td>
<td>0.060</td>
<td>100</td>
</tr>
<tr>
<td>½ in.</td>
<td>0.060</td>
<td>250</td>
</tr>
<tr>
<td>1 in.</td>
<td>0.075</td>
<td>500</td>
</tr>
<tr>
<td>2 in.</td>
<td>0.100</td>
<td>1000</td>
</tr>
<tr>
<td>2-1/2 in.</td>
<td>0.135</td>
<td>1500</td>
</tr>
</tbody>
</table>

(b) Weigh the filled sand cone apparatus and place over the base plate with the cone down. A match mark on the cone of the apparatus and the base plate is required to ensure that the apparatus is placed on the base plate the same way every time.

(c) Make sure there is no construction equipment operating in the immediate vicinity as any vibrations will cause a false volume determination.

(d) Open the valve all the way and let the sand flow freely, being careful not to jar the apparatus while the sand is flowing. When the sand ceases to move in the bottle, close the valve and remove the apparatus.

(e) Weigh the sand cone apparatus with the remaining sand to determine the volume of the hole.
REFERENCE TO METHOD ‘A’ PROCTOR

5. If referencing to Method ‘A’ Proctor continue as follows:

(a) Weigh the material removed from the test hole.

(b) Screen over a 3” and No. 4 sieve.

(c) With a small brush clean as many fines from the rock as possible.

(d) If any rock is retained on the 3” sieve, verify this with a sieve analysis and call this the end point. This sieve analysis shall be reported with a note stating the density is not determinable due to the rock retained on a 3” sieve.

(e) Weigh and record the weight of the material retained on the No. 4 sieve.

(f) Immediately weigh a moisture sample from the passing No. 4 material to be run either by ‘Speedy’ or Hot Plate Method.

(g) Determine the percent of rock by the following equation.

\[
% \text{ Rock} = \left( \frac{\text{wt. of +#4 material}}{\text{total wt. of material removed from hole}} \right) \times 100
\]

(h) If the rock content is greater than 50% (or 60% in the case of Aggregate Base) report the sieve analysis with a note stating that the density is not determinable due to excess rock.

Note: When conditions prevent density determination in areas due to the presence of excessive rock or rock retained on the 3” sieve, an attempt shall be made to compact these areas comparable to those surrounding locations where the required compaction was found through testing to be satisfactory.

(i) If less than 50% (or 60% in the case of Aggregate Base) is retained on the No. 4 sieve, proceed with the following calculations.
CALCULATIONS

6.  (a) Weight of sand, in lbs., to fill hole and funnel \(W_s\):

\[
W_s = \frac{W_o - W_f}{453.6 \text{ g/lb}}
\]

Where:

\(W_o\) = original wt. of sand and apparatus, g.
\(W_f\) = final wt. of sand and apparatus, g.

Example:

\[
\frac{(8560 \text{ g}) - (4314 \text{ g})}{453.6 \text{ g/lb}} = \frac{4246 \text{ g}}{453.6 \text{ g/lb}} = 9.36 \text{ lbs}
\]

(b) Volume, in cubic feet, of hole \((V)\):

\[
V = \frac{W_s}{D_s} - V_c
\]

Where:

\(W_s\) = wt. of sand to fill hole and funnel, lb.
\(D_s\) = density of sand, lb/cu. ft.
\(V_c\) = volume of cone and base plate

Example:

\[
\frac{(9.36 \text{ lbs})}{(96.4 \text{ lb/cu ft})} - (.0407 \text{ cu ft}) = .0564 \text{ cu ft}
\]
(c) Percent moisture of pass No. 4 material may be determined by utilizing the Speedy Test Method, (AASHTO T 217), or by oven-dry Method, (AASHTO T 265).

\[
w = \frac{W_w - W_d}{W_d} \times 100
\]

Where:
- \(W_w\) = weight of wet soil, g.
- \(W_d\) = weight of dry soil, g.
- \(w\) = % moisture in pass No. 4 material

Example:

\[
W = \frac{(322 \text{ g}) - (289 \text{ g})}{(289 \text{ g})} \times 100 = \frac{33 \text{ g}}{289 \text{ g}} = 11.4\%
\]

(d) Moisture content of the total sample expressed in percentage shall be calculated as follows:

\[
W = \frac{w(100 - R) + R}{100}
\]

Where:
- \(W\) = % moisture in total sample
- \(w\) = % moisture in Pass No. 4 material
- \(R\) = % rock (Plus No. 4 material)

Example:

\[
w = 11.4\%
R = 29\%
W = \frac{11.4 (100 - 29) + 29}{100} = \frac{838.4}{100} = 8.4\%
\]
The formula assumes that the rock has a moisture content of 1% and is sufficiently accurate to use in most cases. If the moisture content of the rock is appreciably above 2% as on absorbent materials, the central laboratory should be contacted for instructions.

(e) Wet density, $D_w$ in lb/cu. ft. of material:

$$D_w = \frac{W_t}{V}$$

Where:

$W_t =$ weight of total sample, lb.

Example:

$$D_w = \frac{7.41 \text{ lbs}}{0.0564 \text{ cu ft}} = 131.4 \text{ lb/ cu ft}$$

(f) Field dry density, $D_d$ in lb/ cu. ft, of material:

$$D_d = \frac{D_w}{100 + W} \times 100$$

Example:

$$D_d = \frac{131.4 \text{ lb/ cu ft}}{100 + 8.4} \times 100 = 121.2 \text{ lb/ cu ft}$$

(g) % compaction = $\frac{D_d \times 100}{\text{Maximum Density (Corrected)}}$

Example:

Maximum Dry Density (pass No. 4 material) = 114.0 lb/ cu ft

Percent rock = 29 %
Corrected Maximum Dry Density = 122.0 lb/ cu ft

Note: Compaction shall be reported to the nearest whole percent

% compaction = \frac{121.2 \text{ lb/ cu ft} \times 100}{122.0 \text{ lb/ cu ft}}

= 99 \%

REFERENCE TO ALTERNATE METHOD ‘D’ PROCTOR

7. If referencing to Alternate Method ‘D’ Proctor continue as follows:
   (a) Weigh the material removed from the test hole.
   (b) Screen over a 3” and No. 3/4 sieve.
   (c) With a small brush clean as many fines from the rock as possible.
   (d) If any rock is retained on the 3” sieve, verify this with a sieve analysis and call this the end point. This sieve analysis shall be reported with a note stating the density is not determinable due to the rock retained on a 3” sieve.
   (e) Weigh and record the weight of the material retained on the 3/4” sieve.
   (f) Screen the material that passes the 3/4” sieve over the No. 4 sieve. Weigh the material that is retained on the No. 4 sieve and add this weight to the weight of the material retained on the 3/4” sieve.
   (g) Immediately weigh a moisture sample from the passing No. 4 material to be run either by ‘Speedy’ or Hot Plate Method.
   (h) Determine the percent of rock by the following equation.

% Rock = \frac{\text{wt. of +3/4” material}}{\text{total wt. of material removed from hole}} \times 100
(h) If the rock content is greater than 40% report the sieve analysis with a note stating that the density is not determinable due to excess rock.

Note: When conditions prevent density determination in areas due to the presence of excessive rock or rock retained on the 3” sieve, an attempt shall be made to compact these areas comparable to those surrounding locations where the required compaction was found through testing to be satisfactory.

(i) If less than 40% is retained on the 3/4” sieve, proceed with the following calculations.

**CALCULATIONS**

8. (a) The calculations are the same as those for referencing to a Method ‘A’ Proctor. Section 6.
MOISTURE - DENSITY RELATIONSHIP USING
TYPICAL MOISTURE - DENSITY CURVES
(ONE POINT PROCTOR) METHOD A

(An Arizona Method)

SCOPE

1. (a) This method of test is for the determination of the optimum moisture content and maximum dry density of a soil or soil-aggregate mixture utilizing one moisture-density determination on the portion of the sample passing the No. 4 sieve. Some materials may be more appropriately tested by Arizona Test Method 246, “Moisture-Density Relationship using Typical Moisture-Density Curves (One Point Proctor) Alternate Method D”.

(b) The one-point proctor is used with the typical moisture-density curves, shown on the back of the One Point Proctor Density Test Card (Figures 1 and 2); or by utilizing a family of moisture-density curves developed for the immediate local conditions.

(c) This method is not to be used for volcanic cinders or light porous material on which the specific gravity cannot be determined with consistency or when the absorption of the coarse aggregate is greater than 4.0%.

(d) This method may be used to determine if an existing proctor maximum density determination is valid for the soil being tested. If the existing proctor maximum density determination is not valid, a full proctor according to Arizona 225 should normally be run to determine the maximum density required for that soil type.

(e) An example is provided in Section 7, and Figures 3 and 4, for the calculations and determinations referenced herein.

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

2. The apparatus shall consist of the following:

(a) The general apparatus utilized for this test method shall conform to the apparatus requirements of Arizona Test Method 225.

(b) In place of the oven listed in the general apparatus, a hot plate or stove capable of maintaining a temperature of approximately 230° F. may be used. A Speedy Moisture Tester with a conversion table or calibration curve may also be used for moisture determinations made in the field. Finally, a microwave oven may be used in accordance with Arizona Test Method 719.

(c) Instead of the scale or balance capable of measuring the weight to be determined to at least one gram, a scale capable of measuring the weight to at least 0.01 pound may be utilized.

CALIBRATION OF MOLD

3. Molds shall be calibrated in accordance with APPENDIX A of ARIZ 225.

SAMPLE

4. A representative sample of passing No. 4 material weighing approximately 2500 grams shall be obtained for each one-point proctor.

PROCEDURE

5. (a) The approximate 2500 gram sample of passing No. 4 material shall be thoroughly mixed with sufficient water to bring the sample to slightly less than its optimum moisture content.

(b) Form a specimen by compacting the prepared soil in the four inch mold (with extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. Compact each layer with 25 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation.
NOTE: Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: A block of concrete, weighing not less than 200 lbs., supported by a stable foundation; a sound concrete floor; and for field application, such surfaces as found in concrete box culverts, bridges, and pavements.

(c) When compacting granular, free-draining materials, at moisture contents which are at or above optimum, the mold shall be prepared by first sealing the bottom of the mold with waterproofing grease. All excess grease shall be wiped from the mold and base plate.

(d) Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the soil in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted soil even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Determine the weight of compacted specimen and mold. Determine the wet density, “WD”, of the compacted soil by the following:

\[
WD = \frac{M1 - M2}{VM \times 453.6 \text{ (grams/lb)}}
\]

Where:
- \(WD\) = Wet density of compacted soil, lb./cu. ft.
- \(M1\) = Weight of compacted specimen and mold, grams or lbs.
- \(M2\) = Weight of the mold, grams or lbs.
- \(VM\) = Volume of the mold, cu. ft. (See Section 3 of this procedure).

* If the weights of the compacted specimen and mold, \(M1\), and the empty mold, \(M2\), are measured in pounds, eliminate 453.6 (grams/lb.) from the denominator of the above equation.

(e) Remove the material from the mold and slice vertically through the center. Take a representative minimum 300 gram sample from the full length and width of one of the cut faces. Record the weight of wet soil to the nearest 0.1 gram as “WW”. Dry the sample to constant weight at approximately 230° F. and record weight of dry soil to the nearest 0.1 gram as “DW”. The percent moisture shall be recorded to the nearest 0.1 percent. The equation below is used when the percent moisture is determined by drying the sample. For testing performed in the field, the Speedy Moisture Tester (AASHTO T 217), may be used. For the Speedy Moisture Tester, a representative
sample of the pass No. 4 material shall be utilized and tested in accordance with the instructional manual for that apparatus.

\[
\text{% Moisture} = \frac{\text{WW} - \text{DW}}{\text{DW}} \times 100
\]

MAXIMUM DENSITY DETERMINATION

6. (a) The point representing the wet density and moisture content (dry basis) is then plotted on the Typical Moisture-Density Curves (Figure 2) and the maximum wet density and optimum moisture content determined. When this plotted point falls between two moisture-density curves, a minor interpolation is necessary. The maximum dry density in lb/cu. ft. and the corresponding percent optimum moisture is then read directly or interpolated from the chart. The family of typical moisture-density curves provided should be periodically verified for the local conditions. If it is ascertained that the family of curves provided by Figure 2 is of questionable reliability for given local conditions, then an independent family of curves should be established and used.

(b) The plotted point for wet density and moisture content should be on the dry side of the curve at or near optimum, as it is difficult to interpolate between curves for friable soils when on the wet side of the peak.

(c) If the plotted point representing the wet density and moisture content of the compacted material is on the right of the peak, the test should be repeated using a lower moisture content. An exception to this rule must be made for those soils having high clay contents and relatively flat curves. These soils cannot readily be dried to optimum in the field due to the creation of a cloddy condition which will cause voids in the proctor. Proctors for these materials should be made as near to optimum as possible.

EXAMPLE

7. An illustration of determining the maximum dry density and optimum moisture content is described below, and shown in Figures 3 and 4:

For:

\[
\begin{align*}
\text{Wet Density} & = 120.0 \text{ lb./cu. ft.} \\
\text{Moisture Content} & = 20.0\%
\end{align*}
\]
By plotting this point on the Typical Moisture-Density Curves and interpolating, it shows a point which is approximately 60 percent of the distance from Curve Q to Curve R. From the chart, the dry density for Curve Q is 102.4 @ 20.3% moisture and Curve R is 99.9 @ 21.5% moisture.

By interpolation:

Density: 102.4 - 99.9 = 2.5
   \[.60 \times 2.5 = 1.5 \text{ lb./cu. ft. difference}\]

Moisture: 21.5 - 20.3 = 1.2
   \[.60 \times 1.2 = 0.7\% \text{ difference}\]

Therefore:

Maximum dry density = 102.4 - 1.5
   = 100.9 lb./cu. ft.

Optimum Moisture = 20.3 + 0.7 = 21.0%

* As an alternate to performing the interpolation procedure, Table 1 below can be used to determine the maximum dry density and optimum moisture content when the plotted point falls between two moisture-density curves.

NOTE: The optimum moisture and maximum dry density determinations above are for the material passing the No. 4 sieve. When testing field samples for comparison to proctor optimum moisture and maximum dry density, a correction to the proctor optimum moisture and maximum dry density must be made, in accordance with ARIZ 227, for the percent rock which the field sample contains.

REPORT

8. Record the moisture and density data on the laboratory test form along with the laboratory number, material source and type, and other information required.
ARIZ 232b
November 14, 2008
Page 6
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TABLE 1

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# ONE POINT PROCTOR DENSITY

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<th>Org No.:</th>
<th>Date:</th>
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<td>Type of Material:</td>
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<td>Coarse Agg. Bulk O.D. Sp. Gr.:</td>
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<td>Alternate Method D</td>
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<td>Supervisor:</td>
<td>Date:</td>
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## Wet Density Determination

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<th>Volume of Mold</th>
<th>= VM =</th>
<th>Cu. ft.</th>
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<tr>
<td>Weight of Mold</td>
<td>= M2 =</td>
<td>grams, pounds</td>
</tr>
<tr>
<td>Weight of Sample and Mold</td>
<td>= M1 =</td>
<td>grams, pounds</td>
</tr>
</tbody>
</table>

\[
\text{Wet Density} = \frac{M1 - M2}{VM \times 453.6 \text{(grams/lb.)}} \text{lb./cu.ft.}
\]

*If M1 and M2 are in pounds, eliminate “453.6 (grams/lb.)” from denominator in above equation.

## Percent Moisture Determination

**For either Method A or Alternate Method D, when sample is oven dried:**

- Wet Weight of Moisture Sample = WW = grams
- Dry Weight of Moisture Sample = DW = grams

\[
\% \text{ Moisture} = \frac{WW - DW}{DW} \times 100 = \%
\]

**For Method A, when Speedie Moisture Tester is used:**

\[
\% \text{ Moisture} = \frac{WR4}{PR4} \times 100 = \%
\]

**For Alternate Method D, when Speedie Moisture Tester is used:**

\[
W = \frac{W(l00 - PR4) + PR4}{100}
\]

\[
\text{Total % Moisture} = TW = \frac{W(l00 - PR4) + PR4}{100} = \%
\]

## From Typical Moisture-Density Curves:

- Maximum Dry Density = MD = lb./cu. ft.
- Percent Optimum Moisture = OM = %

**REMARKS:**

---

**FIGURE 1**
TYPICAL MOISTURE-DENSITY CURVES

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<th>Max Dry Wt. lbs/cu.ft.</th>
<th>Optimum Moisture</th>
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<td>O</td>
<td>107.1</td>
<td>18.1</td>
</tr>
<tr>
<td>P</td>
<td>104.7</td>
<td>19.2</td>
</tr>
<tr>
<td>Q</td>
<td>102.4</td>
<td>20.3</td>
</tr>
<tr>
<td>R</td>
<td>99.9</td>
<td>21.5</td>
</tr>
<tr>
<td>S</td>
<td>97.4</td>
<td>22.7</td>
</tr>
<tr>
<td>T</td>
<td>94.6</td>
<td>24.4</td>
</tr>
<tr>
<td>U</td>
<td>92.1</td>
<td>25.8</td>
</tr>
<tr>
<td>V</td>
<td>89.9</td>
<td>27.4</td>
</tr>
<tr>
<td>W</td>
<td>87.5</td>
<td>29.5</td>
</tr>
<tr>
<td>X</td>
<td>85.0</td>
<td>30.5</td>
</tr>
<tr>
<td>Y</td>
<td>83.0</td>
<td>31.5</td>
</tr>
<tr>
<td>Z</td>
<td>81.1</td>
<td>32.5</td>
</tr>
</tbody>
</table>
ONE POINT PROCTOR DENSITY

Lab. No: ___________________ Org No.: ___________________ Date: ___________________
Project No. ___________________ Tracs No. ___________________ Date: ___________________
Original Source: ___________________ Type of Material: ___________________
Coarse Agg. % Absorp.: ___________________ Coarse Agg. Bulk O.D. Sp. Gr.: ____________
Proctor Method Used: Method A X Alternate Method D _____________________
Test Operator: ___________________ Date: ___________________
Supervisor: ___________________ Date: ___________________

Wet Density Determination

Volume of Mold = VM = 0.0335 Cu. ft.
Weight of Mold = M2 = ____________ grams 10.23 pounds
Weight of Sample and Mold = M1 = ____________ grams 14.25 pounds

Wet Density = WD = \( \frac{M1 - M2}{VM \times 453.6 (\text{grams/lb.})} \) = 120.0 lb./cu.ft.

*If M1 and M2 are in pounds, eliminate “453.6 (grams/lb.)” from denominator in above equation.

Percent Moisture Determination

For either Method A or Alternate Method D, when sample is oven dried:

Wet Weight of Moisture Sample = WW = 340.4 grams
Dry Weight of Moisture Sample = DW = 283.7 grams

% Moisture = \( \frac{WW - DW}{DW} \) x 100 = 20.0 %

For Method A, when Speedie Moisture Tester is used:

% Moisture = \( \frac{WR4}{WT} \) x 100 %

For Alternate Method D, when Speedie Moisture Tester is used:

WT = ____________ WR4 = ____________ PR4 = \( \frac{WR4}{WT} \) x 100 = ____________ %

% Moisture in Pass No. 4 material from Speedie = W = ____________ %

Total % Moisture = TW = \( \frac{W(l00 - PR4) + PR4}{100} \) = ____________ %

From Typical Moisture-Density Curves:

Maximum Dry Density = MD = 100.9 lb./cu. ft.
Percent Optimum Moisture = OM = 21.0 %

REMARKS: ___________________

FIGURE 3
**Figure 4**

**Typical Moisture-Density Curves**

<table>
<thead>
<tr>
<th>Curve</th>
<th>Max Dry Wt. lbs/cu. ft.</th>
<th>Optimum Moisture</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>141.8</td>
<td>6.6</td>
</tr>
<tr>
<td>B</td>
<td>139.1</td>
<td>7.2</td>
</tr>
<tr>
<td>C</td>
<td>136.3</td>
<td>7.9</td>
</tr>
<tr>
<td>D</td>
<td>134.1</td>
<td>8.5</td>
</tr>
<tr>
<td>E</td>
<td>132.1</td>
<td>9.0</td>
</tr>
<tr>
<td>F</td>
<td>129.3</td>
<td>9.7</td>
</tr>
<tr>
<td>G</td>
<td>126.6</td>
<td>10.5</td>
</tr>
<tr>
<td>H</td>
<td>124.2</td>
<td>11.2</td>
</tr>
<tr>
<td>I</td>
<td>121.7</td>
<td>11.9</td>
</tr>
<tr>
<td>J</td>
<td>119.3</td>
<td>12.7</td>
</tr>
<tr>
<td>K</td>
<td>117.0</td>
<td>13.5</td>
</tr>
<tr>
<td>L</td>
<td>114.6</td>
<td>14.6</td>
</tr>
<tr>
<td>M</td>
<td>112.0</td>
<td>15.8</td>
</tr>
<tr>
<td>N</td>
<td>109.6</td>
<td>16.9</td>
</tr>
<tr>
<td>O</td>
<td>107.1</td>
<td>18.1</td>
</tr>
<tr>
<td>P</td>
<td>104.7</td>
<td>19.2</td>
</tr>
<tr>
<td>Q</td>
<td>102.4</td>
<td>20.3</td>
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<tr>
<td>R</td>
<td>99.9</td>
<td>21.5</td>
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<tr>
<td>S</td>
<td>97.4</td>
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<tr>
<td>T</td>
<td>94.6</td>
<td>24.4</td>
</tr>
<tr>
<td>U</td>
<td>92.1</td>
<td>25.8</td>
</tr>
<tr>
<td>V</td>
<td>89.9</td>
<td>27.4</td>
</tr>
<tr>
<td>W</td>
<td>87.5</td>
<td>29.5</td>
</tr>
<tr>
<td>X</td>
<td>85.0</td>
<td>30.5</td>
</tr>
<tr>
<td>Y</td>
<td>83.0</td>
<td>31.5</td>
</tr>
<tr>
<td>Z</td>
<td>81.1</td>
<td>32.5</td>
</tr>
</tbody>
</table>

Interpolated Max. Dry Density: **102.4 (Curve Q)**

\[
\text{Approx. 60% Below Curve Q}
\]

\[
\frac{2.5 \times 0.60}{1.5} = 1.5 \text{ lb./cu. ft.}
\]

\[
102.4 - 1.5 = 100.9 \text{ lb./cu. ft.}
\]

Interpolated % Optimum Moisture: **21.5 (Curve R)**

\[
\frac{1.2 \times 0.60}{0.7} = 20.3 + 0.7 = 21.0%
\]
FLAKINESS INDEX OF COARSE AGGREGATE

(An Arizona Method)

1. SCOPE

1.1 This test method describes the procedure for determining the "Flakiness Index" (flatness) of coarse plus No. 8 aggregate. The Flakiness Index is the percentage of particles having a least dimension smaller than 60 percent of the mean size of each of one or more of the coarse sieve fractions. The lower the index for any sample of aggregate, the more nearly the aggregate particles approximate a cubical shape.

1.2 Once the Flakiness Index is known, the average least dimension of the aggregate can be determined if required, for example as in Arizona Test Method 819, “Design of Exposed Aggregate Seal Coats”. The procedure for determining the average least dimension of the aggregate is described in Section 6 of this test method.

1.3 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.4 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 For each size of material to be tested, a 16 gauge steel plate (1/16 inch nominal thickness) with a slotted opening approximately 4 inches in length and having a width conforming to the applicable requirements given below. If desired, the same plate may contain all or some of the required slots, rather than a separate plate for each.
Size of Material | Slot Width (inches) | Slot Width Tolerance (inches)
--- | --- | ---
1-1/2" | 0.738 | ± 0.023
1" | 0.520 | ± 0.016
3/4" | 0.372 | ± 0.012
1/2" | 0.260 | ± 0.008
3/8" | 0.187 | ± 0.006
1/4" | 0.131 | ± 0.004
No. 4 | 0.084 | ± 0.003

2.3 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least one gram.

2.4 Sieves conforming to the requirements of ASTM E11, of sizes 1-1/2", 1", 3/4", 1/2", 3/8", 1/4", No. 4, and No. 8, as necessary for the material being tested.

3. SAMPLE PREPARATION

3.1 A representative sample of the aggregate for the specified use shall be obtained. The size of sample shall be at least the size required by Arizona Test Method 201, and larger if necessary to provide adequate material for the applicable required individual size fractions indicated below. See Subsection 3.3 below for determination of the individual size fractions to be tested.

Size of Material | Minimum Test Sample (grams)
--- | ---
1-1/2" | 3000
1" | 1500
3/4" | 1000
1/2" | 500
3/8" | 200
1/4" | 100
No. 4 | 50
No. 8 | 50

3.2 The sample shall be subjected to sieve analysis in accordance with Arizona Test Method 201. If both the coarse and fine sieve analysis of the material are performed, material passing the No. 4 sieve shall be separated into No. 8 and
passing No. 8 fractions by sieving over a No. 8 sieve. Reference can also be made to Section 3 of Arizona Test Method 248 for utilizing a No. 8 sieve in the coarse sieving, and a fine sieve analysis not being required. The coarse aggregate size fractions of No. 8 and larger shall be placed in individual containers. The material passing the No. 8 sieve may be discarded.

3.3 From each size fraction that has a percent retained value from sieve analysis which is equal to or greater than 10%, obtain a representative test sample of the weight specified in Subsection 3.1 above.

4. FLAKINESS INDEX TEST PROCEDURE

4.1 Weigh each test sample to the nearest gram and record as the "Weight of Test Sample", for the respective size fraction.

4.2 The particles from the test sample for each size fraction shall be individually tested for their ability to pass through the appropriate slot, as specified in Subsection 2.2. Weigh the material which passes the appropriate slot, and record to the nearest gram as the "Weight Passing Slot", for the respective size fraction.

5. CALCULATIONS FOR FLAKINESS INDEX DETERMINATION

5.1 Figure 1 is an example of the calculations. Figure 2 is a blank Flakiness Index form which contains the required calculations.

5.2 Calculate the "Percent Passing Slot", for each respective size fraction, and record to the nearest percent.

5.3 Calculate the "Flakiness Index", and report the result to the nearest percent.

6. DETERMINATION OF AVERAGE LEAST DIMENSION OF AGGREGATE

6.1 When it is required (such as in Arizona Test Method 819), the average least dimension of the aggregate may be determined by using Figures 3 and 4. (These figures include an example which illustrates the procedure described in the paragraphs below.)
6.2 The median size of the aggregate is determined as shown in the example in Figure 3. Plot the % passing from sieve analysis for the two sieve sizes sufficient to locate the 50% line intercept. Draw a line between the two points. From the 50% passing point on the right side of the chart, proceed horizontally to the left until the line drawn between the two points is intercepted. Draw a line vertically from this point to intercept the "Median Size" scale at the bottom of the chart. Read the resultant median size to the nearest 0.01 inch.

6.3 The average least dimension of the aggregate is determined as shown in the example in Figure 4. Find the point for "Median Size" on the left side of the chart. Proceed horizontally to the right until the corresponding line for Flakiness Index is intercepted. Draw a line vertically from this point to intercept the "AVERAGE LEAST DIMENSION" at the bottom of the chart. Read and record the average least dimension of the aggregate to the nearest 0.01 inch.
## FLAKINESS INDEX CALCULATIONS

(AZARON TEST METHOD 233)

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>1-1/2&quot;</th>
<th>1&quot;</th>
<th>3/4&quot;</th>
<th>1/2&quot;</th>
<th>3/8&quot;</th>
<th>1/4&quot;</th>
<th>#4</th>
<th>#8</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Pass From Sieve Analysis</td>
<td>100</td>
<td>78</td>
<td>27</td>
<td>13</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>% Ret. From Sieve Analysis (F)</td>
<td>0</td>
<td>22</td>
<td>51</td>
<td>14</td>
<td>11</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight of Test Sample</td>
<td>611</td>
<td>263</td>
<td>138</td>
<td>78</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight Passing Slot</td>
<td>194</td>
<td>71</td>
<td>28</td>
<td>13</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Percent Passing Slot (P)

**NOTE:** Only the size fractions which have 10 or more percent retained are tested for passing the appropriate slot, and uses to determine the flakiness Index by the equation below.

\[
\text{Percent Passing Slot (P)} = \frac{\text{Weight Passing Slot}}{\text{Weight of Test Sample}} \times 100
\]

**FLAKINESS INDEX**

\[
\frac{[F \ 1-1/2" \times P \ 1-1/2"] + \ldots + [F \ No. 8 \times P \ No. 8]}{[F \ 1-1/2" + \ldots + [F \ No. 8]}
\]

**FLAKINESS INDEX**

\[
\frac{(22 \times 17) + (51 \times 27) + (14 \times 20) + (11 \times 24)}{(22) + (51) + (14) + (11)} = 23 \%
\]

**REMARKS:**

---

Test Operator and Date: **Joe Tester** 11/20/96  
Supervisor and Date: **Peter Headman** 11/20/96
### FLAKINESS INDEX CALCULATIONS

**ARIZONA TEST METHOD 233**

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>1-1/2&quot;</th>
<th>1&quot;</th>
<th>3/4&quot;</th>
<th>1/2&quot;</th>
<th>3/8&quot;</th>
<th>1/4&quot;</th>
<th>#4</th>
<th>#8</th>
</tr>
</thead>
</table>

- % Pass from Sieve Analysis
- % Ret. From Sieve Analysis (F)
- Weight of Test Sample
- Weight Passing Slot
- * Percent Passing Slot (P)

**NOTE:** Only the size fractions which have 10 or more percent retained are tested for passing the appropriate slot, and uses to determine the flakiness index by the equation below.

\[
\text{Percent Passing Slot (P)} = \frac{\text{Weight Passing Slot}}{\text{Weight of Test Sample}} \times 100
\]

**FLAKINESS INDEX**

\[
\frac{[F \: 1-1/2" \times P \: 1-1/2"] + \ldots + \ldots + [F \: \text{No. 8} \times P \: \text{No. 8}]}{[F \: 1-1/2" + \ldots + \ldots + F \: \text{No. 8}]}
\]

**FLAKINESS INDEX**

\[
\frac{x}{+} + \frac{x}{+} + \frac{x}{+} + \frac{x}{+} = \text{______} \%
\]

**REMARKS:**


Test Operator and Date: ___________________________  Supervisor and Date: ___________________________
FIGURE 4
FIELD DENSITY AND MOISTURE CONTENT
OF SOIL AND SOIL-AGGREGATE
MIXTURES BY THE NUCLEAR METHOD
(An Arizona Method)

SCOPE

1. (a) This method is used to determine the in-place density and
moisture content of compacted soil and soil-aggregate mixtures to a depth of 12
inches with a nuclear gauge. This test method is especially suited for soils of a
specified gradation such as aggregate base course, or fine soils of a consistent
nature. Rock correction of the Proctor maximum density requires excavation of
the soils at the test site to determine rock content, which increases the time
required for the test and decreases test efficiency. This method is acceptable for
normal soil and aggregate density testing, including pumping and heaving soils,
but should not be used for open-graded aggregate.

(b) An example is provided in Figure 1 for the calculations and
determinations referenced herein.

SAFETY

2. (a) This test method involves hazardous materials, operations, and
equipment. This test method does not purport to address all of the safety
problems associated with its use. It is the responsibility of whomever uses this
test method to consult and establish appropriate safety and health practices and
determine the applicability of regulatory limitations prior to use.

(b) Safety procedures for operation, transport and storage of nuclear
gauges shall be in accordance with the manufacturer's recommendations and the
applicable regulations of the Arizona Radiation Regulatory Agency (ARRA).

APPARATUS

3. The apparatus shall consist of the following:
(a) Moisture/Density Nuclear gauge capable of determining densities by the direct transmission method. Calibration of the gauge shall be performed in accordance with AASHTO T238, Subsection 3.1., on an annual basis. Adjustment of the calibration curve for field soil conditions shall be done as needed in accordance with AASHTO T238, Subsection 3.3.

(b) Reference standard block or test stand to obtain standard counts for moisture and density which are used to check the gauge stability.

(c) Nuclear gauge transport case and labels which comply with A.R.R.A. Regulations.

(d) Charging cord, if applicable.

(e) Scraper plate and drill rod guide.

(f) Drill rod.

(g) Hammer for driving the drill rod to make the hole for the direct transmission probe.

(h) A #10 sieve or a supply of dry, fine-sieved sand to be used as a sand blanket for surface irregularities at the test site and a fine brush for sand removal.

(i) A #4 and 3/4 inch sieve for use in removing plus #4 or plus 3/4 inch material for rock correction, depending upon the proctor method.

(j) A 3 inch sieve for determining the presence of oversize rock material.

(k) Safety goggles for eye protection, steel-toed footwear, and the radiation exposure badge.

(l) Information packet for the nuclear gauge which shall contain the following items:

1) Moisture/Density Calibration Tables (if required), and a standard count log book.

3) Applicable documentation necessary to meet requirements of ARRA for gauge safety.

4) Blank test forms for use on the applicable nuclear gauge (See Figure 2).

   (m) Calculator for necessary computations.

   (n) Miscellaneous equipment including watch, pencils, writing paper, ruler, eraser, clip board, and hand cart as required.

**GAUGE STABILITY CHECK**

4. A density standard count and moisture standard count shall be taken at the beginning of each day of testing at the project where the field density testing is to be performed. The gauge stability check shall be performed as follows:

   (a) Place the reference standard block on any asphalt, concrete, compacted aggregate or similar surface which is dry and level. The reference standard block should be at least 24 inches away from any vertical projection, at least 15 feet away from any large object, or vehicle, and at least 50 feet away from another nuclear gauge.

   (b) Seat the nuclear gauge on the reference block in accordance with the gauge operation manual. It is very important that the gauge is seated properly on the standard reference block.

   (c) Remove the lock on the source handle and make sure the source handle is in the safe or stored position (the top notch on the index rod).

   (d) Turn the gauge on (in standby power condition) and allow it to warm-up, if necessary, for the recommended time as given in the gauge operation manual, normally 15 minutes.

   (e) After the warm-up period, take a standard moisture count and a standard density count.

   (f) Record the moisture and density standard counts in the proper columns of the standard count log book along with the appropriate additional information, such as date, time, temperature, and location.
(g) Return the gauge to the standby power condition. The gauge should be left in the standby mode for subsequent testing.

(h) Determine if the standard counts are within the limits for normal operation in accordance with the gauge operation manual. This is usually done by comparing the standard counts to the average of the four previous standard counts or utilizing an internal statistical test which is available on some gauges. Additional standard counts may be necessary if initially the gauge does not appear to be operating properly. If the gauge does not meet the normal operating parameters as specified by the Standard Count procedure in the gauge operation manual, the gauge should not be used for testing. It should be sent in for servicing to determine the problem.

NOTE: Some gauges will store standard counts for later use in calculations performed by the gauge itself. The most recent standard counts will usually be stored automatically over preexisting standard counts.

(i) On a weekly basis, compare the average of the four most recent standard counts with the average of four standard counts immediately after gauge calibration or at least three months previous, whichever is shorter. If the accumulative shift in standard count exceeds 2% for moisture or 3% for density, the nuclear gauge should be recalibrated.

SITE PREPARATION

5. (a) Select a location for the field density test at random where the gauge will be at least 24 inches from any vertical projection such as a trench wall, retaining wall or pipe, at least 15 feet away from any vehicle, and at least 50 feet away from another nuclear gauge. If within 24 inches of a vertical projection, refer to section 8, Trench Correction.

(b) Remove all loose, disturbed, and excess material as necessary to reach the top of the compacted lift to be tested. Prepare a horizontal area sufficient in size to accommodate the gauge using the scraper plate supplied with the gauge. Plane the area to a smooth condition removing loose stones to obtain maximum contact between the gauge and the soil or aggregate being tested. Make sure the gauge sits solidly on the test site without rocking.

(c) Use native fines which pass a #10 sieve, or fine dry sand to fill voids only, and level the excess with the scraper plate. The total area of voids filled with fines or sand should be minimized as much as possible.
OBTAINING NUCLEAR MOISTURE AND DENSITY COUNTS

6. (a) Nuclear Density Counts and Moisture Counts shall be obtained by inserting the probe into the soil at the test site. (Refer to the gauge operation manual.) Prior to density count determination, select the mode of testing as follows based on the lift thickness of the soil or aggregate being tested:

<table>
<thead>
<tr>
<th>LIFT THICKNESS &quot;T&quot; INCHES</th>
<th>TRANSMISSION MODE</th>
</tr>
</thead>
<tbody>
<tr>
<td>T &lt; 2</td>
<td>Backscatter</td>
</tr>
<tr>
<td>2 ≤ T &lt; 4</td>
<td>Direct - 2 inch</td>
</tr>
<tr>
<td>4 ≤ T &lt; 6</td>
<td>Direct - 4 inch</td>
</tr>
<tr>
<td>6 ≤ T &lt; 8</td>
<td>Direct - 6 inch</td>
</tr>
<tr>
<td>8 ≤ T ≤ 12</td>
<td>Direct - 8 inch</td>
</tr>
<tr>
<td>12 &lt; T</td>
<td>See 6(b) Below</td>
</tr>
</tbody>
</table>

(b) Tests which require the density for a lift greater than 12 inches in thickness require an initial surface test to determine the density of the upper portion of the lift. Then the soil must be excavated downward to allow another test or tests to determine the density of the lower portion of the lift.

(c) To prepare the gauge for direct transmission testing, place the scraper plate drill rod guide on the test site so that the access hole for the probe will be at the desired location.

(d) Securely hold the scraper plate in place while driving the drill rod into the material. The hole should be at least 2 inches deeper than the depth to be tested. Safety goggles and steel-toed footwear should be worn while driving the drill rod. Note the depth marks on the drill rod. It would be desirable to turn the drill rod slightly after every other blow to allow easier removal.

(e) Remove the drill rod by pulling straight up in order to avoid disturbing the access hole.

(f) Remove the scraper plate and clear away all loose surface material. Using fine material as stated previously, fill any voids caused by driving the drill rod.

(g) Carefully place the gauge over the access hole and extend the probe into the hole to the desired direct transmission mode depth according to the lift thickness as outlined previously. Do not force the probe into the hole. If the probe will not extend into the hole, pull the probe back up. Lift up the gauge and check for a probe imprint. This will help determine if a slight change in the
position of the gauge is necessary to allow the probe to enter the hole. Once the probe is in the hole, gently push it down. Some minor shifting of the gauge may be required to extend the probe in gravelly soils. However, if an obstruction is encountered, it may be necessary to use the drill rod again to open up the hole.

(h) Once the probe is fully extended to the direct transmission depth, pull the gauge firmly, toward the scaler or readout end of the gauge, so that the probe is in firm contact with the soil or aggregate on the scaler side of the hole.

(i) Take density and moisture counts for a minimum one minute time period by pressing the proper button on the gauge. Both counts normally will occur simultaneously during the count period (Refer to the gauge operation manual).

(j) After the count period, press the proper button to obtain the moisture count. Record the moisture count on the data sheet, as "MC". Also, in a similar manner, obtain the density count. Record the density count on the data sheet, as "DC". It is also possible to get a wet density and moisture content readout in lbs./cu.ft. at this time for gauges capable of storing the standard counts and calculating these values (Refer to the gauge operation manual). If so, obtain these values and record on the test data sheet as "WD" and "M" respectively. The gauge may also be capable of calculating the moisture content in %, "M%", and the dry density in lbs./cu.ft., "DD". If so, these values may be recorded on the test data sheet. However, the gauge internal calculations may require some additional corrections as required by the test location or soil type. Refer to Sections 8 and 9 for further information on the trench and moisture corrections.

(k) In those instances when the soil or soil-aggregate mixture being tested in-place is not homogeneous and/or contains substantial variations in the rock content, it may be necessary to rotate the gauge 90 degrees at each test site and obtain an additional moisture and wet density reading at that position. If the new moisture and wet density readings differ by 5% or less from the original readings, the two readings may be averaged for use in later calculations. If they differ from the original readings by more than 5%, the gauge should be moved to a new test site.

**PROCTOR DENSITY ROCK CORRECTION**

7. (a) For determining the percent compaction based on the maximum proctor densities determined by Method A (Arizona Test Method 225 or 232) or Alternate Method D (Arizona Test Method 245 or 246), a rock correction may be
required for the amount of plus No. 4 (Method A) or plus 3/4 inch (Alternate Method D) rock or coarse aggregate in the material tested by the nuclear gauge.

(b) If it appears that 10% of the material will be retained on the No. 4 sieve (Method A) or 3/4 inch sieve (Alternate Method D), excavate the area occupied by the base of the gauge at the testing location to the depth of the test, which is normally the lift thickness. If an average rock correction is to be utilized, refer to 7(f).

(c) Obtain a minimum 3000 gram (7 pound) sample of excavated material and weigh to the nearest gram. Record the weight of this sample as "A". Sieve this material first over a 3 inch sieve to determine the presence of any oversize rock material. If any oversize rock is encountered, proceed to 7(d). If not, sieve the material over a No. 4 sieve (Method A) or 3/4 inch sieve (Alternate Method D), and record the weight of retained material as "B". Proceed to 7(e).

(d) If any rock is retained on the 3 inch sieve, this shall be reported with a note stating that the density is not determinable due to rock being retained on the 3 inch sieve. Additional attempts should be made to locate an area where the test can be accomplished.

(e) Calculate the percent of coarse particles, "PR" according to the following equation:

\[
PR = \frac{B}{A} \times 100
\]

Where: 

- \( PR \) = Percent of coarse aggregate or rock particles retained on either the No. 4 sieve for Method A or the 3/4 inch for Alternate Method D.
- \( B \) = Weight of coarse aggregate or rock particles retained on the No. 4 sieve (Method A) or 3/4 inch sieve (Alternate Method D).
- \( A \) = Weight total of sample which is sieved.

(f) If the material has from 10 to 50 percent (10 to 60 percent in the case of an Aggregate Base) retained on the No. 4 sieve (Method A) or 10 to 50 percent retained on the 3/4 inch sieve (Alternate Method D), the maximum proctor density will require a rock correction. Record the data at the appropriate locations on the data sheet.
(g) If the percent retained on the No. 4 sieve or 3/4 inch exceeds the maximum values listed above (50 or 60 percent), report the percent rock with a note stating that the density is not determinable due to excess rock. Additional attempts should be made to locate an area where a test can be accomplished. Also, if the proctor is a Method A proctor, an Alternate Method D proctor may be considered for this material.

Note: When conditions prevent the determination of density at a specific location, due to the presence of excessive rock or rock retained on the 3 inch sieve, the compactive effort in those areas should be the same as that performed in surrounding locations where the required density was found through testing to be satisfactory.

(h) If, from the results of a minimum of five samples, it is indicated that no rock is retained on the 3 inch sieve and the percent rock retained on the No. 4 or 3/4 inch sieve is within a 12% range (plus or minus 6% from the average), then an average percent of rock based on the five samples may be utilized. However, if a failing density test results from utilization of this average percent of rock, then the test site must be excavated and the actual percent of rock must be determined in accordance with 7(c) and 7(d) above. If there is a visible change in the material including the percent of rock, then a new proctor is required and a new average percent of rock must be determined.

**TRENCH CORRECTION**

8. (a) Moisture and dry density test results may be affected when a gauge is operated within 24 inches of a vertical projection such as a trench wall, pipe or retaining wall which contains moisture. The density counts, determined in the direct transmission mode, should not be affected, but moisture counts will possibly be affected. Without a trench correction, the moisture content determined could be higher than the actual moisture content which would cause the dry density determined to be too low. If the density test passes without a trench correction, then a trench correction would not be necessary.

(b) When a trench correction is necessary, refer to the gauge operation manual for the proper procedure in making a trench correction. Usually it is necessary that a moisture count be taken on the standard block in the trench at the same position and orientation that the moisture count for the density test is taken.
(c) Based on the original standard moisture count and the moisture count taken on the standard block in the trench, a trench correction may be made internally in most gauges by a calculation process or by an external manual moisture count shift. Refer to gauge operation manual for the proper input of the trench correction. Be sure to delete this correction after each test since each test location would have a different correction.

MOISTURE CORRECTION

9. (a) Moisture and dry density test results may be affected by hydrogen in the soil unrelated to actual moisture. Only the moisture reading is affected. The wet density reading is not affected.

(b) The moisture correction is obtained by determining the difference between the average oven dry moisture contents and the average nuclear gauge moisture contents for five or more samples obtained per paragraph 7(c). The oven-dry moisture content in percent, "TW", will be determined for each representative sample according to AASHTO T265. The sample shall include all plus No. 4 or 3/4 inch material, since the nuclear gauge measures total moisture. If the moisture is determined by the Speedy Moisture Tester and the sample contains material retained on the No. 4 sieve, the "Speedy" results must be adjusted in accordance with the following formula to obtain the percent moisture of the total sample.

\[
TW = \frac{[W(100 - PR)] + PR}{100}
\]

Where:
- \(TW\) = % moisture content in total sample
- \(W\) = % moisture in Pass No. 4 material
- \(PR\) = % rock retained on the No. 4 sieve

(c) A moisture correction is needed if the difference calculated indicates that the gauge moisture results are more than 1% higher or lower than the oven dry or Speedy results. Moisture corrections of 1% or less may be disregarded.

(d) Most gauges have the capability for correction of the moisture content in % by an internal calculation or an external moisture count shift. Simply apply the moisture correction up or down as applicable in the gauge according to the procedures in the gauge operation manual. Future density calculations within the gauge will apply this moisture count shift until it is deleted by the operator.
MOISTURE-DENSITY CALCULATIONS

10. Calculations shall be performed as follows:

(a) Record the standard counts for moisture and density on the data sheet in the spaces for "MS" and "DS" respectively.

(b) Most current gauges have the internal capability to calculate the moisture content in lbs./cu.ft. and %, the wet density in lbs./cu.ft., and the dry density in lbs./cu.ft. If so, record "M", "M%", "WD" and "DD" at the appropriate locations on the test data form. Also, list any corrections which have been input into the gauge and apply to these calculations. If it is necessary to use count ratios and calibration tables to determine the moisture content, wet density, and dry density, this should be done in accordance with the procedures in the gauge operation manual. Any data calculated in this manner should also be recorded on the test data sheet at the proper location.

(c) Determine from the correction data obtained in Section 7 whether a rock correction is necessary. If so, determine the rock corrected maximum density using the maximum proctor dry density and percent rock according to the Rock Correction Method, ARIZ 227.

(d) The percent of compaction, based on the maximum dry density (corrected, if necessary) is determined and reported to the nearest whole percent.

\[
\% \text{ compaction} = \frac{DD \times 100}{MD \text{ or CMD}}
\]

Where:

- \( MD \) = Maximum Proctor Dry Density (if no rock correction)
- \( CMD \) = Corrected Maximum Density (if rock correction is necessary).

NOTE: If a rock correction is not necessary, many current nuclear gauges have the capability to perform all of the other density and moisture calculations internally, with results shown in the display or output to an external printer or computer. Refer to the gauge operation manual to take advantage of all possible gauge options.
11. Record the pertinent moisture and density data on the test form along with the test number, location and other information required. An example is given in Figure 1. A blank test form is provided in Figure 2.
ARIZON A DEPARTMENT OF TRANSPORTATION
FIELD DENSITY/MOISTURE OF SOILS BY THE NUCLEAR METHOD
ARIZONA TEST METHOD 235

TRACS Number H 9999 01C
Project Code 8888
Date JAN. 13, 1993
Tested By JOE DOUGL
Material Type EMBANKMENT
Test Number 22
Proctor Number EM #1
Material Source INPLACE
Test Location S.E. #51, STA. 97+00, 60' CH. E

Density Standard (DS) 3239.4
Moisture Standard (MS) 415.1

Density Count (DC) 1645.2
Wet Density (WD) 126.7 PCF
Dry Density (DD) 124.0 PCF

ROCK CORRECTION [IF NECESSARY] (ARIZ 227)
Wet Wt. of Material (A) 24.72 lb
Wet Wt. Retained (B) 6.07 lb
Percent Rock (PR) = (B/A) x 100 = 25%

TRENCH CORRECTION [IF NECESSARY]
Trench Moist. Stand. (TMS) Trench Correction (TC)
Trench Moist. Count (TMC) TC = TMC - TMS =

MOISTURE CORRECTION [IF NECESSARY]
Moisture Correction Input into gauge [YES/NO]

PROCTOR DATA (ARIZ 232 or 245)
Max. Dry Density (MD) 121.0 PCF
Coarse Agg. Sp. Gr. (SG) 2.609
Corrected Maximum Dry Density (CMD) [For Rock Correction]:

CMD = [(100 - PR) x (MD)] + [(56.2) x (PR) x (SG)]
100

ONE POINT PROCTOR DATA (ARIZ 232 or 246)
Wt. of Mold and Soil (M1) 1lb
Mold Volume (VM) CF
Moisture Content %
Max. Dry Density (MD) PCF

COMPAC TION COMPLIANCE CALCULATION
Required Compaction 95 %
Pass V Fail
Percent Compaction = (DD x 100)/MD or CMD = 97.3 %

Remarks:

Lab Supervisor Ted Headman Date 1-13-93

FIGURE 1
ARIZONA DEPARTMENT OF TRANSPORTATION
FIELD DENSITY/MOISTURE OF SOILS BY THE NUCLEAR METHOD
ARIZONA TEST METHOD 235

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<tr>
<td>Material Source</td>
<td>Test Location</td>
</tr>
</tbody>
</table>

### STANDARD COUNTS

- Density Standard (DS)
- Moisture Standard (MS)

### FIELD COUNTS

- Density Count (DC)
- Moisture Count (MC)
- Wet Density (WD) PCF
- Moisture Content (M) PCF
- Dry Density (DD) PCF
- Moisture Content (Mₚ) %

### ROCK CORRECTION [IF NECESSARY] (ARIZ 227)

- Wet Wt. of Material (A)
- Percent Rock (PR)
- Wet Wt. Retained (B)

\[
PR = \frac{(B/A)}{100}
\]

### TRENCH CORRECTION [IF NECESSARY]

- Trench Moist. Stand. (TMS)
- Trench Correction (TC)

\[
TC = TMC - TMS
\]

### MOISTURE CORRECTION [IF NECESSARY]

- Moisture Correction %
- Input into gauge [YES/NO]

### PROCTOR DATA (ARIZ 225 or 245)

- Max. Dry Density (MD) PCF
- Optimum Moisture %
- Coarse Agg. Sp. Gr. (SG)
- Proctor & Rock Retained %
- Corrected Maximum Dry Density (CMD) [For Rock Correction]:

\[
CMD = \frac{[(100 - PR) \times (MD)] + [(56.2) \times (PR) \times (SG)]}{100}
\]

### ONE POINT PROCTOR DATA (ARIZ 232 or 246)

- Wt. of Mold and Soil (M₁) lb
- Wt. of Mold (M₂) lb
- Mold Volume (VM) CF
- Wet Density (M₁-M₂)/VM PCF
- Moisture Content %
- Density Curve I.D.
- Max. Dry Density (MD) PCF
- Corr. Max. Dry Dens. (CMD) PCF

### COMPACTION COMPLIANCE CALCULATION

- Required Compaction %
- Percent Compaction = 
  \[
  \frac{(DD \times 100)}{MD} \text{ or } CMD \times 100
  \]

### Remarks:

-
-

Lab Supervisor ___________________________ Date ____________

FIGURE 2
DETERMINING pH AND MINIMUM RESISTIVITY OF SOILS AND AGGREGATES

(An Arizona Method)

1. SCOPE

1.1 This test method outlines the procedure for determining the pH and minimum resistivity of soil and aggregate materials.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Drying Apparatus - Any suitable device capable of drying samples at a temperature not exceeding 140 °F.

2.3 100 mL glass beaker or other suitable non-metallic container.

2.4 200 mL minimum capacity cylinder graduated in 1 mL increments.

2.5 Distilled Water.

2.6 pH Meter and combination probe, both having a relative accuracy of ± 0.05 pH units, minimum. The pH meter must be capable of registering a minimum pH range of 4.0 - 10.0, and be equipped with temperature compensation controls.
2.7 Standard Buffer Solutions of pH values 4.0, 7.0, and 10.0. The expiration date of the solution shall be noted on the container. Solution shall not be used past the expiration date.

2.8 Soil Box - Designed for use with the resistance meter. An example of an acceptable soil box is shown in Figure 1. Soil boxes which are commercially available may be used provided they meet the requirements specified herein. Soil boxes may vary in size. A unique soil box factor is determined for each soil box.

2.9 Resistance Meter - Instrument to determine resistance (ohms), readable to at least the nearest 10 ohms.

**Note:** The accuracy of the meter shall be established at these measurement levels: 50, 100, 200, 500, 900, 1500 ohms. The resistance standard(s) will have a tolerance of ± 1%. Any deviation greater than 5% from the known resistance shall require either calibration to within specified limits or the development of a calibration curve. Calibration results shall be recorded, dated, signed, and maintained on file for review.

2.10 No. 8 sieve conforming to ASTM E11.

2.11 A Balance or Scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.12 Thermometer - A thermometer accurate to 1 °F.

2.13 Miscellaneous mixing tools and pans.

### 3. SAMPLE PREPARATION

3.1 The soil sample as received from the field shall be dried thoroughly in air or the drying apparatus at a temperature not exceeding 140 °F.

3.2 A representative test sample of approximately 2000 grams shall then be obtained by splitting or quartering.

3.3 Screen the test sample through a No. 8 sieve.
3.4 From the screened Pass No. 8 Material, split out approximately 1500 grams for the resistivity test and obtain approximately 50 grams for the pH test. Record the weight of the pH material to the nearest 0.1 gram.

3.5 Combine the pH sample with an equal weight of distilled water in the 100 mL beaker. Stir until well mixed into a slurry and then stir at regular intervals of 8 to 10 minutes for an hour.

3.6 The temperature of the standard buffer solutions must be within 2 °F of the pH sample. Also, temperatures of the standard buffer solutions and the pH sample must be within the manufacturer's recommended temperature compensation range of the pH meter.

3.7 Place the resistivity sample in a mixing bowl. Add approximately 200 mL of distilled water to moisten the sample and mix thoroughly.

4. TEST PROCEDURE FOR pH

4.1 Per the manufacturer's instructions, standardize the pH meter using two of the standard buffer solutions: 7.0 and either 4.0 or 10.0, whichever is nearest to the estimated pH of the sample.

4.2 For each standard buffer solution, measure its temperature and adjust the temperature controller of the pH meter before testing for the standard solution pH value. This may not be needed on meters with automatic temperature compensation; follow manufacturer's instructions.

4.3 Stir the slurry mixture in the 100 mL beaker. Measure the temperature of the slurry and adjust the temperature controller of the pH meter before testing for the pH value of the slurry. This may not be needed on meters with automatic temperature compensation; follow manufacturer's instructions.

4.4 Carefully insert the pH probe in the slurry mixture. Do not place the electrode(s) into the soil; place them only into the slurry mixture. Determine the pH reading when the meter reading stabilizes.

4.5 If the pH reading is unstable when the electrode is immersed in the slurry, leave the electrode immersed until the pH reading has stabilized. In some cases, the waiting period for stabilization of the pH reading may take up to 5 minutes.

4.6 Record the pH value of the slurry mixture, to the nearest tenth.
5. TEST PROCEDURE FOR MINIMUM RESISTIVITY

5.1 Place moistened soil in a soil box with a known soil box factor, compact lightly with fingers and level off the top with a straightedge. Connect the resistance meter to the side terminals of the box. Determine resistance as per the manufacturer’s instructions and record the resistance to the nearest 10 ohms.

5.2 Empty the soil back into the mixing bowl and add 50 mL of distilled water at room temperature and mix until all the water is dispersed uniformly through the soil.

5.3 Clean the soil box by rinsing with distilled water after each test application.

5.4 Fill the soil box by lightly hand compacting the wet soil, making sure that the soil completely fills the box. Level off the top of the hand compacted sample with a straightedge. Connect the resistance meter to the box. Read and record the resistance.

5.5 Repeat the above procedure, adding distilled water in increments of 50 mL. Ensure that each addition of water is dispersed evenly through the sample. The resistance readings should decrease, with the increase in moisture content, for several readings before an increase is noted. The lowest resistance reading before an increase is used for calculating the minimum resistivity of the soil.

6. CALCULATIONS FOR MINIMUM RESISTIVITY

6.1 Determine the length and width of the electrodes to the nearest 1 mm.

6.2 The soil box factor (SBF) expressed to the nearest 0.01 centimeter, is determined by the following equation:

\[
SBF = \frac{A}{D} \times 0.10
\]

Where:

- \( A \) = Area of one electrode, mm\(^2\)
- \( D \) = Distance between electrodes, mm
Example: (For electrodes having a length of 152 mm and a width of 45 mm, and at a distance between electrodes of 102 mm.)

\[ A = 152 \text{ mm} \times 45 \text{ mm} = 6840 \text{ mm}^2 \]
\[ D = 102 \text{ mm} \]

\[ \text{SBF} = \frac{6840}{102} \times 0.10 = 6.71 \text{ cm} \]

**Note:** The soil box factor will depend on the soil box dimensions.

6.3 The minimum resistivity is determined by the following equation:

\[
\begin{bmatrix}
\text{Minimum Resistivity} \\
\text{(ohm - cm)}
\end{bmatrix} = \begin{bmatrix}
\text{Resistance} \\
\text{(ohms)}
\end{bmatrix} \times \begin{bmatrix}
\text{SBF} \\
\text{(cm)}
\end{bmatrix}
\]

Example: (For a resistance reading of 480 ohms and a soil box factor of 6.71 cm.)

Minimum Resistivity = (480 ohms) x (6.71 cm)
= 3220.8 ohm-cm
= 3221 ohm-cm

6.4 The minimum resistivity value is reported to the nearest whole number.
EXAMPLE OF AN ACCEPTABLE SOIL BOX

Box Material [6 mm (1/4") Plastic]
- Bottom: 1 piece: 165 mm x 114 mm x 6 mm (6-1/2" x 4-1/2" x 1/4")
- Ends: 2 pieces: 114 mm x 45 mm x 6 mm (4-1/2" x 1-3/4" x 1/4")
- Sides: 2 pieces: 152 mm x 45 mm x 6 mm (6" x 1-3/4" x 1/4")

Electrodes [0.76 mm to 1.21 mm thickness (22 to 18 Gauge) Stainless Steel]
- 2 pieces: 152 mm x 45 mm (6" x 1-3/4")

Hardware
- 2 each No. 8-32 x ¾" (or longer), or metric equivalent, Stainless Steel or Brass Machine Screw with Washer and Nut

Gasket
- Rubber Washer

FIGURE 1
DETERMINING pH AND SOLUBLE SALTS OF SOILS

(An Arizona Method)

1. SCOPE

1.1 This test method describes the procedure for determining the pH and soluble salts content of soils.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Drying Apparatus - Any suitable device capable of drying samples at a temperature not exceeding 140 °F.

2.3 100 mL glass beaker or other suitable non-metallic container

2.4 250 mL glass beaker or other suitable non-metallic container

2.5 200 mL minimum capacity cylinder graduated in 1 mL increments

2.6 Richards or Buchner funnel

2.7 Vacuum Source

2.8 Conductance Meter
2.9 Conductivity Cell (Cell constant = 1.0/centimeter preferred)

2.10 Distilled Water

2.11 pH Meter and Combination Probe, both having a relative accuracy of ± 0.05 pH units, minimum. The pH meter must be capable of registering a minimum pH range of 4.0 - 10.0, and be equipped with temperature compensation controls.

2.12 Standard Buffer Solutions of pH values 4.0, 7.0, and 10.0. The expiration date of the solution shall be noted on the container. Solution shall not be used past the expiration date.

2.13 No. 8 sieve conforming to ASTM E11.

2.14 A Balance or Scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.15 Thermometer - A thermometer accurate to 0.5 °C (1 °F)

2.16 Miscellaneous mixing tools and pans

3. **SAMPLE PREPARATION**

3.1 The soil sample as received from the field shall be dried thoroughly in air or the drying apparatus at a temperature not exceeding 140 °F.

3.2 A representative test sample of approximately 500 grams shall then be obtained by splitting or quartering.

3.3 Screen the test sample through a No. 8 sieve.

3.4 From the screened Minus No. 8 Material, split out approximately 250 grams for the soluble salts test and approximately 50 grams for the pH test. Record the weight of the pH material to the nearest 0.1 gram.
3.5 Combine the pH sample with an equal weight of distilled water in the 100 mL beaker. Stir until well mixed into a slurry and then stir at regular intervals of 8 to 10 minutes for an hour.

3.6 The temperature of the standard buffer solutions must be within 2 °F of the pH sample. Also, temperatures of the standard buffer solutions and the pH sample must be within the manufacturer's recommended temperature compensation range of the pH meter.

3.7 Prepare a saturated soil paste from the soluble salts test sample by placing the soil in the 250 mL beaker. Begin adding distilled water and stirring with a spatula. From time to time tap the container on a work bench to consolidate the soil-water mixture. At saturation, the soil paste glistens as it reflects light, flows slightly when the container is tipped, and slides freely and cleanly off the spatula except for soils containing much clay. Allow the sample to stand overnight and then recheck the preceding criteria for saturation. Free water should not collect on the soil surface nor should the paste stiffen markedly or lose its glistening appearance on standing. If this occurs, remix with more water.

4. TEST PROCEDURE FOR pH

4.1 Per the manufacturer's instructions, standardize the pH meter using two of the standard buffer solutions: 7.0 and either 4.0 or 10.0, whichever is nearest to the estimated pH of the sample.

4.2 For each standard buffer solution, measure its temperature and adjust the temperature controller of the pH meter before testing for the standard solution pH value. This may not be needed on meters with automatic temperature compensation; follow manufacturer's instructions.

4.3 Stir the slurry mixture in the 100 mL beaker. Measure the temperature of the slurry and adjust the temperature controller of the pH meter before testing for the pH value of the slurry. This may not be needed on meters with automatic temperature compensation; follow manufacturer's instructions.

4.4 Carefully insert the pH probe in the slurry mixture. Do not place the electrode(s) into the soil; place them only into the slurry mixture. Determine the pH reading when the meter reading stabilizes.
4.5 If the pH reading is unstable when the electrode is immersed in the soil slurry, leave the electrode immersed until the pH reading has stabilized. In some cases, the waiting period for stabilization of the pH reading may take up to 5 minutes.

4.6 Record the pH value of the soil mixture, to the nearest tenth.

5. TEST PROCEDURE FOR SOLUBLE SALTS

5.1 Transfer the saturated soil paste to a filter funnel with a filter paper in place and apply vacuum until air begins to pass through the filter. Collect the extract in a bottle or test tube. Determine the temperature of the extract. Insert the conductivity cell into the extract, and determine electrical conductance. If the conductance falls above the highest range on the conductance meter, dilute the extract to one-tenth its original concentration of salts by diluting one part extract with 9 parts of distilled water. (The conductance reading obtained on the diluted extract must then be multiplied by ten since the conductance of the diluted extract is one-tenth the conductance of the original extract.)

6. CALCULATIONS FOR SOLUBLE SALTS

6.1 If conductance reading was taken in units other than microsiemens, convert it to conductance in microsiemens. For example, if reading was taken in millisiemens, multiply the reading by 1000 to convert to conductance in microsiemens.

6.2 Convert conductance to electrical conductivity, expressed as microsiemens per centimeter, by multiplying by the cell constant.

6.3 Correct the conductivity at the test temperature to conductivity at 25°C (77 °F), by the following formula:

\[ EC_{25} = (EC) \times (f_t) \]

Where:  
- \( EC_{25} \) = Electrical conductivity (microsiemens/cm), corrected for 25 °C.  
- \( EC \) = Electrical conductivity (microsiemens/cm), at test temperature.  
- \( f_t \) = Factor obtained from Figure 1.
6.4 Calculate the soluble salts content in parts per million (ppm) by the following formula:

\[
\text{Soluble Salts (ppm)} = (\text{EC}_{25}) \times (0.64)
\]

6.5 Report the soluble salts to the nearest 10 ppm.
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<th>°F</th>
<th>f_t</th>
<th>°C</th>
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**FIGURE 1**
PERCENT CARBONATES IN AGGREGATE

(An Arizona Method)

SCOPE

1. (a) This test method describes the procedure for determining the percentage of carbonates in aggregate. A combination of hydrogen peroxide and nitric acid is used to react with the carbonates.

   (b) This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS AND MATERIALS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus and materials for this test procedure shall consist of the following:

   (a) Drying apparatus - Any suitable device capable of drying samples at a temperature of 110 ± 5 °C (230 ± 9 °F).
(b) 1000 mL heavy duty beaker.

(c) Hydrogen Peroxide (3% solution) - $\text{H}_2\text{O}_2$.

(d) Nitric Acid (concentrated) - $\text{HNO}_3$.

(e) Distilled water.

(f) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

(g) Neutral Litmus Paper.

(h) Glass or Plastic Stirring Rod.

SAMPLE PREPARATION

3. Prepare the sample according to the following:

(a) For material samples from stockpile(s), obtain a representative 300 ± 10 gram sample of plus 4.75 mm (No. 4) material. Wash the sample over a 4.75 mm (No. 4) sieve and discard minus 4.75 mm (No. 4) material.

(b) For uncrushed material samples, obtain a representative sample and crush to appropriate grading. Obtain a representative 300 ± 10 gram sample of plus 4.75 mm (No. 4) material. Wash the sample over a 4.75 mm (No. 4) sieve and discard minus 4.75 mm (No. 4) material.

(c) The prepared sample shall be oven dried to constant weight at 110 ± 5 °C (230 ± 9 °F).

TEST PROCEDURE

4. (a) Allow sample to cool and place in a tared 1000 mL beaker. Weigh and record the weight of aggregate as the "weight of original sample" or "A" to the nearest 0.1 gram.
(b) Under a fume hood, add approximately 300 mL of H₂O₂ (3% solution) and stir. When the bubbling subsides, begin adding small amounts (approximately 10 mL) of concentrated HNO₃ to the beaker. Bubbling will be vigorous as the carbonates are being dissolved. Stir occasionally.

(c) When the bubbling has ceased and addition of HNO₃ causes no more bubbles, begin to wash by decantation, using distilled water. Care shall be taken not to lose any of the coarse aggregate. Wash by decantation at least 4 times. At this point, neutral litmus paper in the water should show only slight pink.

(d) Decant the water and oven dry to constant weight at 110 ± 5 °C (230 ± 9 °F).

(e) Let cool, weigh, and record the weight of aggregate as the "weight of non-reactive aggregate" or "B" to the nearest 0.1 gram.

CALCULATIONS

5. (a) Calculate the percent of carbonates as follows:

Percent of Carbonates = \( \frac{A - B}{A} \times 100 \)

Where: A = weight of original sample  
B = weight of non-reactive aggregate

(b) Report the percent of carbonates to the nearest 1%. 
Sieve Analysis and Separation of Salvaged AC Pavement Particles for Recycled Asphaltic Concrete

(An Arizona Method)

Scope

1. (a) This procedure provides a method for separation and sieve analysis of salvaged asphaltic concrete pavement particles which are obtained by crushing, milling, or other processes which renders particles which are to be used in the design and/or construction of recycled asphaltic concrete pavement.

(b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

Apparatus

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of necessary splitting equipment, sieves for the coarse screening of the materials, shaker, balance, and other miscellaneous items as specified in Arizona Test Method 201.
PROCEDURE

3. A representative sample of salvaged pavement particles is subjected to coarse sieving in accordance with Arizona Test Method 201, except the following:

   (a) If separated material is to be used in the design of a recycled mix, the entire nest of coarse [4.75 mm (No. 4) and larger] sieves shall be used. The material is subjected to shaking for 5 minutes ± 15 seconds.

   b) If separation of material on a particular sieve size is desired, for example, the 9.5 mm (3/8 inch) sieve, enough larger size sieves shall be used to prevent overloading of the desired sieve size. The material is subjected to shaking for 5 minutes ± 15 seconds, and the material retained on each sieve is combined.

   NOTE: The use of a specified time of shaking provides for a control on the "breaking down" of the particles of salvaged material into smaller size fractions. A sieving to completion is not attempted in this procedure.

SIEVE ANALYSIS

4. The sieve analysis on the separation of particles is performed as specified in Arizona Test Method 201.
COMPRESSIVE STRENGTH OF MOLDED CEMENT TREATED BASE OR SOIL-CEMENT SPECIMENS

(An Arizona Method)

1. SCOPE

1.1 This test method is used to determine the unconfined compressive strength of cement treated base or soil-cement specimens.

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

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 4 inch split ring proctor molds having a capacity of approximately 1/30 cubic foot, with an internal diameter of 4.000 plus 0.024 or minus 0.016 inches and a height of 4.584 plus 0.005 or minus 0.008 inches shall be used. The molds shall have a nominal wall thickness of 1/4 inch. They shall be equipped with an extension collar approximately 2-3/8 inches high.

2.3 A hand or mechanical rammer weighing 5.50 ± 0.02 pounds, having a flat face, and equipped with a suitable arrangement to control the height of drop to a free fall of 12 ± 0.06 (1/16) inches above the elevation of the soil. The rammer face shall be circular with a diameter of 2.000 plus 0.010 or minus 0.015 inches.

2.4 Hardened-steel straightedge, at least 10 inches in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.01 inch per 10 inches (0.1 percent) of length with the portion used for trimming the soil.
2.5 A circular sample follower with a diameter slightly less than the inner diameter of the proctor mold to aid in the removal of the mold collar.

2.6 Sieves of sizes as required for screening conforming to the requirements of ASTM E11.

2.7 Screw type or hydraulically operated compression testing machine capable of meeting the requirements of Subsection 6.3 below.

2.8 Miscellaneous mixing tools and pans.

3. **SAMPLE PREPARATION**

3.1 A representative sample of approximately 10 kg shall be obtained in a random manner from the completed mixture prior to final compaction, and immediately placed in an air-tight container. Not more than 90 minutes shall lapse between the time of mixing and the final mold fabrication.

3.2 Sieve the material over a 3/4 inch sieve and discard any Plus 3/4 Inch Material.

4. **FABRICATION**

4.1 The required number of specimens shall be promptly fabricated, following the procedure below.

4.2 Form a specimen by compacting the prepared soil in the four inch mold (with the extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. The top of the first and second layers shall be scratched to aid in bonding to subsequent layer. Compact each layer with 25 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation.

**NOTE:** Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: A block of concrete, weighing not less than 100 lbs, supported by a stable foundation; a sound concrete floor; and for field application, such surfaces as found in concrete box culverts, bridges, and pavements.
4.3 Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the specimen in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted specimen even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Level the specimen again with the straightedge, spraying with a fine mist of water as necessary to maintain specimen surface in a moist condition.

4.4 If it is necessary to transport the specimens, transport the specimens in the molds with the baseplate and collar attached, in a secure, moist environment as described in Subsection 5.2.

4.5 Following arrival at the location where curing will occur, remove specimens from molds within one hour, taking proper care in removing the specimens so that the shape of the molded specimens is maintained. To assure that specimens are in a moist condition they may be sprayed with a fine mist of water upon demolding, and promptly placed in the curing environment.

5. CURING

5.1 The specimens to be tested for acceptance shall be cured in a fog room at 100% humidity between 65°F and 80°F for six days. The specimens shall be protected from direct contact with free water to prevent saturation and/or leaching of cementitious material from the specimen.

5.2 If the specimens are cured where a “fog” room is not available, the curing is to be accomplished by surrounding the specimens with pre-saturated perlite or asbestos free vermiculite in a sealed container. Pre-saturation is determined as the maximum amount of water which can be absorbed without the presence of free standing water. This sealed container shall be promptly placed in a 65°F and 80°F environment. Store the specimens on a rigid, non-absorbent surface.

5.3 After six days the specimens to be tested for acceptance shall be immersed in saturated lime water at 65 to 80 °F for 24 hours.

NOTE: Informational specimens utilized for 1-day compressive strength testing are not immersed in saturated lime water.
6. **COMPRESSIVE STRENGTH**

6.1 Carefully remove the specimens from the water and towel dry. Due to the fragility of the specimens they must be handled carefully to assure that they are not damaged. Determine the diameter of each test specimen to the nearest 0.01 inch by averaging two diameter determinations measured at right angles to each other at mid-height of the specimen. The specimens shall be capped in accordance with ASTM C617/AASHTO T 231, and broken as soon as practical while specimens are in their moist condition after capping.

6.2 Place each specimen on the bearing block of the compression machine and center it under the loading head.

6.3 Continuously apply a uniform load to the specimen at the rate of 0.05 ± 0.01 inches per minute for the screw type testing machines, or 20 to 50 psi/sec for the hydraulically operated machines until failure is noted. Record the load at failure and convert to psi using the diameter determined in Subsection 6.1 above. No further correction shall be applied to the results. The average of three results shall constitute one test value for purposes of compressive strength specification compliance.

7. **REPORT**

7.1 Record the load failure point and compressive strength in psi for each specimen, and the average compressive strength for the three specimens.
SAND EQUIVALENT TEST FOR MINERAL AGGREGATE
FOR ASPHALTIC CONCRETE FRICTION COURSE

(A Modification of AASHTO T 176)

This test procedure is the same as specified in AASHTO T 176, except:

1. The following is added: “This procedure is used for determining the sand equivalent value for mineral aggregate to be used in ACFC. The mineral aggregate gradation requirements for ACFC yield a large percentage of material which passes the 4.75 mm (No. 4) sieve and is retained on the 2.36 mm (No. 8) sieve. Due to the amount of material retained on the 2.36 mm (No. 8), the sand equivalent test, when performed as outlined in AASHTO T 176, utilizing minus 4.75 mm (No. 4) material does not lend itself to testing the mineral aggregate required for ACFC.”

2. The following is added: “This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.”

3. The following is added: “See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.”

4. The following is added: “Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

5. The following is added: “Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.”

6. In Section 4, “Sample Preparation”, all references to "4.75 mm (No. 4) sieve" shall be replaced by "2.36 mm (No. 8) sieve."
ARTIFICIAL GRADING OF MINERAL AGGREGATE

(An Arizona Method)

Scope

1. This method may be utilized to adjust composites of mineral aggregate by artificially grading the material to meet desired target values for specified screens.

Preparation of Pass No. 4 Sieve Size Samples

2. After samples of each stockpile or bin have been separated into individual sizes of Plus No. 4 and larger, and Pass No. 4 material by coarse sieving as described in ARIZ 201 the Pass No. 4 material is processed by the following procedures. Figure 1 gives an illustration of the steps necessary which may prove helpful in understanding the procedure.

(a) For each stockpile or bin of mineral aggregate the Pass No. 4 material is separated into No. 8 and Pass No. 8 fractions and weights recorded for each.

(b) The Pass No. 8 fractions for each stockpile or bin are proportionately combined and thoroughly blended together.

NOTE: It is important that the material be completely and accurately sieved into the three separate size fractions.

(c) The combined Pass No. 8 material is screened over a series of sieves which includes a No. 8, No. 40, No. 200 sieve, and other intermediate sizes as necessary to prevent overloading individual sieves.

(d) Material which passes the No. 8 sieve and is retained on the No. 40 sieve and any intermediate sieves is combined and saved.

(e) Material which passes the No. 40 sieve and is retained on the No. 200 sieve and any intermediate sieves is combined and saved.

(f) The material passing the No. 200 sieve is also saved.

(g) An approximate 500 gram sample is obtained from the fraction of material which is Pass No. 8 to Retained No. 40, and another approximate 500 gram sample is obtained from the fraction of material which is Pass No. 40 to Retained No. 200.
(h) Each of the samples are subjected to elutriation and fine screening as specified in ARIZ 201, Section 6 or 7, and 8. The sieve analysis of each is determined in accordance with Section 11 of ARIZ 201, except the % passing each sieve is determined and recorded to the nearest 0.01%.

(i) From the elutriation and fine screening of the Pass No. 8 to Retained No. 40 fraction, the % passing the No. 40 sieve and larger than the No. 200 sieve is determined (% Pass No. 200 subtracted from % Pass No. 40), and recorded as "a". The % passing the No. 200 sieve is recorded as "b".

Example:

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<th>% Passing</th>
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<td>0.34</td>
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<tr>
<td># 200</td>
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</table>

"a" = (0.34 - 0.25) = 0.09%

"b" = 0.25%

(3) From the elutriation and fine screening of the Pass No. 40 to Retained No. 200 fraction, the % passing No. 200 sieve is recorded as "c".

Example:

<table>
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<tr>
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</thead>
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<tr>
<td># 200</td>
<td>1.32</td>
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"c" = 1.32%

NOTE: The size fractions used above for artificial grading and throughout this method are based on meeting desired values for % passing the No. 8, No. 40 and No. 200 sieves. If other or additional sieves are used in artificial grading or adjusting the composite, the procedures of this test method will apply, with appropriate modifications.

Procedure for Adjusting Composite for Pass No. 8 Material

3. The composite of mineral aggregate for Pass No. 8 material is then adjusted to meet desired values. The Pass No. 8 to Retained No. 40 fraction from screening has material clinging to it which will contribute to Pass No. 40 to Retained No. 200 and Pass No. 200 material in the adjusted composite, and the Pass No. 40 to Retained No. 200 fraction from screening has material clinging to it which will contribute to the Pass No. 200 material in the adjusted composite. The procedure below uses results from the elutriation and the fine screening of material from each fraction of material to account for the total contribution to each desired size. Figure 2 provides an illustration which may be useful in understanding the
method. The calculations outlined below are utilized to determine the % of each size fraction from screening that is necessary to be used to obtain desired values. An example is provided with the method for each calculation. The illustration in paragraph (h) is a tabulation of the results for the example provided and should be referred to as the procedure is followed to understand the contributions to each desired value from the individual fractions of material.

(a) The amount of Pass No. 8 to Retained No. 40 material for the desired gradation is determined, (% Pass No. 40 subtracted from % Pass No. 8) and recorded as "d". The amount of Pass No. 40 to Retained No. 200 material for the desired gradation is determined, (% Pass No. 200 subtracted from % Pass No. 40), and recorded as "e". The amount of Pass No. 200 for the desired gradation is recorded as "f".

Example:

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Composite Gradation % Pass Values</th>
<th>Desired Gradation % Pass Values</th>
</tr>
</thead>
<tbody>
<tr>
<td># 8</td>
<td>45</td>
<td>48</td>
</tr>
<tr>
<td># 40</td>
<td>21</td>
<td>18</td>
</tr>
<tr>
<td># 200</td>
<td>6.5</td>
<td>5.7</td>
</tr>
</tbody>
</table>

"d" = (48 - 18) = 30 %
"e" = (18 - 5.7) = 12.3%
"f" = 5.7%

(b) The percent of the Pass No. 8 to Retained No. 40 fraction of material necessary to be utilized to obtain the desired adjusted % Pass No. 8 is determined and recorded as "g".

\[
g = (d) \times \frac{a + b}{1 + \frac{a + b}{100}}
\]

Example:

\[
g = (30) \times \frac{0.09 + 0.25}{1 + \frac{0.09 + 0.25}{100}} = 30.10\%
\]

(c) The proportional amounts of Pass No. 40 to Retained No. 200, and Pass No. 200 material which are in the Pass No. 8 to Retained No. 40 fraction, and which need to be accounted for in the adjusted composite are determined and recorded as "h" and "i" respectively:

\[
h = (g - d) \times \frac{a}{a + b}
\]

\[
i = (g - d) \times \frac{b}{a + b}
\]
Example:

\[ h = (30.10 - 30) \times \frac{0.09}{0.09 + 0.25} = 0.03\% \]

\[ i = (30.10 - 30) \times \frac{0.25}{0.09 + 0.25} = 0.07\% \]

(d) The amount of Pass No. 40 to Retained No. 200 material from the Pass No. 40 to Retained No. 200 fraction that is required to give the desired value for Pass No. 40 to Retained No. 200 in the adjusted composite is determined and recorded as "j":

\[ j = (e - h) \]

Example:

\[ j = (12.3 - 0.03) = 12.27\% \]

NOTE: The sum of the contributions to the Pass No. 8 to Retained No. 40, "h" + "j", will equal the total amount desired in the adjusted composite, "e".

(e) The percent of the Pass No. 40 to Retained No. 200 fraction of material, necessary to be utilized to obtain the desired value for Pass No. 40 to Retained No. 200 is determined and recorded as "k":

\[ k = (j) \times \left[ 1 + \frac{c}{100} \right] \]

Example:

\[ k = (12.27) \times \left[ 1 + \frac{1.32}{100} \right] = 12.43\% \]

(f) The amount of Pass No. 200 material which is in the Pass No. 40 to Retained No. 200 fraction, and which needs to be accounted for in the adjusted composite is determined and recorded as "l":

\[ l = (k - j) \]

Example:

\[ l = (12.43 - 12.27) = 0.16\% \]
(g) The percent of Pass No. 200 material necessary to be utilized to obtain the desired \% Pass No. 200 is determined and recorded as "m":

\[ "m" = (f - i - l) \]

Example:

\[ "m" = (5.7 - 0.07 - 0.16) = 5.47\% \]

NOTE: The sum of the contributions to the \% Pass No. 200 material, "i" + "l" + "m", will equal the total amount desired in the adjusted composite, "f".

(h) The following illustration shows the contributions to the adjusted composite from each of the size fractions:

<table>
<thead>
<tr>
<th>% Pass #8 to Ret. #40 Fraction</th>
<th>Pass #40 to Ret. #200 Fraction</th>
<th>Pass #200 Fraction</th>
<th>Desired Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 (d)</td>
<td>12.27 (j)</td>
<td>5.47 (m)</td>
<td>30 (d)</td>
</tr>
<tr>
<td>0.03(h)</td>
<td>0.16(l)</td>
<td>5.47 (m)</td>
<td>12.3 (e)</td>
</tr>
<tr>
<td>0.07(i)</td>
<td>5.47(m)</td>
<td></td>
<td>5.7 (f)</td>
</tr>
<tr>
<td>30.10(g)</td>
<td>12.43(k)</td>
<td>5.47 (m)</td>
<td></td>
</tr>
</tbody>
</table>

(i) The percentages of each size fraction which are to be used in the adjusted composite to meet desired values are recorded:

Example:

Pass No. 8 to Retained No. 40 = 30.10\%
Pass No. 40 to Retained No. 200 = 12.43\%
Pass No. 200 = 5.47\%

Procedure for Adjusting Composite for Plus No. 8 Material

4. The composite is adjusted for the Plus No. 8 material to obtain desired \% pass values for each size.

(a) The \% retained for each sieve size for the desired adjusted composite is divided by the corresponding \% retained value for each sieve size in the stockpile or bin composite. This value is multiplied by the individual fractions of that size for each stockpile or bin. This will give an adjusted amount of each size fraction in the stockpile or bin composite which will be the gradation of the desired composite. An example of this for a composite for an asphaltic concrete mix design is shown in Figure 3.
Checking Adjusted Composite

5. A sample of appropriate size as specified in ARIZ 201, for the maximum aggregate size, is prepared for the adjusted composite using material from the individual size fractions for Plus No. 8 material, the Pass No. 8 to Retained No. 40 fraction, the Pass No. 40 to Retained No. 200 fraction, and the Pass No. 200 fraction. This sample is subjected to ARIZ 201 and the resultant gradation compared to the desired gradation.

Preparing Samples Using Material from Artificial Grading

6. The samples necessary are prepared and weighed up for testing utilizing the adjusted composite.

   (a) For the Plus No. 8 material, the individual size fractions from the adjusted composite for each stockpile or bin are utilized to prepare samples.

   (b) For Pass No. 8 material the amount of each fraction for Pass No. 8 to Retained No. 40, Pass No. 40 to Retained No. 200, and Pass No. 200 from the adjusted composite is utilized to prepare samples.

   (c) A weigh up sheet which is used for asphaltic concrete mix designs, which illustrates the use of the adjusted composite fractions is shown in Figure 4.
FIGURE 1

Amount of Pass No. 8 to Retained No. 40 fraction to be used to obtain desired amount of Pass No. 8 = "g"

Pass No. 8 to Ret. No. 40 = "d"

Amount of Pass No. 8 to Retained No. 40 in the desired gradation = "d"

Amount of Pass No. 40 to Retained No. 200 fraction to be used to obtain desired amount of Pass No. 40 = "k"

Pass No. 40 to Ret. No. 200 = "h"

Amount of Pass No. 40 to Retained No. 200 in the desired gradation = "e"

Amount of Pass No. 200 to Retained No. 200 fraction to be used to obtain desired amount of Pass No. 200 = "m"

Pass No. 200 = "i"

Amount of Pass No. 200 in the desired gradation = "f"
Stockpile or Bin Pass No. 4 from screening

- No. 8
- Pass No. 8

Proportionately combine the Pass No. 8 from each stockpile or bin, blend thoroughly and screen.

- Pass No. 8 to Retained No. 40 fraction
  - 500 gram sample for elutriation and fine screening.
    - Record % Pass No. 40 to Ret. No. 200 as "a"

- Pass No. 40 to Retained No. 200 fraction
  - 500 gram sample for elutriation and fine screening.
    - Record % Pass No. 200 as "b"

- Pass No. 200 fraction

FIGURE 2
## Composite Adjusted to Desired Composite of Plus No. 8 Material

### Aggregate % of Composite

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>C 30%</th>
<th>I 25%</th>
<th>F 45%</th>
<th>% Ret.</th>
<th>% Ret. Rounded</th>
<th>Pass</th>
</tr>
</thead>
<tbody>
<tr>
<td>1&quot;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>2.6</td>
<td>0.4</td>
<td>0.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>3.0</td>
<td>3.0</td>
<td>3.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>2.5</td>
<td>2.5</td>
<td>2.5</td>
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<td></td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>2.2</td>
<td>2.2</td>
<td>2.2</td>
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<td></td>
</tr>
<tr>
<td>#4</td>
<td>1.6</td>
<td>1.6</td>
<td>1.6</td>
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<tr>
<td>#8</td>
<td>5.0</td>
<td>5.0</td>
<td>5.0</td>
<td></td>
<td></td>
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<tr>
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<td>4.8</td>
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<td>#30</td>
<td>1.4</td>
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<td>0.4</td>
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<tr>
<td>Pass #200</td>
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<td>1.53</td>
<td>4.29</td>
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</table>

### Design Criteria

- **MIX DESIGN REQUIREMENTS:**
  - **IMC Ret.**
  - **Min. Wet Strength**
  - **Voids**
  - **V.M.A.**
  - **Min. Stability**
  - **Flow**

### Composite of Pass No. 8 from Generation of Each Stockpile or Bin:

- **Coarse 0.2**
- **Inter 7.9**
- **Fine 37.2**

### Total = 45.3

### Adjusted % of Composite for Use of Mineral Admixture

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Adjusted % of Composite</th>
<th>Adjusted % of Composite</th>
<th>Adjusted % of Composite</th>
<th>Adjusted % of Mineral Admixture</th>
<th>Calc. Pass</th>
<th>% Ret.</th>
<th>Pass</th>
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<td></td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>&quot;</td>
<td>&quot;</td>
<td>&quot;</td>
<td>&quot;</td>
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<td>&quot;</td>
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</tr>
</tbody>
</table>

### Remarks

- \( \frac{3}{3} = 0.8333 \)
- \( \frac{15}{15.2} = 0.9848 \)
- \( \frac{11}{9.7} = 1.1340 \)
- \( \frac{4}{5.7} = 0.7018 \)
- \( \frac{6}{5.3} = 1.1321 \)
- \( \frac{13}{15.2} = 0.8553 \)

**FIGURE 3**
### ASPHALTIC CONCRETE MIX DESIGN WEIGH UP CARD

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>LAB NO.</th>
<th>DATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>PROJECT NO.</td>
<td>PROJECT NAME</td>
<td></td>
</tr>
<tr>
<td>CONTRACTOR</td>
<td>MIX DESIGN REQUEST</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LAB NO.</th>
<th>SIZE</th>
<th>ACCUM % RET.</th>
<th>RICE</th>
<th>STABS</th>
<th>IMC</th>
<th>COARSE SP. GR</th>
<th>COARSE SAND EQUIV.</th>
<th>CRUSH FACES</th>
<th>COARSE C.R.E</th>
<th>LIME STONE</th>
<th>FLAKINESS INDEX</th>
<th>BLENDED</th>
<th>FINE SAND EQUIV.</th>
<th>FINE SP. GR</th>
<th>MAKE-UP MAT'L</th>
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<tbody>
<tr>
<td>Coarse</td>
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<td>2.6</td>
<td>78</td>
<td>86</td>
<td>88</td>
<td>267</td>
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<td>26</td>
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<td>26</td>
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<td>26</td>
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<tr>
<td>1/2</td>
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<td>1261</td>
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</table>

<table>
<thead>
<tr>
<th>Size</th>
<th>LAB NO.</th>
<th>PASS 8 - RET 40</th>
<th>PASS 40 - RET 200</th>
<th>PASS 200</th>
</tr>
</thead>
<tbody>
<tr>
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<td>1243</td>
<td>1373</td>
<td>1414</td>
</tr>
<tr>
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<tr>
<td>1/4</td>
<td>1560</td>
<td>1716</td>
<td>1768</td>
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<tr>
<td>#8</td>
<td>1592</td>
<td>1792</td>
<td>1836</td>
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**Remarks:**

*FIGURE 4*
MAXIMUM DRY DENSITY AND OPTIMUM MOISTURE OF SOILS
BY PROCTOR ALTERNATE METHOD D

(A Modification of AASHTO Designation T 99)

1. SCOPE

1.1 This test method describes the procedure for determining the maximum dry density and optimum moisture content for a soil by the Proctor Alternate Method D.

1.2 Alternate Method D may be used for all maximum dry density and optimum moisture content determinations except for volcanic cinders or light porous material on which the specific gravity cannot be determined with consistency or when the moisture absorption for the coarse aggregate is greater than 4.0%.

1.3 Alternate Method D may be used except when greater than 40% of the material is retained on the 3/4 inch sieve.

1.4 An example is provided in Figure 2 for the calculations and determinations referenced herein.

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

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 6 inch proctor mold having a capacity of approximately 1/13.33 cubic foot, with an internal diameter of 6.000 plus 0.039 or minus 0.026 inches and a height of
4.584 plus 0.005 or minus 0.008 inches. The mold shall have a nominal wall thickness of 1/4 inch. It shall be equipped with an extension collar approximately 2-3/8 inches high. A baseplate as shown in Figure 1 shall be provided.

2.3 A hand or mechanical rammer weighing 5.50 ± 0.02 pounds, having a flat face, and equipped with a suitable arrangement to control the height of drop to a free fall of 12 ± 0.06 (1/16) inches above the elevation of the soil. The hand rammer face shall be circular with a diameter of 2.000 plus 0.010 or minus 0.015 inches. The mechanical rammer face shall have the shape of a piece of pie, with a radius of approximately 3 inches, and having an area of 3.134 ± 0.039 square inches. If a mechanical apparatus is used, it must be monitored through the ADOT proficiency sample program and maintain a rating of 3 or better based on the results of testing ADOT and AMRL proficiency samples.

2.4 Hard steel straightedge, at least 10 inches in length. It shall have one beveled edge, and at least one longitudinal surface (used for final trimming) shall be plane within 0.01 inch per 10 inches (0.1 percent) of length with the portion used for trimming the soil.

2.5 Scale or balance capable of measuring the maximum weight to be determined, accurate to at least one gram.

2.6 Scale or balance capable of measuring the maximum weight to be determined, accurate to at least 0.1 gram.

2.7 Oven capable of maintaining a temperature of 230 ± 9 °F.

2.8 3/4 inch and 3 inch sieves conforming to the requirements of ASTM E11.

2.9 Miscellaneous mixing tools and pans.

2.10 Sample Extruder (optional) consisting of a jack, lever, frame, or other device for extruding the compacted sample from the mold.

3. CALIBRATION OF MOLD

3.1 Molds shall be calibrated in accordance with APPENDIX A of Arizona Test Method 225.
4. SAMPLE

4.1 Enough soil material shall be provided from the field to make five compacted specimens. A minimum sample size of 45,000 grams (approximately 100 lbs.) is normally required.

4.2 If the soil sample is damp when received from the field, dry it until it becomes friable under a trowel. Drying may be in air or by use of a slow fan or other drying apparatus such that the temperature of the sample does not exceed 140 °F.

4.3 Thoroughly break up the aggregations in such a manner as to avoid reducing the natural size of individual particles.

4.4 Weigh out an approximate 45,000 gram sample of representative soil. Record the weight of the sample, and sieve the material over a 3/4 inch sieve. If the percentage of coarse aggregate or rock retained on the 3/4 inch sieve is not already known from gradation testing, save any material retained on the 3/4 inch sieve and weigh. Calculate the percent of coarse aggregate or rock particles retained on the 3/4 inch sieve according to the following equation:

\[
PR_{3/4} = \frac{WR_{3/4}}{WT} \times 100
\]

Where:

- \( PR_{3/4} \) = Percentage of coarse aggregate or rock particles retained on the 3/4 inch sieve
- \( WR_{3/4} \) = Weight of coarse aggregate or rock particles retained on the 3/4 inch sieve
- \( WT \) = Total weight of material sieved

4.5 If "PR\(_{3/4}\)" is greater than 40%, then too much rock is present to allow for a reasonable maximum dry density determination. If "PR\(_{3/4}\)" is less than or equal to 40%, blend material passing the 3/4 inch sieve thoroughly and proceed to Section 5 of this test method. If a specific gravity and absorption determination, in accordance with AASHTO T 85, is to be made for the plus 3/4 inch material, save an adequate amount of this material, otherwise, discard it.
5. PROCEDURE

5.1 From the thoroughly blended passing 3/4 inch material from Subsection 4.5, split out 5 representative approximate 5000 gram samples.

5.2 Select one sample and thoroughly mix with sufficient water to dampen it to approximately three percentage points below optimum moisture content.

Note: If desired, an additional three samples may be mixed at this time with approximate moisture contents of 1% below optimum, 1% over optimum, and 3% over optimum. The moisture in each of these samples shall be retained by covering with a damp cloth or being sealed in air tight containers until they are compacted. One of the five samples should be retained for future use since it is necessary to have at least two points defined on each side of the moisture-density curve.

5.3 Heavy clay soils or materials which tend to break down, or those in which it is difficult to incorporate water, shall require approximately 12 hours for uniform moisture absorption to be achieved. This shall be accomplished by preparing separate samples for each increment of water to be added, and then placing and sealing these samples in air tight containers for the 12-hour period.

5.4 Form a specimen by compacting the prepared soil in the six inch mold (with extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. Compact each layer with 56 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid and stable foundation.

Note: Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: A block of concrete, weighing not less than 200 lbs., supported by a stable foundation; a sound concrete floor; and for field application, such surfaces as found in concrete box culverts, bridges, and pavements.
5.5 When compacting granular, free-draining materials, at moisture contents which are at or above optimum, the mold shall be prepared by first sealing the bottom of the mold with waterproofing grease. All excess grease shall be wiped from the mold and baseplate.

5.6 Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the soil in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted soil even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Determine the weight of compacted specimen and mold. Determine the wet density, "WD", of the compacted soil by the following:

\[
WD = \frac{M1 - M2}{VM \times 453.6 \text{ (grams/lb.)}}
\]

Where:
- WD = Wet density of compacted soil, lb./cu. ft.
- M1 = Weight of compacted specimen and mold, grams
- M2 = Weight of the mold, grams
- VM = Volume of the mold, cu. ft. (See Section 3 of this procedure.)

5.7 The estimated dry density, "ED", of the compacted soil may be calculated and recorded to the nearest 0.1 lb./cu. ft. as follows:

\[
ED = \frac{WD}{(\text{Approx. } \% \text{ of water added}) + 100} \times 100
\]

Where:
- ED = Estimated dry density of compacted soil, lb./cu. ft.
- WD = Wet density of compacted soil, lb./cu. ft.

Note: These estimated densities are approximate only and will be corrected when final moisture results are obtained.

5.8 The estimated dry density is useful in deciding how much water to add in later trial batches, if the procedure described in the note following Subsection 5.2, for
initially preparing three additional samples with varying moisture contents is not utilized. By carefully observing the estimated dry density of the compacted samples, the operator should be able to vary the moisture content as the test proceeds so that when the final moisture-density results are plotted, a curve will result that rises to a peak and then falls away.

5.9 Remove the material from the mold and slice vertically through the center. Take a representative minimum 600 gram sample from the full length and width of one of the cut faces. Weigh immediately, and dry to a constant weight in an oven at 230 ± 9 °F to determine the moisture content in accordance with AASHTO T 265. Record the weight of wet soil to the nearest 0.1 gram as "WW", and record the weight of oven dry soil to the nearest 0.1 gram as "DW".

5.10 For granular, free-draining materials, the moisture content shall be determined using the entire compacted proctor specimen. Determine the weight of wet soil, "WW", by subtracting the weight of the mold, "M2", from the weight of compacted specimen and mold, "M1". Record the weight of wet soil, "WW", and the weight of oven dry soil, "DW", to at least the nearest gram. All clinging material shall be removed from the inside of the mold and included with the specimen. To facilitate drying, the specimen may be broken up and spread out in a large pan, being careful to not lose any soil particles.

5.11 Select another of the samples which was split in Subsection 5.1, and if not already done, thoroughly mix with water in sufficient amount to increase the moisture content by approximately two percentage points.

5.12 Repeat the procedure in Subsections 5.3 through 5.10 for the sample at each moisture content, as necessary to establish a moisture-density curve which rises to a peak and then falls away.
6. **CALCULATIONS**

6.1 Calculate percent moisture and record to the nearest 0.1 percent by the following:

\[
\text{% Moisture} = \frac{\text{WW} - \text{DW}}{\text{DW}} \times 100
\]

Where:  
- WW = weight of wet soil, grams  
- DW = weight of oven dry soil, grams

6.2 Calculate dry density and record to the nearest 0.1 lb./cu. ft. by the following:

\[
\text{DD} = \frac{\text{WD}}{\text{% Moisture} + 100} \times 100
\]

Where:  
- DD = Dry density of compacted soil, lb./cu. ft.  
- WD = Wet density of compacted soil, lb./cu. ft.

7. **MOISTURE-DENSITY RELATIONSHIP**

7.1 The percent moisture and corresponding dry density for each of the compacted soil specimens shall be plotted on the graph provided on the proctor density test form shown in Figure 3. For a good plot, the majority of the graph is utilized. Normally, three increments on the horizontal axis shall equal one percent of moisture, and three increments on the vertical axis shall equal one lb./cu. ft. of dry density. If another number of increments other than three is utilized, the number of increments for one percent moisture and one lb./cu. ft. dry density shall always be the same.

7.2 On each side of the maximum density curve, at least two points should be utilized to form two straight lines. The intersection point of these two lines defines the peak point of the density-moisture content relationship, or the maximum density and optimum moisture content for the soil. In general it will be found that higher unit mass soils assume steeper slopes with high maximum dry densities at low optimum moisture contents, while the lower unit mass soils
assume flatter, more gently sloped lines with high optimum moisture contents and low maximum dry densities. Figure 4 gives examples of moisture-density plots which show the different slopes associated with different maximum dry density ranges.

7.3 Optimum moisture content - The percent moisture content corresponding to the peak (intersection point of the two lines) of the moisture-density curve shall be termed the "optimum moisture content", and shall be reported as "OM" to the nearest 0.1 percent.

7.4 Maximum dry density - The dry density at optimum moisture content corresponding to the peak (intersection point of the two lines) of the moisture-density curve shall be termed the "maximum dry density", and shall be reported as "MD" to the nearest 0.1 lb./cu. ft.

Note: The optimum moisture and maximum dry density determinations above are for the material passing the 3/4 inch sieve. When testing field samples for comparison to proctor optimum moisture and maximum dry density, a correction to the proctor optimum moisture and maximum dry density must be made, in accordance with ARIZ 227, for the percent rock which the field sample contains.

8. REPORT

8.1 Record the moisture and density data on the laboratory test form along with the laboratory number, material source and type, and other information required. A blank laboratory test form is provided in Figure 3.
CYLINDERICAL MOLD, COLLAR, AND BASEPLATE
(6 INCH MOLD)

FIGURE 1
Arizona Department of Transportation

METHOD A or ALTERNATE METHOD D PROCTOR DENSITY
(Arizona Test Method 225 or 245)

Weight of Mold = M2 = 2840 grams
Volume of Mold = VM = 0.0744 cu ft

<table>
<thead>
<tr>
<th>b</th>
<th>M1</th>
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<th>WD</th>
<th>ED</th>
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<th>DD</th>
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<td></td>
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<td></td>
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<td>4617</td>
<td>136.8</td>
<td>121.1</td>
<td>645.9</td>
<td>572.1</td>
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**METHOD A**

WT = WR4 = 
PR4 = WR4 x 100 = 

**ALTERNATE METHOD D**

WT = 48780 WR 3/4 = 17951
PR 3/4 = WR 3/4 x 100 = 37 %

Coarse Aggregate
Bulk Oven Dry
Specific Gravity : 2.631

Coarse Aggregate
Absorption : 2.28 %

OPTIMUM MOISTURE CONTENT = OM = 10.0 %
MAXIMUM DRY DENSITY (lb./cu. ft.) = MD = 124.6

Remarks:

FIGURE 2
Arizona Department of Transportation

METHOD A or ALTERNATE METHOD D PROCTOR DENSITY
(Arizona Test Method 225 or 245)

<table>
<thead>
<tr>
<th>Approx % of water Added</th>
<th>Wt. of Sample and Mold</th>
<th>Wet Wt. of Sample</th>
<th>Wet Density lb/cu ft</th>
<th>Est. Dry Density</th>
<th>Wet Wt. of Moisture Sample</th>
<th>Dry Wt. of Moisture Sample</th>
<th>Wt. of Water</th>
<th>Percent Moisture</th>
<th>Dry Density lb/cu ft</th>
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<tbody>
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</tr>
</tbody>
</table>

**METHOD A**

\[ WT = \frac{WR4}{WT} \]
\[ PR4 = \frac{WR4 \times 100}{WT} \]

**ALTERNATE METHOD D**

\[ WT = \frac{WR 3/4}{WT} \]
\[ PR 3/4 = \frac{WR 3/4 \times 100}{WT} \% \]

**Coarse Aggregate**
- Bulk Oven Dry
- Specific Gravity:

**Dry Density lb/cu ft**

**Optimum Moisture Content**
\[ OM = \%

**Maximum Dry Density**
\[ MD = \%

**Remarks:**

FIGURE 3
AGGREGATE BASE COURSE

% Moisture  Dry Density
5.1         120.8
7.1         122.1
8.9         123.7
10.8        122.7
12.3        121.3

Moisture Density = 124.1 lb/cu ft
Optimum Moisture = 9.4 %

SILTY SAND
AND GRAVEL

% Moisture  Dry Density
7.2         127.0
8.1         129.6
9.4         127.9
10.1        126.6

Moisture Density = 130.0 lb/cu ft
Optimum Moisture = 8.3 %

FIGURE 4
ARIZONA DEPARTMENT OF TRANSPORTATION

CALIBRATION OF PROCTOR MOLD
ARIZ 225 Appendix A

Four Inch Mold  Six Inch Mold  Mold I. D. #:  6A

Calibration Date: 09/15/15  Calibration Expiration Date: 09/15/16

Temperature of water used for Calibration: 75 ° F

Unit Weight of Water: 62.261 lb./cu. ft.

Test Operator: Joe Tester  Supervisor and Date: Joe Supervisor 09/17/15

<table>
<thead>
<tr>
<th>Weight of Baseplate, Empty Mold, and Glass Plate (grams)</th>
<th>Weight of Baseplate, Mold Filled with Water, and Glass Plate (grams)</th>
<th>Weight of Water to Fill Mold (grams)</th>
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</thead>
<tbody>
<tr>
<td>6185.8</td>
<td>8287.0</td>
<td>2101.2</td>
</tr>
</tbody>
</table>

\[
V = \left( \frac{\text{Weight of Water to Fill Mold (grams)}}{\text{Unit Weight of Water (lb. / cu. ft.)}} \right) \times \left( \frac{\text{Volume of Mold (cu. ft.)}}{453.6 \text{ (grams / lb.)}} \right)
\]

\[
V = \left( \frac{2101.2}{62.261} \right) \times \left( \frac{1}{453.6} \right) = 0.0744 \text{ cu. ft.}
\]

REMARKS:

<table>
<thead>
<tr>
<th>Temp °F</th>
<th>Unit Weight of Water Table</th>
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<tr>
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<td>62.155</td>
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</table>

FIGURE 5
ARIZONIA DEPARTMENT OF TRANSPORTATION

CALIBRATION OF PROCTOR MOLD
ARIZ 225 Appendix A

______ Four Inch Mold  ______ Six Inch Mold  Mold I. D. #: __________

Calibration Date: ______________  Calibration Expiration Date: ______________

Temperature of water used for Calibration: __________ °F

Unit Weight of Water: ______________ lb. / cu. ft.

Test Operator: _____________________  Supervisor and Date: ______________

<table>
<thead>
<tr>
<th>Weight of Baseplate, Empty Mold, and Glass Plate (grams)</th>
<th>Weight of Baseplate, Mold Filled with Water, and Glass Plate (grams)</th>
<th>Weight of Water to Fill Mold (grams)</th>
</tr>
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</table>

\[ V = \frac{\text{Unit Weight of Water}}{453.6 \text{ (grams/lb.)}} \times \text{cu. ft.} \]

\[ V = \left( \frac{\text{Unit Weight of Water}}{\text{lb. / cu. ft.}} \right) \times \left( \text{cu. ft.} \right) \]

REMARKS:


<table>
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<tr>
<th>Unit Weight of Water Table</th>
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FIGURE 6
MOISTURE - DENSITY RELATIONSHIP USING TYPICAL MOISTURE - DENSITY CURVES (ONE POINT PROCTOR) ALTERNATE METHOD D

(An Arizona Method)

1. SCOPE

1.1 This method of test is for the determination of the optimum moisture content and maximum dry density of a soil or soil-aggregate mixture utilizing one moisture-density determination on the portion of the sample passing the 3/4 inch sieve.

1.2 The one-point proctor is used with the typical moisture-density curves, shown on the back of the One Point Proctor Density Test Card (Figures 1 and 2); or by utilizing a family of moisture-density curves developed for the immediate local conditions.

1.3 This method is not to be used for volcanic cinders or light porous material on which the specific gravity cannot be determined with consistency or when the absorption of the coarse aggregate is greater than 4.0%.

1.4 This method may be used to determine if an existing proctor maximum density determination is valid for the soil being tested. If the existing proctor maximum density determination is not valid, a full proctor according to Arizona Test Method 245 should normally be run to determine the maximum density required for that soil type.

1.5 An example is provided in Section 7, and Figures 3 and 4, for the calculations and determinations referenced herein.

1.6 This test method may involve hazardous materials, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of who ever uses this test method to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.
2. **APPARATUS**

2.1 The apparatus shall consist of the following:

2.1.1 The apparatus utilized for this test method shall conform to the apparatus requirements of Arizona Test Method 245.

2.1.1.1 Instead of the 230 ± 9 °F oven, a hot plate or stove capable of maintaining a temperature of approximately 230 °F may be used. A Speedy Moisture Tester with a conversion table or calibration curve may also be used for moisture determinations made in the field. Finally, a microwave oven may be used in accordance with Arizona Test Method 719.

2.1.1.2 Instead of the scale or balance capable of measuring the weight to be determined to at least one gram, a scale capable of measuring the weight to at least 0.01 pound may be utilized.

3. **CALIBRATION OF MOLD**

3.1 Molds shall be calibrated in accordance with APPENDIX A of Arizona Test Method 225.

4. **SAMPLE**

4.1 A representative sample of passing 3/4 inch material weighing approximately 5000 grams shall be obtained for each one-point proctor.

5. **PROCEDURE**

5.1 If the Speedy Moisture Tester is not to be used in making the moisture content determination, proceed to Subsection 5.4.

5.2 For testing performed in the field, the Speedy Moisture Tester (AASHTO T 217) may be used to make the moisture content determination. The approximate 5000 gram sample of pass 3/4 inch material is sieved over a No. 4 sieve. Calculate the percent of coarse aggregate or rock particles retained on the No. 4 sieve by the following:

\[
PR4 = \frac{WR4}{WT} \times 100
\]
Where:  
PR4 = Percentage of coarse aggregate or rock particles retained on the No. 4 sieve.  
WR4 = Weight of coarse aggregate or rock particles retained on the No. 4 sieve.  
WT = Total Weight of material sieved.

5.3 Recombine and thoroughly blend the plus No. 4 material with the passing No. 4 material.

5.4 The approximate 5000 gram sample of passing 3/4 inch material shall be thoroughly mixed with sufficient water to bring the sample to slightly less than its optimum moisture content.

5.5 Form a specimen by compacting the prepared soil in the six inch mold (with extension collar attached) in three equal layers to give a total compacted depth of about 5 inches. Compact each layer with 56 uniformly distributed blows from the rammer, dropping free from a height of 12 inches. While each layer is being compacted, the remainder of material shall be in a pan covered by a damp cloth. During compaction, the mold shall rest firmly on a dense, uniform, rigid, and stable foundation.

Note: Each of the following has been found to be a satisfactory base on which to rest the mold during compaction of the soil: a block of concrete, weighing not less than 200 lbs., supported by a stable foundation; a sound concrete floor; and for field application, such surfaces as found in concrete box culverts, bridges, and pavements.

5.6 When compacting granular, free-draining materials, at moisture contents which are at or above optimum, the mold shall be prepared by first sealing the bottom of the mold with waterproofing grease. All excess grease shall be wiped from the mold and base plate.

5.7 Following compaction, carefully remove the extension collar. It may be necessary to use a follower to retain the soil in the mold while removing the collar to prevent damage or disturbance of the soil below the top of the mold. Carefully trim the compacted soil even with the top of the mold by means of the straightedge. If any voids are created during trimming, these shall be filled with fine material and smoothed off. Determine the weight of compacted specimen and mold. Determine the wet density, “WD”, of the compacted soil by the following:
Where: \( \text{WD} \) = Wet density of compacted soil, lb./cu. ft.
\( \text{M1} \) = Weight of compacted specimen and mold, grams or lbs.
\( \text{M2} \) = Weight of the mold, grams or lbs.
\( \text{VM} \) = Volume of the mold, cu. ft. (See Section 3 of this procedure).

* If the weights of the compacted specimen and mold, M1, and the empty mold, M2, are measured in pounds, eliminate “453.6 (grams/lb.)” from the denominator of the above equation.

5.8 The moisture content of the sample is determined either by drying (See Subsection 5.9); or, when testing is performed in the field, the Speedy Moisture Tester may be used (See Subsection 5.10).

5.9 When the percent moisture is determined by drying, remove the material from the mold and slice vertically through the center. Take a representative minimum 600 gram sample from the full length and width of one of the cut faces. Record the weight of wet soil to the nearest 0.1 gram as “WW”. Dry the sample to constant weight at approximately 230 °F and record weight of dry soil to the nearest 0.1 gram as “DW”. The percent moisture shall be recorded to the nearest 0.1 percent. The equation below is used when the percent moisture is determined by drying the sample.

\[
\% \text{ Moisture} = \frac{\text{WW} - \text{DW}}{\text{DW}} \times 100
\]

Where: \( \text{WW} \) = Wet weight of moisture sample.
\( \text{DW} \) = Dry weight of moisture sample.

5.10 For testing in the field, the percent moisture may be determined using the Speedy Moisture Tester. Remove the material from the mold and slice vertically through the center. Obtain a minimum of 600 grams of material from the full length and width of one of the cut faces. This material is screened over a No. 4 sieve as rapidly as possible to avoid drying of the sample. A representative sample of the pass No. 4 material shall be utilized and tested in accordance with the instructional manual for that apparatus. The percent moisture of the pass No. 4 material determined
by the Speedy Moisture Tester is recorded to the nearest 0.1 percent as “W”. The moisture content of the pass 3/4 inch material is determined and recorded as “TW” to the nearest 0.1 percent by the following:

\[ TW = \frac{[W \times (100 - PR4)] + PR4}{100} \]

Where:
- TW = % moisture in pass 3/4 material.
- W = % moisture in pass No. 4 material (determined by Speedy).
- PR4 = % rock retained on the No. 4 sieve (determined in Subsection 5.2).

6. MAXIMUM DENSITY DETERMINATION

6.1 The point representing the wet density and moisture content (dry basis) is then plotted on the Typical Moisture-Density Curves (Figure 2) and the maximum wet density and optimum moisture content determined. When this plotted point falls between two moisture-density curves, a minor interpolation is necessary. The maximum dry density in lb/cu. ft. and the corresponding percent optimum moisture is then read directly or interpolated from the chart. The family of typical moisture-density curves provided should be periodically verified for the local conditions. If it is ascertained that the family of curves provided by Figure 2 is of questionable reliability for given local conditions, then an independent family of curves should be established and used.

6.2 The plotted point for wet density and moisture content should be on the dry side of the curve at or near optimum, as it is difficult to interpolate between curves for friable soils when on the wet side of the peak.

6.3 If the plotted point representing the wet density and moisture content of the compacted material is on the right of the peak, the test should be repeated using a lower moisture content. An exception to this rule must be made for those soils having high clay contents and relatively flat curves. These soils cannot readily be dried to optimum in the field due to the creation of a cloddy condition which will cause voids in the proctor. Proctors for these materials should be made as near to optimum as possible.
7. EXAMPLE

7.1 An illustration of determining the maximum dry density and optimum moisture content is described below, and shown in Figures 3 and 4:

For:

- Wet Density = 122.5 lb./cu. ft.
- Moisture Content = 18.7%

By plotting this point on the Typical Moisture-Density Curves and interpolating to the peak, it shows a point which is approximately 20 percent of the distance from Curve P to Curve Q. From the chart, the dry density for Curve P is 104.7 lb./cu. ft. @ 19.2% moisture and the dry density for Curve Q is 102.4 lb./cu. ft. @ 20.3% moisture.

By interpolation:

- Density: $104.7 - 102.4 = 2.3$
  $0.20 	imes 2.3 = 0.5$ lb./cu. ft. difference

- Moisture: $20.3 - 19.2 = 1.1$
  $0.20 	imes 1.1 = 0.2$% difference

Therefore:

- Maximum dry density = $104.7 - 0.5$
  $= 104.2$ lb./cu. ft.

- Optimum Moisture = $19.2 + 0.2 = 19.4$%

7.1.1 As an alternate to performing the interpolation procedure above, TABLE 1 can be used to determine the maximum dry density and optimum moisture content when the plotted point falls between two moisture-density curves.
**Note:** The optimum moisture and maximum dry density determinations above are for the material passing the 3/4 inch sieve. When testing field samples for comparison to proctor optimum moisture and maximum dry density, a correction to the proctor optimum moisture and maximum dry density must be made, in accordance with Arizona Test Method 227, for the percent rock which the field sample contains.

8. **REPORT**

8.1 Record the moisture and density data on the laboratory test form along with the laboratory number, material source and type, and other information required.
ARIZ 246b
April 19, 2013
Page 8

TABLE 1


**ONE POINT PROCTOR DENSITY**

<table>
<thead>
<tr>
<th>Lab. No:</th>
<th>Org No.:</th>
<th>Date:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Project No.:</td>
<td>TRACS No.:</td>
<td></td>
</tr>
<tr>
<td>Original Source:</td>
<td>Type of Material:</td>
<td></td>
</tr>
<tr>
<td>Coarse Agg. % Absorp.:</td>
<td>Coarse Agg. Bulk O.D. Sp. Gr.:</td>
<td></td>
</tr>
<tr>
<td>Proctor Method Used: Method A</td>
<td>Alternate Method D</td>
<td></td>
</tr>
<tr>
<td>Test Operator:</td>
<td>Date:</td>
<td></td>
</tr>
<tr>
<td>Supervisor:</td>
<td>Date:</td>
<td></td>
</tr>
</tbody>
</table>

### Wet Density Determination

<table>
<thead>
<tr>
<th>Volume of Mold</th>
<th>VM</th>
<th>=</th>
<th>cu. ft.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight of Mold</td>
<td>M2</td>
<td>=</td>
<td>grams</td>
</tr>
<tr>
<td>Weight of Sample and Mold</td>
<td>M1</td>
<td>=</td>
<td>grams</td>
</tr>
</tbody>
</table>

Wet Density = \( \frac{M1 - M2}{VM \times 453.6 \text{ (grams/lb.)}} \) lb./cu.ft.

*If M1 and M2 are in pounds, eliminate “453.6 (grams/lb.)” from denominator in above equation.

### Percent Moisture Determination

For either Method A or Alternate Method D, when sample is oven dried:

Wet Weight of Moisture Sample = WW = grams
Dry Weight of Moisture Sample = DW = grams

% Moisture = \( \frac{WW - DW}{DW} \times 100 \) %

For Method A, when Speedie Moisture Tester is used:

% Moisture = %

For Alternate Method D, when Speedie Moisture Tester is used:

\[ WT = \frac{WR4 \times PR4}{100} \]

% Moisture in Pass No. 4 material from Speedie = \( W = \) %

Total % Moisture = \( \frac{[W(100 - PR4)] + PR4}{100} \) %

From Typical Moisture-Density Curves:

Maximum Dry Density = MD = lb./cu. ft.
Percent Optimum Moisture = OM = %

**FIGURE 1**
# ONE POINT PROCTOR DENSITY

<table>
<thead>
<tr>
<th>Lab. No:</th>
<th>Org No.:</th>
<th>Date:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Project No.</td>
<td>TRACS No.</td>
<td></td>
</tr>
<tr>
<td>Original Source:</td>
<td>Type of Material:</td>
<td></td>
</tr>
<tr>
<td>Coarse Agg. % Absorp.:</td>
<td>Coarse Agg. Bulk O.D. Sp. Gr.:</td>
<td></td>
</tr>
<tr>
<td>Proctor Method Used: Method A</td>
<td>Alternate Method D</td>
<td>X</td>
</tr>
<tr>
<td>Test Operator:</td>
<td>Date:</td>
<td></td>
</tr>
<tr>
<td>Supervisor:</td>
<td>Date:</td>
<td></td>
</tr>
</tbody>
</table>

### Wet Density Determination

| Volume of Mold | VM = | 0.0758 cu. ft. |
| Weight of Mold | M2 = | 6608 grams pounds |
| Weight of Sample and Mold | M1 = | 10820 grams pounds |

\[
W_D = \frac{M_1 - M_2}{V_M \times 453.6} \text{ (grams/lb.)}
\]

*If M1 and M2 are in pounds, eliminate “453.6 (grams/lb.)” from denominator in above equation.

### Percent Moisture Determination

For either Method A or Alternate Method D, when sample is oven dried:

- Wet Weight of Moisture Sample = WW = ________ grams
- Dry Weight of Moisture Sample = DW = ________ grams

\[
\% \text{ Moisture} = \frac{W_W - W_D}{W_D} \times 100 = \text{______}%
\]

For Method A, when Speedie Moisture Tester is used:

\[
\% \text{ Moisture} = \text{______}%
\]

For Alternate Method D, when Speedie Moisture Tester is used:

\[
W_T = \frac{5736}{W_T} \text{, } WR_4 = \frac{1274}{PR_4} = \frac{WR_4}{W_T} \times 100 = \text{______}%
\]

\[
\% \text{ Moisture in Pass No. 4 material from Speedie} = W = \text{______}%
\]

\[
\text{Total } \% \text{ Moisture} = \frac{[W(100 - PR_4)] + PR_4}{100} = \text{______}%
\]

From Typical Moisture-Density Curves:

- Maximum Dry Density = MD = 104.2 lb./cu. ft.
- Percent Optimum Moisture = OM = 19.4 %

**REMARKS:**

- FIGURE 3
FIGURE 4

TYPICAL MOISTURE-DENSITY CURVES

<table>
<thead>
<tr>
<th>Curve</th>
<th>Max Dry Wt. lb/cu.ft</th>
<th>Optimum Moisture %</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>141.8</td>
<td>6.6</td>
</tr>
<tr>
<td>B</td>
<td>139.1</td>
<td>7.2</td>
</tr>
<tr>
<td>C</td>
<td>136.3</td>
<td>7.9</td>
</tr>
<tr>
<td>D</td>
<td>134.1</td>
<td>8.5</td>
</tr>
<tr>
<td>E</td>
<td>132.2</td>
<td>9.0</td>
</tr>
<tr>
<td>F</td>
<td>129.3</td>
<td>9.7</td>
</tr>
<tr>
<td>G</td>
<td>126.6</td>
<td>10.5</td>
</tr>
<tr>
<td>H</td>
<td>124.2</td>
<td>11.2</td>
</tr>
<tr>
<td>I</td>
<td>121.7</td>
<td>11.9</td>
</tr>
<tr>
<td>J</td>
<td>119.3</td>
<td>12.7</td>
</tr>
<tr>
<td>K</td>
<td>117.0</td>
<td>13.5</td>
</tr>
<tr>
<td>L</td>
<td>114.8</td>
<td>14.6</td>
</tr>
<tr>
<td>M</td>
<td>112.0</td>
<td>15.8</td>
</tr>
<tr>
<td>N</td>
<td>109.6</td>
<td>16.9</td>
</tr>
<tr>
<td>O</td>
<td>107.1</td>
<td>18.1</td>
</tr>
<tr>
<td>P</td>
<td>104.7</td>
<td>19.2</td>
</tr>
<tr>
<td>Q</td>
<td>102.4</td>
<td>20.3</td>
</tr>
<tr>
<td>R</td>
<td>99.9</td>
<td>21.5</td>
</tr>
<tr>
<td>S</td>
<td>97.4</td>
<td>22.7</td>
</tr>
<tr>
<td>T</td>
<td>94.5</td>
<td>24.4</td>
</tr>
<tr>
<td>U</td>
<td>92.1</td>
<td>25.8</td>
</tr>
<tr>
<td>V</td>
<td>89.9</td>
<td>27.4</td>
</tr>
<tr>
<td>W</td>
<td>87.5</td>
<td>29.5</td>
</tr>
<tr>
<td>X</td>
<td>85.0</td>
<td>30.5</td>
</tr>
<tr>
<td>Y</td>
<td>83.0</td>
<td>31.5</td>
</tr>
<tr>
<td>Z</td>
<td>81.1</td>
<td>32.5</td>
</tr>
</tbody>
</table>

Approx. 20% Below Curve P

Interpolated Max. Dry Density: 104.7 (Curve P)

\[
- \quad \frac{102.4 \text{ (Curve Q)}}{2.3 \times 0.20 = 0.5 \text{ lb./cu. ft.}} \quad 104.7 - 0.5 = 104.2 \text{ lb. / cu. ft.}
\]

Interpolated % Optimum Moisture: 20.3 (Curve Q)

\[
- \quad \frac{19.2 \text{ (Curve P)}}{1.1 \times 0.20 = 0.2\%} \quad 19.2 + 0.2 = 19.4\%
\]
PARTICLE SHAPE AND TEXTURE OF FINE AGGREGATE USING UNCOMPACTED VOID CONTENT

(A Modification of AASHTO T 304)

1. SCOPE

1.1 This method covers the determination of the "Uncompacted Void Content" of a fine aggregate for use as a measure of its angularity and texture.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of whoever uses this test method to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 This procedure provides a numerical result in terms of percent void content, determined under standardized conditions, which correlates with the particle shape and texture properties of a fine aggregate. An increase in void content indicates greater angularity and rougher texture. Lower void content results are associated with more rounded smooth particles.

1.4 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 A funnel having a volume of at least 200 cm³, or being equipped with a supplemental container to provide the required volume. (See Figure 1)

2.3 Funnel Stand - A support capable of holding the funnel firmly in position with its axis vertically in line with the axis of the measure, and the funnel opening 4.5 ± 0.1 inches above the top of the cylinder. A suitable arrangement is shown in Figure 1.
2.4 Measure - A cylinder of approximately 100 cm³ capacity. (See Figure 2)

2.5 A flat metal or plastic pan of sufficient size for containing the funnel stand, and preventing loss of material that overflows the measure during filling and strike off. The pan shall not be warped so as to prevent rocking of the apparatus during testing.

2.6 A straight metal spatula at least 1” greater than the diameter of the measure and at least 1/2" wide. The end shall be cut at a right angle to the edges. The straight edge of the spatula is used to strike off the fine aggregate. (See Figure 3)

2.7 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.8 Sieves of sizes No. 8, No. 16, No. 30, No. 50 and No. 100, conforming to the requirements of ASTM E11.

3. CALIBRATION OF MEASURE

3.1 Determine and record the weight of the dry, empty measure and a flat, glass plate slightly larger than it's diameter. Fill the measure with water at a temperature of 77 ± 1 °F. Place the glass plate on the measure, being sure that no air bubbles remain. It may be necessary to lightly coat the top edge of the measure with grease prior to determining the weight of the empty measure and glass plate. Dry the outer surfaces of the measure and determine and record the combined weight of measure, glass plate, and water.

3.2 Determine and record the volume of the measure to the nearest 0.01 cm³ by the following calculation:

\[ V = \frac{w}{0.997} \]

Where: \( V \) = volume of cylinder in cm³
\( w \) = net weight of water in grams
0.997 g/cm³ = the density of water at 77 ± 1 °F
4. SAMPLE PREPARATION

4.1 Obtain a sample of Minus No. 8 Material of sufficient size (but not less than 500 grams) to yield the quantities required in Subsection 4.3 below. The sample used for this test may either be virgin aggregate, or aggregate obtained from the extraction of a bituminous mixture.

4.2 Utilizing either a No. 100 or a No. 200 sieve, wash the sample in accordance with either Section 6 or 7 of Arizona Test Method 201. Dry the material to constant weight and sieve into size fractions as indicated in Subsection 4.3 below. Maintain the material in a dry condition in separate containers for each of the sizes specified. The sieving is to be accomplished in accordance with Arizona Test Method 201.

Note: Processing additional material may be required.

4.3 Weigh out and combine the following quantities of dry fine aggregate from each of the sizes below:

<table>
<thead>
<tr>
<th>PASS</th>
<th>RETAINED</th>
<th>WEIGHT IN GRAMS</th>
<th>ACCUM. WEIGHT</th>
</tr>
</thead>
<tbody>
<tr>
<td># 8</td>
<td># 16</td>
<td>44 ± 0.2</td>
<td>44 ± 0.2</td>
</tr>
<tr>
<td># 16</td>
<td># 30</td>
<td>57 ± 0.2</td>
<td>101 ± 0.4</td>
</tr>
<tr>
<td># 30</td>
<td># 50</td>
<td>72 ± 0.2</td>
<td>173 ± 0.6</td>
</tr>
<tr>
<td># 50</td>
<td># 100</td>
<td>17 ± 0.2</td>
<td>190 ± 0.8</td>
</tr>
</tbody>
</table>

5. PROCEDURE

5.1 If the fine aggregate has become moist, dry to constant weight and cool to room temperature.

5.2 Record the weight of the empty measure to the nearest 0.1 gram, place the funnel and measure in the funnel stand, and place the assembly in the pan described in Subsection 2.5.

5.3 Mix the test sample until it appears homogeneous. Using a finger to block the opening, pour the test sample into the funnel. Lightly level the top of the material using the end of the spatula. Remove the finger and allow the sample to fall freely into the measure.
5.4 After the funnel empties, remove excess fine aggregate from the measure by a single pass of the spatula with the edge of the blade vertical and in light contact with the top of the measure. Until this operation is complete, exercise care to avoid vibration or disturbance that could cause compaction of the fine aggregate in the measure. After strike-off, tap the measure lightly to compact the sample to make it easier to transfer the measure to the balance without spilling any of the sample. Brush adhering material from the outside of the measure and determine and record the weight of the measure and contents to the nearest 0.1 gram. (See Figure 3)

Note: The intent of this process is to allow the sample to flow freely into the measure without any vibrations or disturbance of the cylinder until the operation is complete. The cylinder may be held during strikeoff as long as there is no vibration or disturbance from the strike off process.

5.5 Collect all of the fine aggregate from the pan and measure, and repeat the procedure again.

5.6 For each determination, record the net weight of the fine aggregate in the measure. If the two net weights differ by 0.5 gram or less, average the two weights and record to the nearest 0.1 gram as the "average net weight of fine aggregate in measure", (W). If the two weights differ by more than 0.5 gram, the procedure shall be repeated until any two results are achieved which differ by 0.5 gram or less. The average of these two results is recorded to the nearest 0.1 gram as the "average net weight of fine aggregate in measure", (W).
6. **CALCULATION**

6.1 Determine and record the "Uncompacted Void Content" (U), to the nearest 0.1% by the following calculation:

\[
U = \left( \frac{V - \left( \frac{W}{G} \right)}{V} \right) \times 100
\]

Where:
- \( U \) = Uncompacted Void Content, percent.
- \( V \) = volume of measure in cm³.
- \( W \) = average net weight of fine aggregate in measure.
- \( G \) = bulk oven dry specific gravity of fine aggregate, measured in accordance with Arizona Test Method 211, "Specific Gravity and Absorption of Fine Aggregate".

6.2 For most aggregate sources the fine aggregate specific gravity does not vary much from sample to sample. It is intended that the value used in the above calculation be from a routine specific gravity test which is representative of the fine aggregate. A difference in specific gravity of 0.05 will change the calculated "Uncompacted Void Content" about one percent.

7. **REPORT**

7.1 The "Uncompacted Void Content" (U), to the nearest 0.1%.

7.2 The bulk oven dry specific gravity of the fine aggregate (G), to the nearest 0.001.
FUNNEL, FUNNEL STAND, AND MEASURE

FIGURE 1
MEASURE

1.63 in. approx.

1.52 ± 0.02 in.

Copper Pipe

Nominal 100 cm³ Measure

Epoxy Joint

¾ in. Plate

¾ in. Drilled Hole, Approx.
¾ in. Deep - Used for Centering Container

3.37 ± 0.02 in.

0.150 ± 0.005 in.

FIGURE 2
ALTERNATE PROCEDURES FOR SIEVING OF COARSE AND FINE GRADED SOILS AND AGGREGATES

(An Arizona Method)

SCOPE

1. (a) This test method contains five alternate procedures for sieving and determining the sieve analysis of fine and coarse graded soils and aggregates.

   (b) This test method must be used in conjunction with Arizona Test Method 201. In general, the method for sample preparation, sieving, and calculation of the sieve analysis as described in Arizona Test Method 201 will be followed, with the changes and additional information being outlined and described herein for each alternate procedure shown below.

   1) Alternate #1 (Section 2) - Fine screening of aggregate which contains a small amount of Plus No. 4 (100% pass 3/8") material, e.g., fine aggregate for concrete.

   2) Alternate #2 (Section 3) - Utilizing a No. 8 screen in the coarse sieving, and a fine sieve analysis not being required.

   3) Alternate #3 (Section 4) - Subjecting the entire sample to washing, and a minus No. 4 split for fine screening is not required.

   4) Alternate #4 (Section 5) - Subjecting the entire sample to washing, and a minus No. 4 split for fine screening is required.

   5) Alternate #5 (Section 6) - Washing of coarse aggregate (Plus No. 4) to remove clinging particles and adjusting the fine sieve analysis.

ALTERNATE #1

2. Fine screening of aggregate which contains a small amount of Plus No. 4 (100% pass 3/8") material, e.g., fine aggregate for concrete. An example of this procedure is shown in Figure 1.
## Arizona Department of Transportation
### Soil and Aggregate Tabulation

<table>
<thead>
<tr>
<th>Lab Number</th>
<th>Code</th>
<th>Org Number</th>
<th>Matl Type</th>
<th>Test Code</th>
<th>Test</th>
<th>Size</th>
<th>Size %</th>
<th>Original Source</th>
<th>Project Engineer</th>
<th>Project Number</th>
<th>Tracs Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>93-2222</td>
<td>7777</td>
<td>8888</td>
<td>FA+</td>
<td>A</td>
<td>P</td>
<td>-</td>
<td>+</td>
<td>Pit # 5555</td>
<td>B. Smith</td>
<td>F-099-9 (99)</td>
<td>H999901C</td>
</tr>
</tbody>
</table>

**Example for Fine Screening When Fine Aggregate Contains a Small Amount of Plus No. 4 Material (Alternate #1)**

### ARIZ 201

<table>
<thead>
<tr>
<th>Fraction</th>
<th>% Retained</th>
<th>% Passing</th>
<th>Spec</th>
<th>Modulus</th>
</tr>
</thead>
<tbody>
<tr>
<td>3&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 1/2&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 1/2&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
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<tr>
<td>3/8&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#4</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Wet Sample Pre-Wet Weight (Wet WT. Of #4):**

**Cumulative Retention:**

<table>
<thead>
<tr>
<th>% Ret.</th>
<th>% Pass</th>
<th>Spec</th>
<th>Modulus</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>9%</td>
<td>95%</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Dry Wt. of #4 Split:**

<table>
<thead>
<tr>
<th>Fraction</th>
<th>% Retained</th>
<th>% Passing</th>
<th>Spec</th>
<th>Modulus</th>
</tr>
</thead>
<tbody>
<tr>
<td>#8</td>
<td>13</td>
<td>89</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#10</td>
<td>42</td>
<td>86</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#16</td>
<td>91</td>
<td>70</td>
<td>45-10</td>
<td></td>
</tr>
<tr>
<td>#20</td>
<td>67</td>
<td>12</td>
<td>58</td>
<td></td>
</tr>
<tr>
<td>#24</td>
<td>12</td>
<td>22</td>
<td>36</td>
<td></td>
</tr>
<tr>
<td>#41</td>
<td>41</td>
<td>79</td>
<td>8-30</td>
<td></td>
</tr>
<tr>
<td>#100</td>
<td>121</td>
<td>22</td>
<td>7</td>
<td>0-10</td>
</tr>
<tr>
<td>#200</td>
<td>24</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#220</td>
<td>2</td>
<td>3.0</td>
<td>0-4</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>547</td>
<td>100%</td>
<td>Total Cumulative % Ret.</td>
<td></td>
</tr>
</tbody>
</table>

**Remarks:**

- If total sample is washed:
  - Unwashed Wt. = 547
  - Washed Wt. = 15

**Fine Factor:**

<table>
<thead>
<tr>
<th>% Ret.</th>
<th>% Pass</th>
<th>Spec</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.0</td>
<td>0-4</td>
<td></td>
</tr>
</tbody>
</table>

**FINISHES MODULUS:**

- Total
- White
- Yellow
- Blue

**Test Details:**

- Received Date: 12-08-93
- Test Operator & Date: Joe DiBiaso 12-08-93
- Supervisor & Date: Ted Trudel 12-08-93

**FIGURE 1**
(a) A representative minimum 500 gram sample is obtained and dried to constant weight. Allow the sample to cool, and record the weight to the nearest gram as the "Unwashed Wt", (562 in the example).

(b) Subject the sample to either mechanical or hand washing.

(c) Dry the sample to constant weight and allow to cool. Record weight to the nearest gram as "Washed Wt" and also as the fine sieve "Total Dry Weight", (547).

(d) Determine and record "Elutriation" by subtracting the "Washed Wt" from the "Unwashed Wt", (562 - 547 = 15).

(e) Additional sieves (Plus No. 4) as required are added to the nest of sieves used in the fine screening.

(f) The sample is subjected to screening and the weight retained on each sieve and in the pan is recorded.

(g) The sum of the individual weights retained for each sieve is compared to the weight of sample prior to sieving (Washed Wt. or fine sieve Total Dry Weight). Adjust or resieve as necessary.

(h) Determine the sieve analysis of the material. The factor for calculating the sieve analysis is determined by dividing 100 by the "Unwashed Wt", (100/562 = 0.177936). The calculation of the percent passing each sieve is continuous through the entire sieve analysis.

ALTERNATE #2

3. Utilizing a No. 8 screen in the coarse sieving, and a fine sieve analysis not being required. An example is given in Figure 2.

(a) A representative sample is obtained and the weight recorded as the "Coarse Sieve Total", (14683 in the example).

(b) Subject the sample to coarse sieving using the No. 8 as the smallest sieve. If desired, the material may be screened using a No. 4 as the smallest sieve, and the pass No. 4 material further separated into No. 8 and pass No. 8 fractions.
### ARIZONA DEPARTMENT OF TRANSPORTATION
SOIL AND AGGREGATE TABULATION

<table>
<thead>
<tr>
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<th>PROJ CODE</th>
<th>ORG NUMBER</th>
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**EXAMPLE FOR USING A NO. 8 SIEVE IN THE COARSE SIEVING AND NO FINE SIEVE ANALYSIS REQUIRED - (ALTERNATE #2).**

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**REMARKS**

| EXAMPLE FOR USING A NO. 8 SIEVE IN THE COARSE SIEVING AND NO FINE SIEVE ANALYSIS REQUIRED - (ALTERNATE #2).**

<table>
<thead>
<tr>
<th>PROJECT ENGINEER / SUPERVISOR</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. Smith</td>
</tr>
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**SIEVING**

**TEST OPERATOR & DATE**

| Joe Doood         | 12-03-93         |

**SUPERVISOR & DATE**

| Ted Henderson     | 12-03-93         |

**FIGURE 2**
(c) The weight of material retained on each of the sieves No. 8 and larger, and the weight of pass No. 8 material is recorded on the laboratory sieve analysis card, modifying the card for the pass No. 8 material as shown in the example.

(d) The sum of the individual weights retained for each sieve is compared to the weight of the sample prior to sieving (Coarse Sieve Total). Adjust or resieve as needed.

(e) Determine the sieve analysis of the material. The coarse sieve factor is determined by dividing 100 by the "Coarse Sieve Total", \(100/14683 = 0.006811\). The calculation of the percent passing each sieve is continuous through the entire sieve analysis.

ALTERNATE #3

4. Subjecting the entire sample to washing, and a minus No. 4 split for fine screening is not required. An example of this procedure is given in Figure 3.

(a) A representative sample is obtained, dried to constant weight, allowed to cool, and the weight recorded as the "Coarse Sieve Total" and also as the "Unwashed Weight", (1137 in the example).

(b) Subject the sample to hand washing.

(c) Dry the sample to constant weight and allow to cool. The weight of the sample is recorded as the "Washed Wt", (1118).

(d) Determine "Elutriation" by subtracting the "Washed Wt" from the "Unwashed Wt", \(1137 - 1118 = 19\).

(e) Subject the sample to coarse screening and record the weight retained on each sieve and in the pan, except the weight of pass No. 4 material is not recorded in the coarse sieve area, but rather as the fine sieve "Total Dry Weight", (468).

(f) Determine and record the weight of pass No. 4 for coarse sieve analysis by adding the "elutriation" and the fine sieve "Total Dry Weight", \(468 + 19 = 487\).

(g) The sum of the individual weights retained for each sieve is compared to the weight of sample prior to sieving (Washed Wt.). Adjust or resieve as necessary.
### ARIZONA DEPARTMENT OF TRANSPORTATION

#### SOIL AND AGGREGATE TABULATION

<table>
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<tr>
<th>LAB NUMBER</th>
<th>PROJ CODE</th>
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**Test Information**

- **Test No.**: 3
- **Lot Code**: A
- **Sampled By**: A. Jones
- **Sampled From**: COLD FEED
- **Date**: 10/07/93
- **Time**: 12:10 PM
- **Am/PM**: PM
- **Location**: STATION P
- **Milepost, Exit/OT Number**: 
- **Original Source**: PIT # 4444
- **Project Engineer**: B. Smith
- **Project Number**: F-039-9 (99)
- **Trac Number**: H99990C
- **Remarks**: Example when entire sample is washed and a minus No. 4 split for fine sieving is not required - (ALTERNATE # 3).

#### ARIZ 201

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**Wet Sample Preweight**

- **Wet WT of #4**: -
- **Split Wet WT.**: -

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**Total Sample is Washed, Unwashed WT. = 111.7**

**Liquid Limit (LL)**

- **T-02**: 

**Plastic Limit (PL)**

- **T-90**: 

**Plasticity Index (PI) = LL - PL**

- **T-90**: 

**Abrasion Method (A,B,C,D)**

- **T-96**: 

**Absorption, hyd**: 

- **T-05**: ARIZ 211

**Specific Gravity, SSD**: 

- **T-05**: ARIZ 211

**Specific Gravity, OD**: 

- **T-05**: ARIZ 211

**Proctor Method**: 

- **Optimum Moisture**: 
- **Max. Dry Density**: 

**Sample Equivalent**: 

- **T-176**: ARIZ 242 (MAFC)

**Fractured Faces Weight (WF)**

- **Total Sample Weight (Wt)**

**Fractured Faces (FP) = Wt x 100 ARIZ 212**

**Wet Weight (W)**

- **Dry Weight (D)**

**Moisture Content = WF x 100**: 

- **T-05**: ARIZ 212

**Flakiness Index**: ARIZ 233

**Carbonates**: ARIZ 235

**pH**: ARIZ 236 OR 237

**Relative (ohm-cm)**: ARIZ 236

**Soluble Sulfate (PPM)**: ARIZ 237

**Fineness Modulus**: 

- **Total Cumulative % Ret. 100**: 

**White, Yellow, Blue**

---

**Figure 3**

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**Date**: 10-07-93

**Test Operator & Date**: Joe Doard 10-07-93

**Supervisor & Date**: Ted Hedlund 10-08-93
(h) Sieve the pass No. 4 material and record the weight retained for each sieve and in the pan.

(i) The sum of the individual weights retained for each sieve is compared to the weight of sample prior to sieving (fine sieve Total Dry Weight). Adjust or resieve as necessary.

(j) Determine the sieve analysis of the material. The coarse sieve factor is determined by dividing 100 by the "Coarse Sieve Total", \( \frac{100}{1137} = 0.087951 \). A fine sieve factor is not determined. The calculation of the percent passing each sieve is continuous through the entire sieve analysis.

ALTERNATE #4

5. Subjecting the entire sample to washing, and a minus No. 4 split for fine screening is required. An example of this procedure is given in Figure 4.

(a) A representative sample is obtained, dried to constant weight, allowed to cool, and the weight recorded as the "Coarse Sieve Total" and also as the "Unwashed Weight", (4893 in the example).

(b) Subject the sample to hand washing.

(c) Dry the sample to constant weight and allow to cool. The weight of the sample is recorded as the "Washed Wt", (4674).

(d) The elutriation of the total sample is determined by subtracting the "Washed Wt" from the "Unwashed Wt", (4893 - 4674 = 219).

(e) The sample shall be subjected to coarse screening and the weight retained on each sieve and in the pan is recorded, except the weight of pass No. 4 material is not recorded in the coarse sieve area, but rather as "Wt. of - #4" (2521).

(f) The sum of the individual weights retained for each sieve is compared to the weight of sample prior to sieving (Washed Wt.). Adjust or resieve as necessary.

(g) The passing No. 4 weight for coarse sieve analysis is the combination of the weight of pass No. 4 and the elutriation of the total sample, (2521 + 219 = 2740).

(h) The weight of the passing No. 4 material split for fine sieving is recorded as the fine sieve "Total Dry Weight", (665).
ARIZONA DEPARTMENT OF TRANSPORTATION
SOIL AND AGGREGATE TABULATION

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<td>H-099901C</td>
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EXAMPLE WHEN ENTIRE SAMPLE IS WASHED AND A
MINUS NO. 4 SPLIT FOR FINE SIEVING IS
REQUIRED - (ALTERNATIVE # 4).

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DRIED SAMPLE PRE-WEIGHT | DRIED WT. OF #4 | 665 |
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</table>

FIGURE 4
The elutriation value for fine sieve analysis is determined as follows:

\[
\text{Elutriation for Fine Sieve Analysis} = \frac{\text{Wt. of Passing No. 4 Material Split for Fine Sieving, (Fine Sieve "Total Dry Weight")}}{\text{Wt. of - #4}} \times \frac{\text{Elutriation of Total Sample}}{}
\]

In the example in Figure 4:

Elutriation for Fine Sieve Analysis = (665/2521) x 219 = 58

(j) The "Dry Wt. of Pass #4 Split" is determined by adding the fine sieve total dry weight and the calculated elutriation for fine sieve analysis, (665 + 58 = 723).

(k) The pass No. 4 material is sieved and the weight retained for each sieve and in the pan is recorded.

(l) The sum of the individual weights retained for each sieve is compared to the weight of sample prior to sieving (fine sieve Total Dry Weight). Adjust or resieve as necessary.

(m) Determine the sieve analysis of the material.

**ALTERNATE #5**

6. Washing of coarse aggregate (Plus No. 4) to remove clinging particles and adjusting the fine sieve analysis. An example of this procedure is shown in Figures 5 and 6.

(a) A representative sample shall be prepared, screened, and the sieve analysis calculated as described in Arizona Test Method 201. (When this alternate procedure is used as a referee method, the sample shall be dried to constant weight prior to sieving.)
### ARIZONA DEPARTMENT OF TRANSPORTATION

#### SOIL AND AGGREGATE TABULATION

**FIGURE 5**

**EXAMPLE OF WASHING COARSE (PLUS NO. 4) AGGREGATE AND ADJUSTING FINE SIEVE ANALYSIS — (ALTERNATE #5).**

**ARIZ 201**

<table>
<thead>
<tr>
<th>% OVERSIZE</th>
<th>COARSE FACTOR</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>004918</td>
</tr>
</tbody>
</table>

**WET SAMPLE PREWEIGHT**

| WET WT. OF -44 | 22940 |

**44 SPLIT WET WT.**

| 11480 |

**CUMULATIVE % RET. FINENESS MODULES**

| 5% | 100 |

**WEIGHTS RETAINED**

| 2 1/2" | 0 | 100 |
| 2" | 12 | 11 | 29 |
| 1 1/2" | 4 | 6 | 43 |
| 1" | 5 | 7 | 47 |
| 3/4" | 5 | 13 | 56 | 60-100 |
| 1/2" | 5 | 2 | 51 |
| 3/8" | 2 | 49 |
| 1/4" | 5 | 3 | 53 |
| 1/8" | 4 | 2 |
| # | 4 | 1 |

**TOTAL**

| 22 | 6 | 3 | 3 |

**T = AASHTO TESTS**

| Liquid Limit (LL) | T - 89 |
| Plastic Limit (PL) | T - 90 |
| Plasticity Index (PI) = LL - PL | T - 90 |
| Abrasion Method (A,B,C,D) | T - 96 |

**SPECIFICATIONS**

- @ 100 Revolutions
- @ 500 Revolutions
- A - ARIZ 225
- G - ARIZ 226
- D - ARIZ 228
- AO - ARIZ 245
- AI - ARIZ 233
- AD - ARIZ 246

**FRACTURED FACES WEIGHT (WF)**

- Total Sample Weight (W): 100
- Fractured Faces (WF) = Wf / Wg ≤ 100 ARIZ 212

**WET WEIGHT (W)**

| 100 |

**DRY WEIGHT OF -44 SPLIT**

| 9 | 40 |
| 35-40 |

**FINE FACTOR**

| 542 |

**WEIGHTS RETAINED**

| 101 |
| 12 |
| 30 |
| 33 |
| 44 |
| 42 |
| 35 |
| 2 |

**TOTAL**

| 4 | 1 | 9 |

**FINENESS MODULUS**

| 11.6 |

**RECEIVED DATE**

| 11-15-93 |

**TEST OPERATOR & DATE**

| Joe Doood 11-16-93 |

**SUPERVISOR & DATE**

| Ted Headman 11-16-93 |

**FIGURE 5**
DETERMINATION OF PERCENT MINUS NO. 200 ON COARSE AGGREGATE
AND TOTAL PERCENT PASS NO. 200 IN SAMPLE

Project No. F-099-9(99) / H.99990/C, Lab. No. 93-9999

Unwashed Weight of Plus No. 4 = \( \frac{4967}{4941} \)
Washed Weight of Plus No. 4 = \( \frac{4967}{4941} \)

Calculate the "Percent Minus No. 200 on Coarse Aggregate" by the following, and record result to the nearest 0.01%:

\[
\text{Percent Minus No. 200 on Coarse Aggregate} = \left( \frac{ \text{Unwashed Weight of Plus No. 4} - \text{Washed Weight of Plus No. 4} }{ \text{Unwashed Weight of Plus No. 4} } \right) \times 100
\]

\[
= \left( \frac{4967}{4941} \right) - \left( \frac{4967}{4941} \right) \times 100
\]

Percent Minus No. 200 on Coarse Aggregate = 0.52

Calculate the "Total Percent Pass No. 200 in Sample" by the following, and record result to the nearest 0.1%:

\[
\text{Total Percent Pass No. 200 in Sample} = \left[ \frac{\% \text{ Pass No. 200 from Fine Sieving}}{100} \right] + \left[ \frac{\% \text{ Pass No. 4}}{100} \right] \times \left[ \frac{\text{Percent Minus No. 200 on Coarse Aggregate}}{100} \right]
\]

\[
= \left( 11.3 \right) + \left[ 100 - \left( \frac{49}{49} \right) \right] \times \left( 0.52 \right) \times \left( \frac{100}{100} \right)
\]

Total Percent Pass No. 200 in Sample = 11.6

FIGURE 6
(b) Either the entire amount, or a representative sample of approximately 5000 grams, whichever is less, of the Plus No. 4 material from screening shall be combined, dried to constant weight, allowed to cool, and the weight recorded as the "Unwashed Weight of Plus No. 4", (4967 in the example in Figure 6).

(c) Subject the sample to hand washing.

(d) Dry the sample to constant weight and allow to cool. Record the weight to at least the nearest gram as the "Washed Weight of Plus No. 4", (4941 in the example in Figure 6).

(e) Utilizing the equations shown in Figure 6, determine and record the "Percent Minus No. 200 on Coarse Aggregate" to the nearest 0.01%, (0.52); and the "Total Percent Pass No. 200 in Sample" to the nearest 0.1%, (11.6).

(f) Compare the percent Pass No. 200 from fine sieving with the "Total Percent Pass No. 200 in Sample". If the difference is less than or equal to 1.0, replace the percent Pass No. 200 from fine sieving with the "Total Percent Pass No. 200 in Sample" and, if necessary, adjust the value for percent retained on the No. 200 sieve. (In the example in Figure 5, the 11.3 percent Pass No. 200 from fine sieving has been replaced with the 11.6 "Total Percent Pass No. 200 in Sample", and the percent retained on the No. 200 sieve has been adjusted from 3 to 2.) If the difference is greater than 1.0, another sample shall be obtained and the entire sample subjected to washing and the gradation determined as specified in Section 4 or 5.

REPORT

7. (a) The sieve analysis shall be reported as shown in Figures 1 through 5 for the particular example which is applicable.

(b) When applicable, the determination of "Percent Minus No. 200 on Coarse Aggregate" and "Total Percent Pass No. 200 in Sample" is reported as shown in the example given in Figure 6.

(c) A blank Soils and Aggregate Tabulation laboratory card is provided in Figure 7.

(d) A blank form for the determination of "Percent Minus No. 200 on Coarse Aggregate" and "Total Percent Pass No. 200 in Sample" is provided in Figure 8.
### ARIZ 201

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<th>Weight Retained</th>
<th>% Ret.</th>
<th>% Pass</th>
<th>Coarse Factor</th>
<th>Spec's</th>
<th>% Ret.</th>
<th>Spec's</th>
<th>Cumulative % Ret.</th>
<th>Fineness Modulus</th>
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**Wet Sample Preweight:**

- 3" SPLIT WET WT. =

**T = AASHTO Test:**

- Liquid Limit (LL)  
- Plastic Limit (PL)  
- Plasticity Index (PI) = LL - PL
- Abrasion Method (A,B,C,D)
- Proctor Method
- Optimum Moisture
- Max. Dry Density
- Sand Equivalent

**Dry WT. of #4 SPLIT**

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<th>% Pass</th>
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</table>

**Wet WT. of 3" SPLIT WET WT. =**

**FRAC**

- Fractured Faces Weight (WF)
- Total Sample Weight (Ws)
- Fractured Faces (FF) = Wf x 100
- Wet Weight (W)
- Dry Weight (O)
- Moisture Content = Wfd x 100
- Flatness Index
- Carbonates
- pH
- Resistivity (ohm-cm)
- Soluble Sulfur (PPM)

**Fineness Modulus = Total Cumulative % Ret. / 100**

**FIGURE 7**
DETERMINATION OF PERCENT MINUS NO. 200 ON COARSE AGGREGATE AND TOTAL PERCENT PASS NO. 200 IN SAMPLE

Project No. ___________________________ Lab. No. _________

Unwashed Weight of Plus No. 4 = _________

Washed Weight of Plus No. 4 = _________

Calculate the "Percent Minus No. 200 on Coarse Aggregate" by the following, and record result to the nearest 0.01%:

\[
\text{Percent Minus No. 200 on Coarse Aggregate} = \frac{\text{Unwashed Weight of Plus No. 4} - \text{Washed Weight of Plus No. 4}}{\text{Unwashed Weight of Plus No. 4}} \times 100
\]

Percent Minus No. 200 on Coarse Aggregate = _________

Calculate the "Total Percent Pass No. 200 in Sample" by the following, and record result to the nearest 0.1%:

\[
\text{Total Percent Pass No. 200 in Sample} = \left[ \frac{\% \text{ Pass No. 200 from Fine Sieving}}{100} \right] + \left[ \frac{\% \text{ Pass No. 4}}{100} \times \frac{\text{Percent Minus No. 200 on Coarse Aggregate}}{100} \right]
\]

Total Percent Pass No. 200 in Sample = _________

FIGURE 8
REMOLDED RING SAMPLES FOR DIRECT SHEAR, SWELL, AND CONSOLIDATION

(An Arizona Method)

SCOPE

1. (a) This procedure describes the method used to prepare remolded ring samples of soil for further testing of direct shear, swell, and consolidation. Samples are generally remolded at 95% or 97% of proctor maximum dry density and at minus 3% of optimum moisture. These values can be changed to suit the person ordering the test.

(b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:
(a) Brass ring(s) with an inside diameter of 61.47 ± 0.03 mm (2.420 ± 0.001 inches), a height of 25.40 ± 0.05 mm (1.000 ± 0.002 inch), and a nominal wall thickness of 1.02 mm (0.040 inches). The brass ring(s) shall fit snugly inside the loading cylinder with no discernible free play in any direction.

(b) Baseplate and loading cylinder, as shown in Figure 1.

(c) Metal ram with a nominal outside diameter of 63.5 mm (2.5 inches).

NOTE: The brass ring(s), baseplate, loading cylinder, and ram should be free of bumps, dents, scratches, rust, dirt, and corrosion.

(d) Rubber mallet.

(e) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.1 gram.

SAMPLING

3. (a) Obtain 34 to 45 kilograms (75 to 100 pounds) of material for a proctor density test.

(b) For the remolded ring samples, obtain 600 grams of minus 4.75 mm (No. 4) material from material for proctor density test. This amount of material is sufficient to remold 3 ring samples. Material should be split in accordance with AASHTO R 76.

NOTE: Direct shears require 3 rings. Swell and consolidation tests require only 1 ring.

(c) Obtain approximately 100 grams of the minus 4.75 mm (No. 4) material for determination of percent hydroscopic moisture content.

PROCEDURE

4. (a) Determine the percent hydrosopic moisture content of the minus 4.75 mm (No. 4) material in accordance with AASHTO T 265.
(b) Perform a Method A Proctor Density test in accordance with Arizona Test Method 225 or 232.

(c) When the weight of soil and water "$W_2$" has been calculated as shown in the examples given in Figure 2 and Figure 3 (a blank worksheet is given in Figure 4), the soil and water must be thoroughly mixed, put into a sealed container, and left to cure for at least 12 hours.

(d) After the soil has cured, the ring is ready to be loaded. Place ring in the bottom of the assembled baseplate and loading cylinder. Using a funnel, pour the soil mix into the cylinder. If it is necessary, gently tamp the material with your fingers. When all of the soil is loaded into the cylinder, insert the ram.

(e) Using the rubber mallet, strike the ram 3 or 4 times. Remove the ram from the cylinder and, using a spatula, clean the soil from edge of ring. Insert ram and strike it 2 or 3 times.

(f) Remove the ram from the cylinder and inspect the ring. If the ring is not showing, clean the soil from the edge of the ring using spatula, insert ram, and strike 2 or 3 times. If the ring is showing, remove loading cylinder from baseplate, and push the ring and soil gently out using the ram.

(g) From this point on, direct shears follow ASTM D 3080, swell tests follow ASTM D 4546, and consolidation tests follow ASTM D 2435.
Baseplate and Loading Cylinder

ALL DIMENSIONS ARE NOMINAL.
REMOLDED SHEAR / SWELL / CONSOLIDATION
WORKSHEET

Lab No.: 96-999 Date: 10-25-96

Proctor Maximum Dry Density: 1884 kg/m³ ——— lb./cu. ft.

Optimum Percent Moisture: 12.3

Maximum Dry Density at 95% or 97% (D): 1789 kg/m³ ——— lb./cu. ft.

Percent Optimum Moisture minus 3% (w₁): 9.3

Percent Hydroscopic Moisture Content (w): 1.9

Brass Ring Volume/Density Factor (F): 0.0754 (for density in kg/m³) or 1.2074 (for density in lb./cu. ft.)

W = Weight of Dry Soil, grams

W₁ = Weight of Soil with Hydroscopic Moisture, grams

W₂ = Weight of Soil and Water, grams

w₂ = Weight of Water Added/Required, grams

W = D x F

W₁ = W x (1.00 + w₁/100)

W₂ = W x (1.00 + w₁/100)

w₂ = W₂ - W₁

W = 134.9

W₁ = 137.5

W₂ = 147.4

w₂ = 9.9

Test Operator: Joe DoGood

FIGURE 2
## REMOLDED SHEAR / SWELL / CONSOLIDATION WORKSHEET

<table>
<thead>
<tr>
<th>Lab No.</th>
<th>96-999</th>
<th>Date</th>
<th>10-25-96</th>
</tr>
</thead>
</table>

Proctor Maximum Dry Density: \( \frac{kg}{m^3} \) \( 117.6 \) lb./cu. ft.

Optimum Percent Moisture: 12.3

Maximum Dry Density at 95% or 97% (D): \( \frac{kg}{m^3} \) \( 111.7 \) lb./cu. ft.

Percent Optimum Moisture minus 3% \( (w_1) \): 9.3

Percent Hydroscopic Moisture Content (w): 1.9

Brass Ring Volume/Density Factor (\( F \)) 0.0754 (for density in \( kg/m^3 \)) or 1.2074 (for density in lb./cu. ft.)

\[
W = \text{Weight of Dry Soil, grams} \quad W = D \times F \quad W = 134.9
\]

\[
W_1 = \text{Weight of Soil with Hydroscopic Moisture, grams} \quad W_1 = W \times (1.00 + w/100) \quad W_1 = 137.5
\]

\[
W_2 = \text{Weight of Soil and Water, grams} \quad W_2 = W \times (1.00 + w_1/100) \quad W_2 = 147.4
\]

\[
w_2 = \text{Weight of Water Added/Required, grams} \quad w_2 = W_2 - W_1 \quad w_2 = 9.9
\]

Test Operator

Joe Doood

FIGURE 3
REMOVED SHEAR / SWELL / CONSOLIDATION
WORKSHEET

Lab No.: __________________  Date: __________________

Proctor Maximum Dry Density: ___________ kg/m$^3$ ___________ lb./cu. ft.

Optimum Percent Moisture: ___________

Maximum Dry Density at 95% or 97% (D): ___________ kg/m$^3$ ___________ lb./cu. ft.

Percent Optimum Moisture minus 3% ($w_1$): ___________

Percent Hydroscopic Moisture Content ($w$): ___________

Brass Ring Volume/Density Factor (F): 0.0754 (for density in kg/m$^3$) or 1.2074 (for density in lb./cu. ft.)

$W$ = Weight of Dry Soil, grams $W = D \times F$ $W = __________$

$W_1$ = Weight of Soil with Hydroscopic Moisture, grams $W_1 = W \times (1.00 + w/100)$ $W_1 = __________$

$W_2$ = Weight of Soil and Water, grams $W_2 = W \times (1.00 + w_1/100)$ $W_2 = __________$

$w_2$ = Weight of Water Added/Required, grams $w_2 = W_2 - W_1$ $w_2 = __________$

Test Operator __________________________

FIGURE 4
COMBINED COARSE AND FINE AGGREGATE 
SPECIFIC GRAVITY AND ABSORPTION

(An Arizona Method)

1. SCOPE

(a) This procedure describes the method which is used to determine the combined coarse and fine aggregate specific gravity and absorption when the specific gravity and absorption of the fine aggregate and the coarse aggregate are known. Arizona Test Method 210 describes the procedure for determining the coarse aggregate specific gravity and absorption. Arizona Test Method 211 describes the procedure for determining the fine aggregate specific gravity and absorption.

(b) This procedure also contains provisions for calculating the combined specific gravity and combined absorption when it is desired to perform those calculations for coarse aggregate, fine aggregate, and mineral admixture.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. CALCULATIONS

(a) Combined Bulk (Oven Dry), Bulk (SSD), or Apparent specific gravity for coarse aggregate and fine aggregate is determined by the following:

\[
\text{Combined Specific Gravity} = \frac{P_c + P_f}{\frac{P_c}{G_c} + \frac{P_f}{G_f}}
\]

Where:
- \( P_c \) = Percent of Coarse Aggregate
- \( P_f \) = Percent of Fine Aggregate
- \( P_c + P_f = 100 \)
- \( G_c \) = Specific Gravity of Coarse Aggregate
- \( G_f \) = Specific Gravity of Fine Aggregate
Example:

\[ P_c = 41 \]
\[ P_f = 59 \]
\[ G_c = 2.597 \]
\[ G_f = 2.626 \]

Combined Specific Gravity = \[ \frac{41 + 59}{2.597 + 2.626} = 2.614 \]

(b) Combined Bulk (Oven Dry), Bulk (SSD), or Apparent specific gravity for coarse aggregate, fine aggregate, and mineral admixture, is determined by the following:

\[
\text{Combined Specific Gravity} = \frac{P_c + P_f + P_{\text{admix}}}{G_c + G_f + G_{\text{admix}}} \]

Where:
- \( P_c \) = Percent of Coarse Aggregate
- \( P_f \) = Percent of Fine Aggregate
- \( P_{\text{admix}} \) = Percent of Mineral Admixture, by weight of the mineral aggregate

\[ P_c + P_f + P_{\text{admix}} = 100 + \% \text{ Mineral Admixture} \]

\[ G_c = \text{Specific Gravity of Coarse Aggregate} \]
\[ G_f = \text{Specific Gravity of Fine Aggregate} \]
\[ G_{\text{admix}} = \text{Specific Gravity of Mineral Admixture} \]

Type I or II Cement = 3.14
Type IP Cement = 3.00
Hydrated Lime = 2.20

Example:

\[ P_c = 41 \]
\[ P_f = 59 \]
\[ P_{\text{admix}} = 1.0\% \]
\[ G_c = 2.597 \]
\[ G_f = 2.626 \]

Type of Mineral Admixture = Hydrated Lime
Mineral Admixture Specific Gravity = 2.20

Combined Specific Gravity = \[ \frac{41 + 59 + 1.0}{2.597 + 2.626 + 2.20} = 2.609 \]
(c) Combined absorption for coarse aggregate and fine aggregate is determined by the following:

\[
\text{Combined Absorption} = \frac{(P_c \times A_c) + (P_f \times A_f)}{P_c + P_f}
\]

Where:
- \(P_c\) = Percent of Coarse Aggregate
- \(P_f\) = Percent of Fine Aggregate
- \(P_c + P_f\) = 100
- \(A_c\) = Percent water absorption of coarse aggregate
- \(A_f\) = Percent water absorption of fine aggregate

NOTE: The equation shown above for combined absorption differs from that currently shown in Arizona Test Methods 210 and 211. In the future, appropriate changes in these test methods will be made to reflect the above equation, but in the interim period the calculation shall be performed as shown above.

Example:
- \(P_c = 41\)
- \(P_f = 59\)
- \(A_c = 1.51\)
- \(A_f = 1.43\)

\[
\text{Combined Absorption} = \frac{(41 \times 1.51) + (59 \times 1.43)}{41 + 59} = 1.46\%
\]

(d) Combined absorption for coarse aggregate, fine aggregate, and mineral admixture, is determined by the following:

\[
\text{Combined Absorption} = \frac{(P_c \times A_c) + (P_f \times A_f) + (P_{admix} \times A_{admix})}{P_c + P_f + P_{admix}}
\]
Where: \( P_c \) = Percent of Coarse Aggregate
\( P_f \) = Percent of Fine Aggregate
\( P_{\text{admix}} \) = Percent of Mineral Admixture, by weight of the mineral aggregate
\( P_c + P_f + P_{\text{admix}} \) = 100 + % Mineral Admixture
\( A_c \) = Percent water absorption of coarse aggregate
\( A_f \) = Percent water absorption of fine aggregate
\( A_{\text{admix}} \) = Percent water absorption of mineral admixture (assumed to be 0.0%)

Example:
\[
\begin{align*}
P_c &= 41 \\
P_f &= 59 \\
P_{\text{admix}} &= 1.0 \\
A_c &= 1.51 \\
A_f &= 1.43 \\
A_{\text{admix}} &= 0.0
\end{align*}
\]

Combined Absorption = \( \frac{(41 \times 1.51) + (59 \times 1.43) + (1.0 \times 0.0)}{41 + 59 + 1.0} \) = 1.45%

3. REPORT

(a) Report combined Bulk (Oven Dry), Bulk (SSD), or Apparent specific gravity to the nearest 0.001, and indicate the type of specific gravity, whether Bulk (Oven Dry), Bulk (SSD), or Apparent.

(b) Report combined absorption to the nearest 0.01%.

(c) Report whether the combined specific gravity and absorption is for coarse aggregate and fine aggregate, or whether the combined specific gravity and absorption is for coarse aggregate, fine aggregate, and mineral admixture.
SERIES 300
CONCRETE
METHOD OF ADJUSTING CONCRETE MIXES FOR VARIATION IN AGGREGATE MOISTURE CONTENT
(An Arizona Method)

Scope

1. This method is intended for determining the field batch weights for a concrete mix on a Saturated Surface-Dry, or an Oven Dried basis.

The objective is to correct the field mix design for the percentage of moisture in the aggregates, either on an SSD or OD basis.

Apparatus and Procedure

2. If oven dried basis is desired refer to paragraph (a). For S.S.D. basis, paragraphs (b) and (c) shall be adhered to.

(a) Use the apparatus and follow the procedure for determining the percent total moisture content and the percent surface moisture content of coarse and fine aggregate on a oven dried basis. (AASHTO T 255)

(b) Determine the percent surface moisture of the coarse aggregate on a S.S.D. basis as shown below:

1) Weigh and record the weight of a representative sample of wet coarse aggregate as sampled from the stockpile. The sample should weigh at least 1500 g.

2) Place the sample on a towel or absorbent cloth and dry by rubbing or rolling until the sample still appears damp but no sheen caused by reflected light from free water on the surface of the particles. This is the "saturated surface-dry" state.

3) Weigh and record the weight of the "Saturated Surface Dry" sample. Care shall be taken during the weighing operation to prevent loss of moisture by evaporation.

4) The calculations are as follows:

\[
\text{\% Surface Moisture} = \frac{\text{Wet wt.} - \text{SSD wt.}}{\text{SSD wt.}} \times 100
\]

Example:
Wet wt. = 2056 g.
SSD wt. = 2040 g.
\[
\text{\% Surface Moisture} = \frac{2056 - 2040}{2040} \times 100 = 0.8\%
\]

(c) Determine the percent surface moisture of the fine aggregate (sand) on a S.S.D. basis as shown below:

1) Fill a Chapman flask with water to the 200 ml. mark on the lower neck. The temperature of the water shall be from 65° F to 85° F.

2) Pour a moist sample weighing exactly 500 g. down the neck of the flask.

3) Agitate or roll so as to remove all air bubbles.

4) After all air bubbles have been removed, record the level of the liquid on the scale on the upper neck of the flask.

5) The percent surface moisture shall be calculated as follows:

\[
M = \frac{V - \left( \frac{W_w}{\text{Sp. Gr.}} \right) - V_w}{W_w + V_w - V} \times 100
\]

Where:

\[M = \text{Percent Surface Moisture}\]
\[V = \text{Volume of combined water and sand, ml.}\]
\[\text{Sp. Gr.} = \text{Bulk Specific Gravity of Sand (S.S.D.)}\]
\[W_w = \text{Wet wt. of sand} = 500 \text{ g.}\]
\[V_w = \text{Volume of water} = 200 \text{ ml.}\]

Example:

Volume of water and sand = 399 ml.
Specific Gravity of Sand = 2.653

\[
\frac{500}{2.653} - 200 = 399
\]

\[
M = \frac{399}{500 + 200 - 399} \times 100 = \frac{10.5}{301} = 3.5\%
\]

NOTE: The specific gravity of sand from any one source usually may be considered as constant; however, periodic determinations should be made to verify this. The specific gravity reported with the mix design may be used in the calculation. If no specific gravity has been recorded it may be determined for the sample in accordance with AASHTO T-84, Specific Gravity and absorption of fine aggregate.
Calculations and Example (for Oven-dry Basis)

3. The following serves to illustrate adjustments on an oven-dried (OD) basis:

(a) Moisture contents as determined in 2(a) on an oven dried basis:

Total: Sand = 5.9% water
Coarse = 3.0% water
Surface: Sand = 4.9% water
Coarse = 1.0% water
Asportion: Sand = 1.0%
Coarse = 2.0%

(b) Mix Design (Oven dry basis):
Cement = 94 lb. (one sack)
Sand = 203 lb.
Coarse = 301 lb.
Water = 7.6 gal. (@ 8.33 lbs/gal) = 63 lbs.

(c) Batch weight adjustment:
Cement = 94 lb. (unchanged)
Sand = \( \frac{(O.D. \text{ wt.}) \times (100 + \% \text{ total water})}{100} \)
= \( \frac{(203) \times (105.9)}{100} \)
= 215 lb.
Coarse = \( \frac{(O.D. \text{ wt.}) \times (100 + \% \text{ total water})}{100} \)
= \( \frac{(301) \times (103.0)}{100} \)
= 310 lb.
Surface moisture in sand = \( \frac{\% \text{ surface water} \times (O.D. \text{ wt.})}{100} \)
= \( \frac{(4.9) \times (203)}{100} \)
= 10 lb.
Surface moisture in coarse = \( \frac{\% \text{ surface water} \times (O.D. \text{ wt.})}{100} \)
= \( \frac{(1.0) \times (303)}{100} \)
= 3 lb.
Total surface moisture = 10 + 3 = 13 lb.
Gallons of surface moisture = \( \frac{13 \text{ lb.}}{8.33 \text{ lb. gal.}} \)
= 1.6 gals.

Mix water required = 7.6 — 1.6 = 6.0 gals.

(d) Adjusted batch weights:
Cement = 94 lb.
Sand = 215 lb.
Coarse = 310 lb.
Water = 6.0 gals. (@ 8.33 lbs/gal) = 50 lbs.

Calculations and Example (for S.S.D. basis)

4. The following serves to illustrate adjustments on a SSD basis:

(a) Percent surface moisture as determined in 2(b) and (c) on a SSD basis:
Sand = 3.5% water
Coarse = 0.8% water

(b) Mix Design (Saturated Surface-Dry basis):
Cement = 94 lb. (1 sack)
Sand = 205 lb.
Coarse = 307 lb.
Water = 6.5 gal. (@ 8.33 lbs/gal) = 54 lbs.

(c) Batch weight adjustment:
Cement = 94 lb. (unchanged)
Sand = \( \frac{(SSD \text{ wt.}) \times (100 + \% \text{ water})}{100} \)
= \( \frac{(205) \times (103.5)}{100} \)
= 212 lb.
Coarse = \( \frac{(SSD \text{ wt.}) \times (100 + \% \text{ water})}{100} \)
= \( \frac{(307) \times (100.8)}{100} \)
= 309 lb.
Surface moisture in sand = 212 — 205 = 7 lb.
Surface moisture in coarse = 309 — 307 = 2 lb.
Total Surface Moisture = 7 + 2 = 9 lb.
Gallons of surface moisture = \( \frac{9 \text{ lb.}}{8.33 \text{ lb. gal.}} \)
= 1.1 gal.
Gallons of mixing water required = 6.5 — 1.1
= 5.4 gal.
(d) Adjusted batch weights:

- Cement = 94 lb. (unchanged)
- Sand = 212 lb.
- Coarse = 309 lb.
- Water = 5.4 gal. (@ 8.33 lb./gal/) = 45 lbs.

Report

5. Report adjusted batch weights to the nearest 1 lb., and water content to the nearest 0.1 gal.
TESTING IMPERVIOUS MATERIALS & COMPOUNDS
FOR CURING CONCRETE

(A Modification of AASHO Designation T 155)

REFeree Test

Scope

1. This method is intended for use in determining the efficiency of liquid membrane-forming compounds for curing concrete, as measured by their ability to prevent loss of moisture during the early hardening period.

Apparatus

2. The apparatus shall consist of the following:

(a) Molds. – Molds shall be made of metal, glass, hard rubber, or plastic and shall be watertight, of such construction that distortion is prevented. Interior dimensions shall be: 6 x 12 in. at the top, 5½ x 11½ in. at the bottom, within ± ¼ in., and 2 ± ½ in. depth. A flat rim is required at the top and on all sides and shall be ¼ in. in width.

(b) Curing Cabinet. – A cabinet capable of a temperature of 100°F ± 2°F and relative humidity of 32 ± 2%.

Specific Gravity Determination

3. (a) Fill a specific gravity flask with a well-stirred sample of curing compound.

(b) Gently lower a selected hydrometer into the compound, taking care the hydrometer does not touch the sides of the flask. If the trial hydrometer comes to rest and floats too high, a hydrometer of greater specific gravity is necessary; if the hydrometer sinks below the scale, a lighter hydrometer is required. Continue to change hydrometers until one is found which comes to rest and floats in the sides of the flask. If the trial hydrometer comes to rest on the scale, a lighter hydrometer is required.

(c) Allow the hydrometer to settle to a constant level. (For clear, wax and varnish compounds, this occurs in a very few minutes; but the heavy-pigmented types may require an hour or more). Read and record the specific gravity of the compound.

(d) From Table I which follows, find the quantity in grams of compound to use for the test. For unusual compounds which do not occur on the chart, consult the supervisor.

NOTE: The amount of compound is calculated to represent 150 sq. ft. per gallon coverage. For clear type compounds this works out to 10.0 ml per sample, but other types, and especially the pigmented types, may require more.

Proportioning and Mixing Mortar

4. (a) Proportioning. – The proportions of cement and sand shall be determined by adding dry sand to a cement paste having a water-cement ratio of 0.40 by weight, to produce a flow of 35 ± 3. The flow test shall be made as described in the Standard Method of Test for measuring Mortar-Making Properites of Fine Aggregate (AASHO Designation: T 71).

(b) Mixing. – The mortar shall be mixed at room temperature, preferably 73.4°F ± 3°F (23°C ± 1°C) if possible, and at a relative humidity of 40 to 60 percent. The temperature of the mortar at the time of molding shall be 73.4°F ± 3°F (23°C ± 1°C). The cement and water shall be placed in a nonabsorbent vessel and the cement permitted to absorb water for 1 minute. These materials shall then be mixed with a spoon into a smooth paste. The sand shall be added to the paste and final mixing shall be accomplished by continuous kneading and squeezing with the hands for 2 min. Rubber gloves shall be worn during the mixing operation. A suitable mortar-mixing machine may be used in lieu of hand mixing.

(c) Three specimens shall be molded for testing with a given curing compound.

(d) The mix used to determine the portion of sand to cement to produce the specified consistency shall be discarded and shall not be used for making test specimens.

Molding Specimens

4. (a) Coat the insides of the molds with a very thin coating of a light lubricating oil.

(b) Place a layer of mortar approximately 1 inch thick in the mold, and tamp 50 times with a wooden rod 1-in. square. Fill the mold with mortar and again tamp 50 times.
(c) Immediately after completing the tamping, strike off the specimen level with the top of the mold, with a sawing motion of a wooden screed having a flat surface 1-inch in width. One pass only shall be made in the direction of the long axis of the mold.

**Storage of Specimens**

5. (a) Immediately after molding, weigh each full mold to the nearest 1 g. and place in the curing cabinet. Specimens shall be arranged so as to provide a clear space on all sides of from 2 to 7 inches.

**NOTE:** The movement of conditioned air within the cabinet shall be such that the solvent from curing compounds will readily evaporate. During the first day of drying, sufficient fresh air shall be admitted into the cabinet to eliminate solvent vapors.

**Application of Curing Materials**

7. (a) After approximately 1½ hours, check the specimens for surface water. If surface water is evident, leave the specimens in the cabinet until all surface water has disappeared. Then lightly brush the surface of the specimens with a stiff bristle brush. If surface water appears upon brushing, return the specimen to the curing cabinet. Remove the specimen immediately upon disappearance of the surface water brought to the surface, and brush again. Repeat the process until no surface water appears when the specimen is brushed.

(b) When the proper surface condition has been attained, form a V-shaped groove approximately ¼ in. deep and not more than 1/16 in. wide between the edge of the mortar and the mold, extending all around the specimen. Fill the groove with a sealing compound that will not be affected by the curing material under test. This compound shall not extend more than ⅛ in. from the mold onto the surface of the specimen.

(c) Weigh the specimen to the nearest 1 g. Then with a spray gun (for clear compounds), or with a soft-bristle brush 1 in. in width or a pipette (for pigmented compounds) apply the amount of curing compound determined from Section 3. Take care that the entire surface of the specimen is coated uniformly. Record the weight to the nearest 1 g. Place the specimen in the curing cabinet.

(d) After 3 hours, remove the specimens from the cabinet and weigh to the nearest 1 g. An unusually heavy loss in weight indicates a leaking mold or faulty seal. In such a case, the specimen shall be discarded. Return the specimens to the curing cabinet without delay.
Duration of Curing

8. The specimens shall be cured for 72 hours with occasional daily checks, and the amount of water lost shall be determined by weighing the specimen.

 Corrections for Loss in Weight of Liquid Curing Materials During Test

9. (a) Take an identical quantity of curing compound and spread evenly over a clean, dry, tared metal pan approximately 6 in. x 12 in. x ½ in.

(b) Place the pan in the curing cabinet and dry it to constant weight. Reweight to the nearest 0.1 g. Record the loss in weight of the curing compound as volatile matter.

Calculations

10. (a) Calculate the loss of water and volatiles in the specimen and curing compound by the following:

\[ W = S - S_c \]

Where:

\[ W \]: combined loss of water and volatile g.
\[ S \]: original weight of mold and specimen, with curing compound, g.
\[ S_c \]: weight of cured specimen, plus mold, plus curing compound, g.

(b) Calculate the net loss of water from the specimen by the following:

\[ L = W - W_v \]

Where:

\[ L \]: The net loss of water from specimen, g.
\[ W_v \]: weight of volatile matter, as determined in section 9.(b), g.

(c) Measure the width of the mold in cm. in three areas and use the average with the length to acquire the surface area (A) of the mold.

(d) Calculate the weight of water lost per sq. cm. of surface area by the following:

\[ W_s = L/A \]

Example

11. The following shall serve to illustrate calculations:

- Orig. wt., Mold + Sample + Compound (S) = 5502 g.
- Cured wt. of above (S_c) = 5490 g.
- Loss of water + Volatile (W) = 12 g.
- Volatile in 10 g. Curing Compound (W_v) = 7.2 g.
- Net loss of water (L) = 4.8 g.
- Avg. width of mold = 14.08 cm.
- Avg. length of mold = 29.50 cm.
- Surface Area of mold (A) = 415.4 cm².

Unit loss of water (W_s) = \[ \frac{L}{A} = \frac{4.8}{415.4} = 0.012 \text{ g/cm}² \]

Report

12. Report the following:

(a) Original weight of mold and specimen with curing compound, to nearest 1 g.

(b) Weight of mold, specimen and compound after curing, to nearest 1 g.

(c) Weight of loss of volatiles, to nearest 0.1 g.

(d) Loss of water and volatiles in cured specimen to the nearest 0.1 g.

(e) Net weight of water lost to the nearest 0.1 g.

(f) Average width of mold, to nearest 0.01 cm.

(g) Length of mold to nearest 0.01 cm.

(h) Surface area of mold, to nearest 1 sq. cm.

(i) Unit loss of water, to nearest 0.001 g. per sq. cm.

ALTERNATE METHOD

Scope

1. This method is a procedure for the determination of the non-volatile residue of curing compounds. It is applicable for all types of membrane-forming curing compounds. The percent residue is used in determining the moisture retention efficiency of the curing compound.
Apparatus

2. Apparatus shall consist of the following:

(a) Balance, accurate to 0.1 gram.

(b) Hot Plate, with a minimum of 3 heat variations.

(c) Ointment Tins, 6 oz.

Procedure

3. (a) Thoroughly stir the sample until it is completely free of lumps. Place 5 to 10 g. (weighed to the nearest 0.1 gram) into each of three 6 ounce ointment tins. Place the samples on a medium temperature hot plate (300° to 400° F) and heat until the volatile fraction is completely evaporated. This point can be determined when the formation of bubbles ceases and the residue starts to darken in color.

(b) Remove the cans from the hot plate and allow to cool to room temperature. Weigh each can to the nearest 0.1 gram and average the residue weights for the 3 cans.

Calculation

4. Using the average weight, calculate the percent non-volatiles with the following formula:

\[ \text{Percent} = \frac{B}{A} \times 100 \]

Where:

A = Weight of compound before heating.

B = Weight of residue.

Report

5. Report the result to the nearest 0.1 gram.

Notes

(1) Pigmented compounds should be allowed to settle until the clear liquid can be decanted. The clear liquid is then tested as above.

(2) Emulsion types should be heated slowly to avoid spattering.
MEASURING TEXTURE DEPTH OF
PORTLAND CEMENT CONCRETE
WITH METAL TINE FINISH

A Modification of Louisiana
Dept. of Hwys. Designation TR 229

SCOPE

1. This method describes the procedure for measuring texture depth of fresh or hardened concrete finished with a metal tine.

APPARATUS

2. (a) A tire tread depth measuring gauge with 1/32 of an inch gradations similar to the one shown in Figure 1, or a similar device approved by the Engineer.

(b) Wire brush.

(c) Steel straightedge approximately 1/4" x 1" x 12".

PROCEDURE

3. (a) The depth of texture shall be measured from the original concrete surface. Any projections above the original surface shall be removed by wire brushing or with the steel straightedge prior to taking a measurement on hardened concrete. If measurements are being made on fresh concrete, the depth gauge shall be pressed down until substantially at the level of the original concrete surface. If measurements are being made on hardened concrete such that the depth gauge cannot be pressed down until substantially of the level of the original concrete surface, due to surface irregularities caused by the texturing operation, the height of these surface irregularities shall be measured and subtracted from the measured depth of texture.
(b) With the depth gauge guides in contact with the original concrete surface, the plunger is depressed until contact is made with the bottom of the groove in the concrete. The gauge is then removed from the surface with care being taken to prevent the plunger from being disturbed. The texture depth is then read to the nearest 1/32 of an inch on the calibrated plunger.

(c) The plunger is then re-zeroed and the procedure is repeated until the necessary measurements are completed.

(d) Sufficient measurements shall be taken at random locations along and across the paving width to ensure that the specification requirements are met. In most cases, one test per 12 foot width per 500 foot length should be sufficient. The results should be noted in the inspectors daily diary.
METHOD OF TEST FOR FLOW OF GROUT MIXTURES (FLOW CONE METHOD)

(A Modification of California Test Method 541)

SCOPE

1. (a) This method is intended to be used for determining the flow of grout mixtures as described in this test method.

   (b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

   (a) Flow cone conforming to the dimensions indicated in Figure 1.

   (b) Stop watch accurate to 0.1 second.
(c) Rubber stoppers.

(d) Sample container of four liter minimum capacity [a 152.4 mm x 304.8 mm (6 inch x 12 inch) concrete cylinder mold is adequate].

(e) Supporting ring for flow cone and stand [a 19 liter (5 gallon) bucket may be used], see figure 2.

SAMPLE

3. A representative sample shall be approximately 4 liters of grout.

PRECAUTIONS

4. (a) This test must be performed at a location that is free from vibration.

(b) The cone must be kept clean from cement build-up, especially in or near the orifice and nozzle.

PROCEDURE

5. (a) Determination of Efflux Time

1) Dampen flow cone and allow any excess water to drain. Place the cone in the supporting ring and insert the rubber stopper.

2) Level the cone, then pour the grout from the sample container into the cone until the grout Surface is level with the bottom of the holes in the side of the cone.

3) Remove the stopper and start the stopwatch simultaneously.

4) Stop the stopwatch at the first break or change in the continuous flow of grout from the discharge tube.

5) Dispose of the tested grout sample; rinse the equipment.
(b) Determination of Efflux After Quiescence

1) Fill cone with grout as previously described, using the remainder of the 4 liter sample.

2) Allow grout to rest in cone for 20 minutes ± 15 seconds from the instant the cone is filled. After the 20 minute quiescent period, remove the stopper and determine efflux time as described above.

EXAMPLE

6. Quiescent time \( T_Q \) is the amount of time that a sample of grout remains undisturbed (quiescent) in the flow cone and is expressed in minutes. Efflux time \( T_E \) is the amount of time that a sample of grout requires to run out of the flow cone after the plug is removed, expressed in seconds.

   (a) Efflux time at the pump discharge:

   \[ T_E \geq 11 \text{ seconds (when } T_Q = 0 \text{ minutes)} \]

   (b) Efflux time of grout sample at \( T_Q = 20 \) minutes:

   \[ T_E \text{ (at } T_Q = 20) \geq T_E \text{ (at } T_Q = 0) + 3 \text{ seconds,} \]

   and

   \[ T_E \text{ (at } T_Q = 20) \leq T_E \text{ (at } T_Q = 0) + 8 \text{ seconds} \]

   NOTE: The above mathematical expressions for quiescent time of 20 minutes are expressed as follows: "The efflux time after 20 minutes must be at least 3 seconds greater than the initial efflux time (Quiescent Time = Zero) and not more than 8 seconds greater than the initial efflux time."

REPORT

7. Report the efflux time to the nearest 0.1 seconds for both \( T_Q=0 \) and \( T_Q=20 \).
Grout Flow Cone

FIGURE 1

A = 177.8 mm (7 inches)
B = 50.8 mm (2 inches)
C = 190.5 mm (7-1/2 inches)
D = 12.7 mm (1/2 inch)
E = 38.1 mm (1-1/2 inches)
Grout Flow Test Apparatus

FIGURE 2
COMPRESSION STRENGTH OF CYLINDRICAL CONCRETE SPECIMENS

(A Modification of AASHTO Designation T 22 and ASTM Designation C 39)

1. This test procedure is the same as specified in AASHTO T 22 and ASTM C 39.

2. A blank ADOT “Concrete Test Report” laboratory card is provided in Figure 3 (front of card) and Figure 4 (back of card). An example of the use of the ADOT “Concrete Test Report” laboratory card is given in Figures 5 and 6.
FIGURE 3
**FIGURE 4**

- **Type 1**: Reasonably well-formed cones on both ends, less than 1 in. [25 mm] of cracking through caps.
- **Type 2**: Well-formed cone on one end, vertical cracks running through caps, no well-defined cone on other end.
- **Type 3**: Columnar vertical cracking through both ends, no well-formed cones.
- **Type 4**: Diagonal fracture with no cracking through ends: tap with hammer to distinguish from Type 1.
- **Type 5**: Sidewall fractures at top or bottom (occur commonly with unbonded caps).
- **Type 6**: Similar to Type 5, but end of cylinder is pointed.

---

**AGE:**

Were unbonded caps used? Yes [ ] No [ ] If Yes, what type of unbonded cap was used? Neoprene [ ] Other [ ]

**SPECIMEN IDENTIFICATION**

<table>
<thead>
<tr>
<th>INDICATE TYPE OF FRACTURE PATTERN (✓)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TYPE 1</td>
</tr>
<tr>
<td>A</td>
</tr>
<tr>
<td>B</td>
</tr>
<tr>
<td>C</td>
</tr>
</tbody>
</table>

* If the type of fracture pattern is other than Type 1 to Type 6, sketch and briefly describe the fracture pattern.

Remarks:

---

**AGE:**

Were unbonded caps used? Yes [ ] No [ ] If Yes, what type of unbonded cap was used? Neoprene [ ] Other [ ]

**SPECIMEN IDENTIFICATION**

<table>
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<tr>
<td>B</td>
</tr>
<tr>
<td>C</td>
</tr>
</tbody>
</table>

* If the type of fracture pattern is other than Type 1 to Type 6, sketch and briefly describe the fracture pattern.

Remarks:
**FIGURE 5**

**ARIZONIA DEPARTMENT OF TRANSPORTATION**

**CONCRETE TEST REPORT**

**DATE BATCHED**: 03/07/11

**TRUCK OR BATCH QUANTITY**: 10 CY

**PRODUCT CODE**: 14070

**REQUIRED 28-DAY STRENGTH**: 4000 PSI

---

**AT PLANT** (AT SITE WHEN NO PLANT INSPECTOR)

**DATE SAMPLED**: 03/07/11

**SAMPLE BY**: Joe Sampler

**AT SITE**

**PLACED IN**: -12 FT OF $\text{E}$

**DIR**: NB

**STATION**: 22 + 25

**MAIN ROAD**: -12 FT OF $\text{E}$

**AIR CONTENT**: SPEC

**MEASURED SLUMP/SPREAD**: 0.00 TO 4.50

**MEASURED SLUMP/SPREAD**: 2.75

---

**LAB NUMBER**: 165

**DATE RECEIVED**: 03/08

**TIME REC'D IN LAB**: 13:25

**DATE TESTED**: 04/04

**TIME TESTED**: 05:49

**INDICATE TYPE OF FRACTURE PATTERN ON BACK**

**LAMINAR SIGNATURE AND DATE**: Kim Supervisor 04/04/11

**DOUBLE CHECK**

---

**COMPRESSION STRENGTH IF CYLINDER 165C ELIMINATED - DIFFERS BY MORE THAN 10% FROM THE AVERAGE OF THE 3 CYLINDERS**

**LABORATORY SIGNATURE AND DATE**:

**SUPERVISOR SIGNATURE AND DATE**:

---

**SEE BACK ALSO**
FIGURE 6

AGE: 28 Day

Were unbonded caps used? Yes ☑ No ☐ If Yes, what type of unbonded cap was used? Neoprene ☑ Other ☐

<table>
<thead>
<tr>
<th>SPECIMEN IDENTIFICATION</th>
<th>TYPE 1</th>
<th>TYPE 2</th>
<th>TYPE 3</th>
<th>TYPE 4</th>
<th>TYPE 5</th>
<th>TYPE 6</th>
<th>OTHER*</th>
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<tbody>
<tr>
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<td></td>
<td></td>
<td>✗</td>
<td></td>
<td></td>
<td>✗</td>
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</tr>
<tr>
<td>B</td>
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</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>✗</td>
</tr>
</tbody>
</table>

* If the type of fracture pattern is other than Type 1 to Type 6, sketch and briefly describe the fracture pattern.

Remarks: ____________________________________________

AGE: __________________

Were unbonded caps used? Yes ☐ No ☑ If Yes, what type of unbonded cap was used? Neoprene ☑ Other ☐

<table>
<thead>
<tr>
<th>SPECIMEN IDENTIFICATION</th>
<th>TYPE 1</th>
<th>TYPE 2</th>
<th>TYPE 3</th>
<th>TYPE 4</th>
<th>TYPE 5</th>
<th>TYPE 6</th>
<th>OTHER*</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
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<td>B</td>
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<tr>
<td>C</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* If the type of fracture pattern is other than Type 1 to Type 6, sketch and briefly describe the fracture pattern.

Remarks: ____________________________________________
PRECAST MORTAR BLOCKS TEST

(An Arizona Test Method)

SCOPE

1. (a) This test method covers the accepted procedure for determining the compressive strength of precast mortar spacer blocks which are used to maintain the proper clearance on reinforcing steel bars. Precast mortar spacer blocks are prepared utilizing portland cement, aggregates, water, and admixtures.

(b) The compressive strength of the precast mortar spacer blocks shall be equal to or greater than the required twenty-eight day strength of the concrete that will encase the spacer blocks.

(c) The normal compressive strength of the precast mortar spacer blocks shall be in a range between 15 to 40 MPa (2000 to 6000 psi).

(d) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(e) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(f) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.
APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

   (a) A testing machine that conforms to the description given in AASHTO T 22, Section 5.

   (b) A saw having a diamond or silicon-carbide cutting edge capable of cutting specimens which conform to the prescribed dimensions, without excessive heating or shock.

   (c) Alignment devices and melting pots for sulfur mortars, conforming to the applicable sections of AASHTO T 231.

PREPARATION

3. (a) The length of the specimen, when capped, shall be as nearly as practical twice the diameter or least width. A specimen having a length which is less than the diameter or least width after capping shall not be tested.

   (b) The ends of the specimens to be tested in compression shall be essentially smooth, perpendicular to the longitudinal axis, and of the same cross-sectional area as the body of the specimen. If necessary, saw or tool the ends of the specimens until the following requirements are met:

       1) Projections, if any, shall not extend more than 5 mm (0.2 inch) above the end surfaces.

       2) The end surfaces shall not depart from the perpendicular to the longitudinal axis by more than 5 degrees.

   (c) Before making the compression test, cap the ends of the specimens in conformance with the procedure prescribed in the applicable section of AASHTO T 231. The capped surfaces of the specimens shall conform to the planeness requirements of AASHTO T 231.
(d) Prior to testing, measure the length of the capped specimen to the nearest 2.5 mm (0.1 inch) and use this length to compute the ratio of length to diameter or least width. Determine the dimension of the diameter or each width by averaging two measurements taken at the 1/3 points of the length of the specimen. Measure the diameter or each width and determine the average dimension to at least the nearest 2.5 mm (0.1 inch). If the ratio of length to diameter or least width of the specimen exceeds 2.10, the specimen shall be reduced in length to have a ratio of 1.94 to 2.10.

TEST PROCEDURE

4. (a) Test the specimens in accordance with the applicable provisions of AASHTO T 22.

Note: Short specimens fail at a greater load because the steel platens of the testing machine restrain lateral expansion throughout the specimen more effectively. The effect of end restraint is conventionally assumed to be negligible for a standard specimen with a ratio of length to diameter or least width of two. Short specimens are clearly defined in both AASHTO T 22 and T 24.

CALCULATIONS

5. (a) Calculate the compressive strength of each specimen using the computed cross-sectional area based on the average area of the specimen.

(b) Specimens which have a ratio of length to diameter or least width of 1.94 to 2.10 require no correction.

(c) If the ratio of length to diameter or least width of the specimen is less than 1.94, apply correction factors shown in the following table:

<table>
<thead>
<tr>
<th>Ratio of Length to Diameter or Least Width</th>
<th>Strength Correction Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.75</td>
<td>0.98</td>
</tr>
<tr>
<td>1.50</td>
<td>0.96</td>
</tr>
<tr>
<td>1.25</td>
<td>0.93</td>
</tr>
<tr>
<td>1.00</td>
<td>0.87</td>
</tr>
</tbody>
</table>

Values not given in the table shall be determined by interpolation.
REPORT

6. (a) The compressive strength of the tested specimen at ultimate load shall be reported to the nearest 50 kPa (10 psi), after correction for length to diameter or least width ratio when appropriate.
OBTAINING AND TESTING DRILLED CORES
AND SAWED BEAMS OF CONCRETE

(A Modification of AASHTO Designation T 24)

1. This test procedure is the same as specified in AASHTO T 24, with the following exceptions:

   (a) Section 7.5 of AASHTO T 24 is revised to read:

   7.5 Measurement – Prior to testing, measure the length of the capped specimen to the nearest 0.1 inch (2.5 mm) and use this length to compute the length-diameter ratio. Each test specimen shall be measured to determine its diameter. The diameter shall be the average of two measurements, taken using a caliper, at right angles to each other at about mid-height of the specimen. The individual diameter measurements shall be determined to the nearest 0.01 inch (0.25 mm). Determine and record the average of the two diameter measurements to the nearest 0.01 inch (0.25 mm). That value shall be the diameter used to calculate the cross-sectional area of the test specimen. Do not test the core if the difference between the largest diameter measurement and the smallest diameter measurement exceeds 5% of their average.

   (b) Section 7.6 of AASHTO T 24 is revised to read:

   7.6 Testing – Test the specimens in accordance with Arizona Test Method 314, except unbonded caps (neoprene or other elastomeric materials) with metal retainers shall not be used. Unless otherwise specified, the specimens shall be tested within seven days after coring.

   (c) Section 7.8 of AASHTO T 24 is revised to read:

   7.8 Report – Report the results as required by Arizona Test Method 314, with the addition of the following information:
(d) Section 7.8.1 of AASHTO T24 is revised to read:

7.8.1 The location where the core was obtained. The length of the core
as drilled to the nearest 0.2 inch (5 mm),

(e) Section 7.8.2 of AASHTO T 24 is revised to read:

7.8.2 The length of the test specimen before and after capping or end
grinding to the nearest 0.1 inch (2.5 mm), and the average
diameter of the core to the nearest 0.01 inch (0.25 mm),

(f) Section 7.8.3 of AASHTO T 24 is revised to read:

7.8.3 The compressive strength of the test specimen to the nearest 10 psi (50 kPa), after correction for the length-diameter ratio when required. In addition, if the average compressive strength is determined for multiple specimens, the average compressive strength to the nearest 10 psi (50 kPa),
ESTIMATING THE DEVELOPMENT OF CONCRETE STRENGTH BY THE MATURITY METHOD

(An Arizona Method)

1. SCOPE

1.1 This test method provides a procedure for estimating the development of concrete strength by means of the maturity method.

1.2 This procedure is normally used, when desired, for Class S and Class P concrete; however, it may be used for other classes of concrete.

1.3 The concept of the maturity method is based on the combined effects of concrete age and temperature during the hydration process versus the rate of strength gain for a specific mix.

1.4 This method requires the establishment of a relationship curve between compressive strength and calculated maturity index for a specific concrete mixture (mix design) prior to placement of the mixture in the field.

1.5 This procedure may not be used to estimate the strength of mass concrete unless the strength-maturity relationship is developed at elevated temperatures. Mass concrete is defined as concrete which has a least dimension of three feet or more.

1.6 The maturity method shall not substitute for compressive strength acceptance testing (28-day test cylinder breaks).

1.7 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.8 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.
2. REFERENCED DOCUMENTS

2.1 AASHTO Documents

- AASHTO T325 Estimating the Strength of Concrete in Transportation Construction by Maturity Tests
- AASHTO T23 Making and Curing Concrete Test Specimens in the Field
- AASHTO R39 Making and Curing Concrete Test Specimens in the Laboratory

2.2 ASTM Documents

- ASTM C 1074 Estimating Concrete Strength by the Maturity Method

2.3 Arizona Documents

- ARIZ 314 Compressive Strength of Cylindrical Concrete Specimens
- PPD No. 15 Submittal and Approval of Portland Cement Concrete Mix Designs

3. TERMINOLOGY

3.1 Maturity - The extent of development of concrete properties that are dependent on the hydration of cementitious materials.

3.2 Maturity Function - A mathematical expression that converts the temperature history of concrete to an index, which indicates its maturity.

3.3 Maturity Index - An index, calculated by using a maturity function, which can be used as an indicator of strength development in concrete.

3.4 Strength-Maturity Relationship - An empirical relationship between concrete strength and its maturity index, determined by comparing the strength of concrete cylinders, made from a specific concrete mix, to their maturity index at the time of strength testing.
4. SIGNIFICANCE AND USE

4.1 The estimated strength of in-place concrete determined by this procedure provides guidance useful in making decisions concerning:

4.1.1 Removal of formwork and reshoring.

4.1.2 Post-tensioning.

4.1.3 Opening of the roadways to traffic.

4.1.4 Initiation of strength tests, such as coring and pullout tests, on the in-place concrete.

4.2 The most critical limitations of the procedures presented are:

4.2.1 Concrete must be cured in a condition that supports the reaction of cementitious materials.

4.2.2 Batching or placement errors that are not detected.

4.2.3 Curing errors other than those that affect temperature that are not detected.

4.2.4 The actual strength of the concrete that is not measured.

4.3 This method provides technical personnel with a coordinated procedure for:

4.3.1 Developing a strength-maturity relationship for the approved concrete mix design in the laboratory.

4.3.2 Determining the temperature history of the in-place concrete.

4.3.3 Determining the maturity index of the in-place concrete.

4.3.4 Using the strength-maturity relationship and the maturity index to estimate the strength of the in-place concrete.

5. MATURITY METHOD

5.1 The temperature-time factor (Nurse-Saul) method shall be used to determine the maturity index.
5.2 Maturity digital data loggers shall be set at a datum factor of 0 °C.

5.3 The strength-maturity relationship shall be plotted using a linear chart or a logarithmic chart. In support of using either a linear chart or a logarithmic chart, a printout of the strength-maturity relationship test data shall be used.

6. APPARATUS

6.1 A maturity meter capable of reading a maturity digital data logger that has a secure method of collecting un-interruptible and un-alterable data, subsequent to the concrete pour and initial concrete set, for a minimum of 28 days. The system shall be accurate to at least ± 1 °C.

6.2 Maturity digital data loggers shall contain a real time clock and be capable of recording the current temperature and temperature history in real time with an associated time stamp.

6.3 Documentation shall be provided by the manufacturer that certifies the accuracy of the maturity meter and digital data loggers.

7. PROCEDURE TO DEVELOP STRENGTH-MATURITY RELATIONSHIP

7.1 A strength-maturity relationship curve shall be developed for a specific concrete mix prior to using this method on the project.

7.2 As described in Subsection 7.3 below, the strength-maturity relationship curve is developed utilizing compressive strength cylinders which have been fabricated, cured, and tested from a trial batch as described in ADOT Materials Policy and Procedure Directive No. 15, “Submittal and Approval of Portland Cement Concrete Mix Designs”. The concrete mixture shall meet the specification (design mix) requirements in order to determine the strength-maturity relationship curve. The concrete mixture shall be at or above the target air content.

7.3 The compressive strength testing for the development of the strength-maturity relationship curve shall be performed as follows:
7.3.1 This procedure requires the initial casting of at least 17 compressive strength test cylinders. The test cylinders shall be fabricated in accordance with the requirements of AASHTO T23.

7.3.2 Two cylinders will be outfitted with maturity digital data loggers. Readings are taken from the data loggers in each of these cylinders. The readings are averaged, and the results are correlated with the compressive strength test results to establish the maturity curve.

7.3.3 A minimum of fifteen cylinders will be tested for compressive strength as described below.

7.3.3.1 Compressive strength tests shall be performed on three cylinders, at a minimum of five different ages encompassing the age at which the desired strength is needed. For example, compressive strength tests may be performed at 1, 2, 3, 5 and 7 days for Class S concrete or 12, 24, 36, 48 and 72 hours for Class P concrete. More cylinders may be fabricated, cured, and tested at additional ages if desired. Compressive strength testing shall be performed in accordance with the requirements of Arizona Test Method 314.

Note: Tests should be scheduled to result in approximately equal increments of strength gain between ages. Additionally, compressive strength tests should bracket the estimated target strength.

7.3.3.2 The compressive strength of each of the three test cylinders at each age and their average compressive strength shall be determined. However, if the compressive strength of any one of the three test cylinders differs by more than 10 percent from the average of the three, its result shall be discarded and the compressive strength shall be the average of the remaining two cylinders. Should the individual compressive strength of any two of the three test cylinders differ by more than 10 percent from the average of the three, the results of both will be discarded and the compressive strength shall be the strength of the remaining cylinder.

8. PLACEMENT OF DIGITAL DATA LOGGERS IN THE FIELD

8.1 Maturity digital data loggers shall be placed as follows, either prior to concrete placement (with maturity digital data loggers properly secured in their proper location) or into the fresh concrete following the placement of concrete.
### Class of Concrete

<table>
<thead>
<tr>
<th>Class of Concrete</th>
<th>Frequency of Logger Placement</th>
<th>Location of Logger Placement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class P (PCCP)</td>
<td>Every 100 cubic yards</td>
<td>Mid-depth of pavement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>≥ 12 inches from edge of pavement</td>
</tr>
<tr>
<td>Class S (Structural)</td>
<td>Every 50 cubic yards*</td>
<td>Critical locations per exposure and/or structural requirements</td>
</tr>
<tr>
<td>Other than Class S or Class P</td>
<td>Every 50 cubic yards*</td>
<td>Critical locations per exposure and/or structural requirements</td>
</tr>
</tbody>
</table>

* At least two additional maturity data loggers should be available for placement as needed.

### 8.2

Activate the maturity digital data loggers as soon as practicable after concrete placement, but not before the maturity digital data loggers are fully encapsulated in concrete.

### 9.

**FIELD VALIDATION OF STRENGTH-MATURITY RELATIONSHIP**

**9.1** The maturity method does not account for variations in strength due to proportioning, mixing, placing, and moisture conditions of the field-placed concrete. Proper curing of the concrete and minimizing moisture loss is critical in achieving reliable strength-maturity relationship results.

**9.2** During the initial use of this method for the concrete mixture for which the strength-maturity relationship has been developed, a field validation of the strength-maturity relationship shall be performed. A minimum of three compressive strength test cylinders shall be fabricated, cured, handled, protected, and transported in accordance with the requirements of AASHTO T23. In addition, a minimum of one test cylinder shall be fabricated and cured in
the same manner as the three compressive strength test cylinders, and at least two maturity digital data loggers shall be embedded in the additional test cylinders, as close to the center (±1/2 inch) of the cylinders as possible. The data logger readings are used in conjunction with the average compressive strength for validation of the strength-maturity relationship.

9.3 The three compressive strength cylinders shall be tested in accordance with the requirements of Arizona Test Method 314 at approximately the estimated temperature-time factor target. The average compressive strength of the three test cylinders shall be determined. However, if the compressive strength of any one of the three test cylinders differs by more than 10 percent from the average of the three, its result shall be discarded and the compressive strength shall be the average of the remaining two cylinders. Should the individual compressive strength of any two of the three test cylinders differ by more than 10 percent from the average of the three, the results of both will be discarded and the compressive strength shall be the strength of the remaining cylinder.

9.4 Compare the data logger readings and the average compressive strength of the field validation test cylinders to the original strength-maturity relationship curve at the estimated temperature-time factor target. The average compressive strength of the field validation cylinders is used only for comparison to the original strength-maturity relationship curve.

9.4.1 If the average compressive strength of the validation test cylinders is equal to or differs by less than ±10 percent of the original strength-maturity relationship which has been developed for the concrete mixture, the development of a new strength-maturity relationship curve may not be required.

9.4.2 If the average compressive strength of the validation test cylinders differs by greater than ±10 percent of the original strength-maturity relationship, the development of a new strength-maturity relationship curve is required.

9.5 During placement of the concrete mixture, field validations of the strength-maturity relationship curve shall be performed in accordance with the requirements of Subsections 9.2 through 9.4. The minimum frequency of field validations will be as follows:

9.5.1 For Class P concrete: every 10 days of concrete placement.

9.5.2 For Class S concrete: every 300 cubic yards of concrete placement.

9.5.3 For other than Class P or Class S: every 300 cubic yards of concrete placement.
10. REPORT FOR STRENGTH MATURITY RELATIONSHIP

10.1 Project number

10.2 Concrete Product Code and the source of each material (aggregate, cement, supplementary cementitious material, and admixtures).

10.3 Date(s) and times(s) that compressive strength testing of the cylinders is performed.

10.4 The air content, slump, water content, and the water/cementitious material ratio for each batch of concrete tested.

10.5 The amount and type of admixtures used in the concrete mixture.

10.6 The strength of each test cylinder and the average strength of the test cylinders at each test age.

10.7 The maturity index for each instrumented test cylinder and the average maturity index for the instrumented test cylinders at each test age.

10.8 A graph of the plotted points, on either a linear chart or a logarithmic chart, for the average compressive strength at each age versus the average maturity index at each age, and the best-fit curve. An example of such a linear graph is shown in Figure 1. An example of a logarithmic graph is shown in Figure 2. To be acceptable, the plotted points shall produce a correlation coefficient (R2) of at least 0.90.

10.8.1 In support of providing either a linear chart or a logarithmic chart, a printout of the strength-maturity relationship test data used shall be provided. An example of such a printout is shown in Figure 3.
Example Strength-Maturity Relationship Curve
Linear Scale

$R^2 = 0.980$

FIGURE 1
Example Strength-Maturity Relationship Curve
(Logarithmic Scale)

FIGURE 2
### Example Printout of Strength-Maturity Relationship Test Data

<table>
<thead>
<tr>
<th>Temperature-Time Factor, °C-Hours</th>
<th>Compressive Strength, psi</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>2,300</td>
</tr>
<tr>
<td>1,100</td>
<td>3,500</td>
</tr>
<tr>
<td>2,300</td>
<td>4,500</td>
</tr>
<tr>
<td>4,000</td>
<td>5,000</td>
</tr>
<tr>
<td>7,900</td>
<td>5,600</td>
</tr>
<tr>
<td>16,000</td>
<td>6,300</td>
</tr>
</tbody>
</table>

**FIGURE 3**
SERIES 400
BITUMINOUS MIXTURES
MOISTURE CONTENT
OF BITUMINOUS MIXTURES

(An Arizona Method)

1. SCOPE

1.1 This method is used to determine the percent moisture in bituminous mixtures. The option of using a conventional oven or a microwave oven is provided. In case of dispute, the conventional oven shall be utilized.

1.2 This test method may involve hazardous material, operations, and equipment. This test method does not purport to address all of the safety problems associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Suitable sample containers for use in testing with the conventional oven or microwave oven.

2.3 Oven - A thermostatically controlled oven capable of maintaining a temperature of 290 ± 10 °F; or, a microwave oven capable of variable heat intensity settings.

2.4 A balance or scale capable of measuring the maximum weight to be determined, accurate to at least 0.1 gram.

3. PROCEDURE (CONVENTIONAL OVEN)

3.1 Obtain a representative 1000 ± 50 gram sample in accordance with ARIZ 416.

3.2 Record the tare weight of the container to the nearest 0.1 gram.
3.3 Place sample in the container and weigh. Determine and record the wet weight of sample to the nearest 0.1 gram as "f".

3.4 Place container and sample in a 290 ± 10 °F oven and initially dry for a minimum of 1 hour. Weigh the container and sample. Record the weight to the nearest 0.1 gram.

3.5 Continue drying and weighing until a constant weight is obtained; being the weight at which further drying does not alter the weight more than 0.1 gram at intervals of a minimum of 30 minutes.

3.6 After constant weight is obtained, cover sample and allow to cool 30 ± 10 minutes at room temperature. Weigh and determine and record the dry weight of sample to the nearest 0.1 gram as "g".

3.7 Proceed to section 5 for moisture content calculation.

4. PROCEDURE (MICROWAVE OVEN)

4.1 Obtain a representative 1000 ± 50 gram sample in accordance with ARIZ 416.

4.2 Record tare weight of the container to the nearest 0.1 gram.

4.3 Place sample in the container and weigh. Determine and record the wet weight of sample to the nearest 0.1 gram as "f".

4.4 Dry sample until a constant weight is obtained. The sample is considered to be at constant weight when further drying causes, or would cause, a difference in weight of not more than 0.1 gram. The sample shall be heated in such a manner that controls the intensity of heat generated to prevent splattering, aggregate breakage, and asphalt being "burned off". The method used with a microwave oven shall give results similar to those achieved with a conventional oven.

4.5 After constant weight is obtained, cover sample and allow to cool 30 ± 10 minutes at room temperature. Weigh and determine and record the dry weight of sample to the nearest 0.1 gram as "g".
5. **CALCULATION**

5.1 Calculate the percent moisture, "h", and record to the nearest 0.01% as shown below.

\[ h = \frac{f - g}{f} \times 100 \]

Where:
- \( h \) = Percent Moisture
- \( f \) = Wet weight of sample
- \( g \) = Dry weight of sample
1. SCOPE

1.1 This method covers the procedure for compaction and testing bituminous mixtures utilizing four inch Marshall apparatus.

1.2 This procedure is used for bituminous mixtures with a mix design gradation target of at least 85% passing the 3/4 inch sieve.

1.3 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.4 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Compaction Mold Assembly 4 inch diameter cylindrical mold, baseplate, and extension collar constructed of steel and conforming to the requirements of Figure 1. (Three compaction mold assemblies are normally utilized.)

2.3 Specimen Extruding Device - Extrusion jack or press for extruding specimens from molds.
2.4 Compaction Hammer:

2.4.1 The compaction hammer shall either be a mechanical or hand compaction hammer having a flat, circular tamping face with a nominal diameter of (3-7/8 inches), and a (10 ± 1/4 pound) sliding weight with a free fall of (18 ± 1/2 inches).

2.4.2 Compaction hammers must be monitored through the ADOT proficiency sample program. To be qualified, compaction hammers must produce specimens with an average density of no greater than ± 1.0 lb./cu. ft. from the average bulk density for the most recent set of proficiency samples. If two samples are required for the proficiency samples, both of the bulk density sets must meet the ± 1.0 lb./cu. ft. criteria, if not, the hammer is not qualified.

2.4.3 As an alternate to qualifying a compaction hammer through the proficiency sample program, a compaction hammer may be qualified by correlating with a hammer which has been approved through comparison with proficiency sample results. When qualified in this manner, results must be no greater than ± 0.5 lb./cu. ft.

2.4.4 Hammers which have had adjustments or repairs made to them after being qualified, must be requalified by correlating with another qualified hammer and yield results within ± 0.5 lb./cu. ft.

Note: Marshall compaction equipment can go out of calibration at any time, and each laboratory is encouraged to establish a method of ensuring that their equipment remains in calibration. Alternate methods that can be used include regular comparisons with other approved hammers or compaction of samples which have a known density.

2.4.5 Hammers which do not meet the above requirements may be adjusted by modifying the weight, or the height of fall, within the given criteria; by adjusting the number of blows a maximum of ± 10 from the specified 75 blows; or by a combination of adjustments to weight, height of fall, or number of blows.

2.4.6 Should a compacton pedestal be moved or replaced, the compaction hammer(s) shall be requalified.
2.5 Compaction pedestal - The compaction pedestal shall consist of a 8" x 8" x 18" wooden post capped with a 12" x 12" x 1" steel plate. The steel cap shall be firmly fastened to the post. The wooden post shall have a dry weight of 42 to 48 lbs./cu. ft. and shall rest squarely on, and be firmly secured to, a solid concrete slab. The pedestal assembly shall be installed so that the post is plumb and the cap is level.

2.6 Specimen Mold Holder - Mounted on the compaction pedestal so as to center the compaction mold over the center of the post. It shall hold the compaction mold, collar, and base plate securely in position during compaction of specimen.

2.7 Oven for heating bituminous mixtures and specimen mold assemblies at required temperature.

2.8 Hot plate for heating compaction hammer, spoon and spatula.

2.9 A flat spatula with blade approximately 1 inch wide and at least 6 inches long, stiff enough to penetrate the entire bituminous mixture.

2.10 Calibrated/verified thermometers, for determining temperatures of bituminous mixtures, with a range of 50 to 400 °F and increments of not greater than 5 °F. For digital thermometers, increments shall not be greater than 1 °F.

2.11 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least one gram.

2.12 If Marshall stability and flow are to be determined, the following additional apparatus is required:

2.12.1 Breaking Head and Water Bath, conforming to the requirements specified in AASHTO T 245.

2.12.2 Marshall stability and flow testing apparatus, with operating instruction manual. The apparatus shall be capable of applying a load with a constant rate of travel of 2.0 ± 0.1 inches per minute.

2.12.3 Height gauge capable of measuring the height of specimens to the nearest 0.001 inch.
3. PROCEDURE

3.1 Obtain three representative samples for Marshall specimens, as described in Arizona Test Method 416. If the Combined Aggregate Bulk (O.D.) Specific Gravity is known, the weight calculated by the following equation (± 50 grams) will normally give specimens which meet the height requirement of 2.300 to 2.700 inches. (1150 ± 50 grams is generally adequate.)

\[
\text{Weight of each Sample} = \frac{\text{Combined Aggregate Bulk (O.D.) Specific Gravity}}{2.520} \times 1150
\]

3.2 Before placing the mixture in the mold, the mixture and a mold assembly (baseplate, mold, and collar) shall be at approximately 290 °F. The face of the compaction hammer shall be thoroughly cleaned and heated on a hot plate set at approximately 290 °F. The temperature of the laboratory during compaction of the specimens shall be between 68 and 86 °F.

3.3 Place a 4-inch paper disc in the bottom of the mold before the mixture is introduced. Place the entire batch in the mold in one lift. Care should be taken to avoid segregation of material in the mold. Spade the mixture vigorously, penetrating the entire mix, with the heated spatula 15 times around the perimeter and 10 times at random into the mixture. Smooth the surface of the mix to a slightly rounded shape.

3.4 The compaction temperature shall be the laboratory compaction temperature shown on the mix design.

3.5 If necessary, the mixture and mold assembly shall be returned to an oven at the required temperature for the minimum time necessary to achieve the laboratory compaction temperature ± 5 °F; however, in no case shall the mixture be reheated longer than 60 minutes.

3.6 Place a 4-inch paper disc on top of material, place the mold assembly on the compaction pedestal in the mold holder, and apply 75 blows [or adjusted number, as determined in Subsection 2.4] with the compaction hammer. When a hand hammer is utilized, the operator shall hold the handle by one hand so that the axis of the compaction hammer is as nearly perpendicular to the base of the mold assembly as possible while compaction is accomplished. Care shall be taken not to add body weight to the hammer by leaning or pressing down on the hammer. When using a hand hammer, no mechanical device of any kind is
to be used to restrict movement of the handle during compaction. Compaction shall be performed at a minimum rate of 40 blows per minute. The compaction hammer shall apply only one blow with each fall, that is, there shall not be a rebound impact. Remove the base plate and collar, and reverse and reassemble the mold. Apply 75 (or adjusted number) compaction blows to the face of the reversed specimen.

3.7 Remove collar, baseplate, and paper discs, and allow specimen to cool. Cooling may be accomplished at room temperature, in a 77 °F air bath, or if more rapid cooling is desired the mold and specimen may be placed in front of a fan until cool.

3.8 Extrude the specimen from the mold. Care shall be taken in extruding the specimen from the mold, so as not to develop tensile stresses in the specimen or tear the sides of the specimen.

4. SPECIMEN TESTING

4.1 If Marshall stability and flow are to be determined, measure height of specimens to the nearest 0.001 inch. Prior to measurement of height, excess material shall be brushed from the edges of the specimens. Compacted specimens shall be 2.300 to 2.700 inches in height. If this criteria is not met, the entire set of specimens shall be discarded and a new set prepared after necessary adjustments in sample weight have been made.

4.2 Determine the specific gravity of the specimens in accordance with Arizona Test Method 415, Method A. (Assume specimen is at constant weight after cooling.)

4.3 Determine the bulk density of each of the specimens, by multiplying the respective specific gravity by 62.3 lbs./cu. ft. Record the individual bulk densities to the nearest 0.1 lb./cu. ft. The densities of the three specimens shall not differ by more than 2.5 lbs./cu. ft. for 1/2", 3/4", or recycle mixes; and 3.0 lbs./cu. ft. for Base mixes. If this density requirement is not met, the entire set of specimens shall be discarded and a new set of specimens prepared.

4.4 Determine the average specific gravity of the specimens and record to the nearest 0.001. Calculate the average bulk density of the specimens, by multiplying the average specific gravity by 62.3 lbs./cu. ft. Record the average bulk density to the nearest 0.1 lb./cu. ft.
4.5 If the stability and flow are to be determined, the steps in Subsections 4.6 through 4.11 below are followed, utilizing the apparatus in accordance with the operating instructions for that apparatus.

4.6 Bring the specimens to $140 \pm 2 \, ^\circ$F by immersing in the water bath 30 to 40 minutes. Prior to testing, it shall be assured that the inside of the test heads are clean, and that the guide rods are clean and lubricated so that the upper test head slides freely over them.

4.7 The breaking head temperature shall be maintained between 70 to 100 $^\circ$F, using a water bath when required. Remove the specimen from the water bath, quickly towel dry specimen and place in the lower segment of the breaking head. Place the upper segment of the breaking head on the specimen, and place the complete assembly in position on the testing machine.

4.8 Apply the load to the specimen with a constant rate of $2.0 \pm 0.1$ inches per minute until the maximum load is reached and the load decreases. The maximum load is defined as the last point in the load/time curve before the load decreases. The elapsed time for the test from removal of the test specimen from water bath to maximum load determination shall not exceed 30 seconds.

4.9 Record the stability of each specimen to the nearest 10 pounds force, and the flow to the nearest 0.01 inch.

4.10 Correct the stability obtained for each specimen, for the height of the specimen, by the table in Figure 2. Record the corrected stability to the nearest 10 pounds force.

4.11 Determine and record the average corrected stability to the nearest 10 pounds force, and the average flow to the nearest 0.01 inch.
Mold Symmetrical about this centerline

**LEGEND**

A: 4-3/4 in.  
B: 9/16 in.  
C: 3-63/64 in.  
D: 1/8 to 9/32 in.  
E: 4-1/2 in.  
F: 3/16 in.  
G: 4-1/8 in.  
H: 3-7/16 in.  
I: 1/4 in.  
J: 2-3/4 in.  
K: 4-5/32 in.  
L: 4-9/32 in.  
M: 4-19/64 in.  
N: 9/32 in.  
O: 20 degrees  
P: 3.990 to 4.010 in.

All dimensions are nominal, except where tolerances are indicated.

**Four Inch Compaction Mold, Extension Collar, and Baseplate**

FIGURE 1
### STABILITY CORRELATION RATIOS*

For 4 inch Diameter Specimens

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<th>Height of Specimen (Inches)</th>
<th>Correlation Ratio</th>
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<td>2.698 - 2.700</td>
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</tbody>
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* The measured stability of a specimen multiplied by the correlation ratio for the height of the specimen equals the corrected stability for a 2-1/2 inch specimen.
DETERMINATION OF BITUMINOUS DISTRIBUTOR TRUCK TRANSVERSE SPREAD RATE

(A Modification of California Test 339)

1. SCOPE

1.1 This procedure describes the method for determining the transverse spread rate, in gallons per square yard, of bituminous distributor trucks. This procedure is applicable for all bituminous materials other than asphalt-rubber and cutback asphalts.

Note: Three Alternate Sampling Procedures are outlined in this test method.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 Balance - A balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.1 gram.

2.1.2 Oven - An oven capable of maintaining a temperature of 140 ± 5 °F.

2.1.3 Metal Sheets (20 gauge galvanized) - 7-7/8 inches in width and with a length to accommodate a sixteen foot spray bar. Any combination of lengths of metal plates may be used. (For example, three plates each 68
inches long; or two plates each 84 inches long and one plate 36 inches long.)

2.1.4 Testing Machine (when Alternate Sampling Procedure #1 is used.) - A device with a sliding platform which holds the test plates as they are moved under and through the bituminous material while it is being sprayed. See Figure 4 for an illustration of a typical "sliding platform" device.

2.1.5 When Alternate Sampling Procedures #2 or #3 are used, a provision to raise the plates and pads off the surface must be provided. (For example, 2” x 4” or 1” x 6” boards in any combination of lengths for a total length of 17 feet.)

2.1.6 Stapler - For fastening cotton pads to paper strips.

3. MATERIALS

3.1 100% cotton gauze pads - 4” x 8”, 12 ply.

3.2 5” x 12” strips cut from heavy wrapping paper.

3.3 Masking tape, minimum 1” width.

3.4 Scotch tape, 3/4” width

4. PREPARATION OF TEST PADS AND TEST PLATES

4.1 Prepare each test pad by attaching two cotton pads (laid on top of each other) to each paper strip with a staple at each corner of the cotton pads. The cotton pads are attached to the paper strips as shown in Figure 1.

4.2 Place a strip of scotch tape along each end of the paper strip as shown in Figure 1. This will allow the masking tape to be easily removed from the paper strip during the removal of the test pad from the test plate.

4.3 Place an identification number on the paper back of each test pad. To facilitate the assembly of the test plates at the test site, it is helpful to also place the identification number in the middle of either end on the front of each test pad. (Fifty-one consecutively numbered test pads are required to fabricate the three test plates for each test.)
4.4 Weigh each test pad and record its tare weight to the nearest 0.1 gram on the test record shown in Figure 7.

4.5 The test pads are attached to the metal sheet in numerical order so that the cotton pads are snug against each other by overlapping the adjacent strips 1 inch. Figure 2 provides an illustration of test pads attached to the test plate. Twenty-one test pads are required to fabricate an 84-inch test plate. Nine test pads are required to fabricate a 36-inch test plate. Each end of the test pad is folded over to the backside of the test plate and secured with masking tape.
Figure 2 – Test Pads Attached To Test Plate
4.6 Stack the assembled test plates on one another, with the corresponding test record containing the test pad tare weights. Wrap the bundle in heavy wrapping paper, as shown in Figure 3, for protection and to keep them clean while being stored or transported to the test site.

![Wrapped Test Plates](image)

**Figure 3 – Wrapped Test Plates**

5. **ALTERNATE SAMPLING PROCEDURE #1**

5.1 Alternate Sampling Procedure #1 utilizes the sliding platform testing machine described in Subsection 2.1.4. In this procedure, the distributor truck remains stationary while the test is being performed. Side shields shall not be used when testing is performed under this procedure.
5.1.1 Securely attach the test plates to the testing machine platform by wire tying. The test plates are placed end-to-end so that the test pads are in numerical order with test pad No. 1 being on the driver's side of the truck.

5.1.2 Position the distributor truck so that its spray bar is over the testing machine, but far enough away from the test plates to prevent any bituminous material being sprayed on the test pads. Initiate the spraying. When the spray is judged to be normal, energize the platform so that it moves under and through the spray, as illustrated in Figure 4. Terminate the spray and pull the distributor truck away from the testing machine.

5.1.3 Proceed to Section 8.

Typical "Sliding Platform" Testing Machine
With Bituminous Material Being Sprayed

Figure 4
6. ALTERNATE SAMPLING PROCEDURE #2

6.1 If desired, Alternate Sampling Procedure #2 may be used instead of Alternate Sampling Procedure #1 described in Section 5 or Alternate Sampling Procedure #3 described in Section 7. Side shields shall not be used when testing is performed under this procedure.

6.1.1 Place the boards used to raise the plates off the surface in a suitable location.

**Note:** The boards should be protected from the bituminous material by wrapping them in heavy paper.

6.1.2 Place the test plates end-to-end on top of the boards so that the test pads are in numerical order with test pad No. 1 being on the driver's side of the truck.

6.1.3 Back the distributor truck up to test plates so that the back wheels of the truck are a few inches from the test plates.

6.1.4 Drive the distributor truck forward at a constant speed and spray bituminous material across the test plates.

6.1.5 Proceed to Section 8.

7. ALTERNATE SAMPLING PROCEDURE #3

7.1 If desired, Alternate Sampling Procedure #3 may be used instead of Alternate Sampling Procedure #1 described in Section 5 or Alternate Sampling Procedure #2 described in Section 6. Side shields shall not be used when testing is performed under this procedure.

7.1.1 Place the boards used to raise the plates off the surface in a suitable location, which will facilitate the distributor truck backing up at a constant speed with the ability to stop prior to its wheels hitting the test plates.

**Note:** The boards should be protected from the bituminous material by wrapping them in heavy paper.
7.1.2 Place the test plates end-to-end on top of the boards so that the test pads are in numerical order with test pad No. 1 being on the driver's side of the truck.

7.1.3 With the distributor truck starting at an adequate distance in front of the test plates, it is backed up at a constant speed so that the bituminous material is sprayed across the test plates. The distributor truck must be stopped prior to its wheels hitting the test plates.

7.1.4 Proceed to Section 8.

8. **REMOVAL OF TEST PADS FROM THE TEST PLATES**

8.1 After the spraying operation is complete, leave the test plates in place a short time to allow the bituminous material to set up enough to prevent running.

8.2 Remove the test plates and transport them to the laboratory.

8.3 Place each test plate in turn on an adequate size work area. Starting with the highest number test pad and proceeding to the lowest number, remove each test pad taking care to not remove any of the paper when removing the masking tape. Fold each test pad, and stack them in a drying pan as shown in Figure 5.

8.4 Dry the test pads to constant weight at 140 ± 5°F. Allow them to cool to room temperature prior to weighing.

**Note:** If PG asphalt binder is used, drying of the test pads at 140 °F is not required.

8.5 Weigh each test pad and record its weight to the nearest 0.1 gram on the test record shown in Figure 8.

9. **CALCULATIONS**

9.1 Omit end test pads that are outside the spray bar width.

9.2 Determine the weight of bituminous material on each of the remaining test pads, by subtracting the corresponding test pad tare weight, and record the weight to the nearest 0.1 gram on the test record shown in Figure 7.
Folded and Stacked Test Pads

Figure 5

9.3 Multiply the weight of bituminous material on each test pad by 0.0107, or use the table found in Figure 6, to determine the spread rate to the nearest 0.001 gal./sq. yd. for each test pad.

9.4 Determine the average spread rate to the nearest 0.001 gal./sq. yd.

9.5 Determine the acceptable range as shown below:

9.5.1 Acceptable Range = Average spread rate ± 10%, or ± 0.020 gal/sq. yd., whichever is less.

9.6 Determine the number of test pads that are outside the acceptable range. More than three pads outside the acceptable range constitutes a failing test. Three or fewer test pads outside the acceptable range constitutes a passing test.
10. REPORT AND EXAMPLE

10.1 Report the following:

10.1.1 The spread rate, to the nearest 0.001 gal./sq. yd., for each test pad used in the spread rate calculations.

10.1.2 The average spread rate, to the nearest 0.001 gal./sq. yd.

10.1.3 Any test pads outside the acceptable range.

10.1.4 The person performing the test, the test date, the contractor's name, and identifying information on the truck tested (the license plate number and the truck number).

10.1.5 The Alternate Sampling Procedure used.

10.1.6 Whether the test result indicates a passing result or a failing result.

10.2 A blank test report form is shown in Figure 7. An example of a completed test report is given in Figure 8.
## CONVERSION TABLE

(Weight of bituminous material on a 4" by 8" cotton pad to gals./sq. yd.)

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*Figure 6*
ARIZONA DEPARTMENT OF TRANSPORTATION

BITUMINOUS DISTRIBUTOR TRUCK TRANSVERSE SPREAD RATE
(Arizona Test Method 411)

OWNER/CONTRACTOR: ___________________________  TESTED BY: _______________  DATE: ______

TRUCK #: _______  TRUCK LICENSE PLATE #: _______________  SPRAY WIDTH ______ FT.

AVERAGE SPREAD RATE: ______ GALS./SQ. YD.  ACCEPTABLE RANGE: UPPER ______ LOWER: ______

TYPE OF BITUMINOUS MATERIAL USED IN TEST: ______  TEST RESULT: PASS ______ FAIL ______

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* SPREAD RATE = GALS/SQ. YD.

REMARKS: ____________________________________________________________
### ARIZONA DEPARTMENT OF TRANSPORTATION

**BITUMINOUS DISTRIBUTOR TRUCK TRANSVERSE SPREAD RATE**  
(Arizona Test Method 411)

**OWNER/CONTRACTOR:** ACME ASPHALT TRANSPORT  
**TESTED BY:** JOHN DOE  
**DATE:** 09/28/12

**TRUCK #:** 960  
**TRUCK LICENSE PLATE #:** C 4568 A2  
**SPRAY WIDTH:** 16 FT.

**AVERAGE SPREAD RATE:** 0.139 GALS./SQ. YD.  
**ACCEPTABLE RANGE:**  
**UPPER:** 0.153  
**LOWER:** 0.125

**TYPE OF BITUMINOUS MATERIAL USED IN TEST:** SS-1

**TEST RESULT: PASS**

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* SPREAD RATE = GALS./SQ. YD.

**REMARKS:**
DENSITY OF COMPACTED BITUMINOUS MIXTURES - NUCLEAR METHOD

(An Arizona Method)

SCOPE

1. (a) This method is used to determine the in-place density of compacted layers of bituminous mixtures by use of nuclear apparatus.

     (b) This test method involves hazardous materials, operations, and equipment. This test method does not purport to address all of the safety problems associated with its use. It is the responsibility of whoever uses this test method to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

     (c) Safety procedures for operation, transport and storage of nuclear gauges shall be in accordance with the manufacturer’s recommendations and the applicable regulations of the Arizona Radiation Regulatory Agency (ARRA).

APPARATUS

2. The apparatus shall consist of the following:

     (a) Moisture/Density Nuclear gauge capable of determining densities by the backscatter method. Calibration of the gauge shall be performed in accordance with AASHTO T310, Annexes A1, A2 and A3, on an annual basis.

     (b) Reference standard block or test stand to obtain standard counts for moisture and density which are used to check the gauge stability.

     (c) Nuclear gauge transport case and labels which comply with A.R.R.A. Regulations.

     (d) Charging cord, if applicable.
(e) Radiation exposure badge (if required by license .with A.R.R.A.)

(f) Information packet for the nuclear gauge which shall contain the following items:

(g) Moisture/Density Calibration Tables (if required), and a standard count log book.

(h) Manufacturer’s Gauge Operation Manual for the nuclear gauge.

(i) Applicable documentation necessary to meet requirements of ARRA for gauge safety.

(j) Blank test forms for use on the applicable nuclear gauge.

(k) Calculator for necessary computations.

(l) Miscellaneous equipment including watch, pencils, writing paper, ruler, eraser, clip board, and hand cart as required.

GAUGE STABILITY CHECK

3. A density standard count and moisture standard count shall be taken at the beginning of each day of testing at the project where the field density testing is to be performed. The gauge stability check shall be performed as follows:

(a) Place the reference standard block on any asphalt, concrete, compacted aggregate or similar surface which is dry and level. The reference standard block should be at least at least 15 feet away from any large object, or vehicle, and at least 50 feet away from another nuclear gauge.

(b) Seat the nuclear gauge on the reference block in accordance with the gauge operation manual. It is very important that the gauge is seated properly on the standard reference block.

(c) Remove the lock on the source handle and make sure the source handle is in the safe or stored position (the top notch on the index rod).

(d) Turn the gauge on (in standby power condition) and allow it to warm-up, if necessary, for the recommended time as given in the gauge operation manual.
(e) After the warm-up period, take a standard moisture count and a standard density count in accordance with the gauge operation manual.

(f) Record the moisture and density standard counts in the proper columns of the standard count log book along with the appropriate additional information, such as date, time, temperature, and location.

(g) Return the gauge to the standby power condition. The gauge should be left in the standby mode for subsequent testing.

(h) Determine if the standard counts are within the limits for normal operation in accordance with the gauge operation manual. This is usually done by comparing the standard counts to the average of the four previous standard counts or utilizing an internal statistical test which is available on some gauges. Additional standard counts may be necessary if initially the gauge does not appear to be operating properly. If the gauge does not meet the normal operating parameters as specified by the Standard Count procedure in the gauge operation manual, the gauge should not be used for testing. It should be sent in for servicing to determine the problem.

NOTE: Some gauges will store standard counts for later use in calculations performed by the gauge itself. The most recent standard counts will usually be stored automatically over pre-existing standard counts.

(i) On a weekly basis, compare the average of the four most recent standard counts with the average of four standard counts immediately after gauge calibration or at least three months previous, whichever is shorter. If the accumulative shift in standard count exceeds 2% for moisture or 3% for density, the nuclear gauge should be recalibrated.

PROCEDURE

4. (a) At each location to be tested, two one-minute readings shall be obtained by taking the first reading and recording the wet density to the nearest 0.1 lb. per cu./ft. then rotating the gauge 180° (making sure that the gauge is set in the same footprint as the first reading) and taking another reading and again recording the wet density to the nearest 0.1 lb. per cu./ft. The two readings are then averaged.

(b) Normally the preparation of the surface for taking readings at each location shall not include the removal of any material for the purpose of making it more
smooth, except that particles which are completely unattached and merely lying loose on top of the compacted and bound mixture shall be brushed away. Not more than one pound of dry fine sand (minus #10 material) shall be spread over each location and then scraped away with a straightedge so that the mixture is visible over the majority of the surface.

**PRECAUTIONS**

5. (a) Except when actually in use performing tests, the gauge and its accessories are to be kept within the A.R.R.A. (Arizona Radiation Regulatory Agency) approved carrying case, to protect it from damage and to provide better radiation shielding for persons in its vicinity.
1. SCOPE

1.1 This procedure covers the determination of bulk specific gravity and bulk density of specimens of compacted bituminous mixtures.

1.2 This test procedure consists of three methods for determining bulk specific gravity and bulk density. These methods are referred to as "Method A", "Method C (Rapid Test)", and "Vacuum Method".

Note: If it is desired to use "Method B", which is not included in this procedure, AASHTO T 166 shall be referred to. Method B should not be used with samples that contain open or interconnecting voids and/or absorb more than 2.0 percent water by volume. Method B is not acceptable for specimens that have more than six percent air voids.

1.3 Method A should not be used with samples that contain open or interconnecting voids and/or absorb more than 2.0 percent water by volume as determined in Subsection 7.4. If the percent water absorbed by the specimen exceeds 2.0 percent, either Method C, Vacuum Method, or AASHTO T 275 “Bulk Specific Gravity (Gmb) of Compacted Hot Mix Asphalt (HMA) Using Paraffin-Coated Specimens” shall be used.

1.4 Referee testing shall be performed in accordance with "Method C".

1.5 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.6 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.
2. TEST SPECIMENS

2.1 Test specimens may be either laboratory molded specimens or samples from an existing bituminous pavement.

2.2 Size of specimens - It is recommended that: (1) the diameter of cylindrically molded or cored specimens, or the length of the sides of sawed specimens, be at least four times the maximum size of the aggregate and, (2) the thickness of specimens be at least one and one half times the maximum size of the aggregate.

2.3 Specimens obtained from bituminous pavements shall be taken with a core drill, diamond or carborundum saw, or by other suitable means.

2.4 Care shall be taken to avoid distortion, bending, or cracking of specimens during and after the removal from the pavement or mold. Specimens shall be stored in a safe, cool place.

2.5 Test specimens shall be free from paper, tape, soil, and other foreign materials not intended to be tested as part of the specimen.

2.6 If desired, specimens may be separated from other pavement layers by sawing or other suitable means. Care shall be exercised to ensure that specimens are not damaged during the separation process.

3. APPARATUS

3.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

3.1.1 Balance - A balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.1 gram. The balance shall be equipped with a suitable suspension apparatus and holder to permit weighing the specimen while suspended from the center of the balance.
3.1.2 Suspension Apparatus - The wire(s) suspending the holder shall be the smallest practical size to minimize any possible effects of a variable immersed length. The suspension apparatus shall be constructed to enable the container to be immersed to a depth sufficient to cover it and the test sample during weighing.

3.1.3 Water bath - The water bath for immersing the specimen in water while suspended under the balance shall be equipped with an overflow outlet or a clearly marked “fill line” for maintaining a constant water level. The level of the water shall be adjusted before testing each specimen, if necessary, to maintain a constant water level.

3.1.4 Oven - An oven capable of drying specimens at either $125 \pm 5 \, ^\circ F$ (Method A) or $290 \pm 10 \, ^\circ F$ (Method C).

3.1.5 Vacuum Drying Device (For Vacuum Method) – Conforming to the requirements of AASHTO PP 75. Automatic controls of the unit shall be calibrated by the manufacturer prior to initial use.

### METHOD A

#### 4. PROCEDURE

4.1 Dry the specimen to constant mass. Constant mass shall be defined as the mass at which further drying at $125 \pm 5 \, ^\circ F$ does not alter the mass by more than 0.1 gram. Samples saturated with water shall initially be dried overnight at $125 \pm 5 \, ^\circ F$ and then weighed at two-hour drying intervals. Recently molded laboratory samples, which have not been exposed to moisture, do not require drying.

4.2 Cool the specimen to room temperature at $77 \pm 9 \, ^\circ F$ and record the dry mass to the nearest 0.1 gram as "A".

4.3 Immerse the specimen in water at $77 \pm 2 \, ^\circ F$ for $4 \pm 1$ minutes and record the immersed mass to the nearest 0.1 gram as "C". Care shall be exercised to ensure that no trapped air bubbles exist under the specimen.

4.4 Remove the specimen from the water, quickly damp dry the specimen surface by blotting with a damp towel, and as quickly as possible determine and record the surface-dry mass to the nearest 0.1 gram as "B". Any water that seeps from the specimen during the surface-dry
A terry cloth towel has been found to work well. Damp is considered to be when no water can be wrung from the towel.

Note: If desired, the sequence of testing operations may be changed to expedite the test results. For example, first the immersed mass, "C", can be taken, then the surface-dry mass, "B", and finally the dry mass, "A".

4.5 Calculate the bulk specific gravity, bulk density, and percent water absorption by volume in accordance with Section 7.

### METHOD C (RAPID TEST)

**5. PROEDURE**

5.1 This procedure can be used for testing specimens which are not required to be saved and which contain a substantial amount of moisture. Specimens obtained by methods that introduce moisture, such as wet coring or sawing, can generally be tested the same day by this method.

5.2 Ensure the specimen is at room temperature (77 ± 9 °F).

5.3 Immerse the specimen in water at 77 ± 2 °F for 4 ± 1 minutes and record the immersed mass to the nearest 0.1 gram as "C". Care shall be exercised to ensure that no trapped air bubbles exist under the specimen.

5.4 Remove the specimen from the water, quickly damp dry the specimen surface by blotting with a damp towel, and as quickly as possible determine and record the surface-dry mass to the nearest 0.1 gram as "B". Any water that seeps from the specimen during the surface-dry weighing operation is considered as part of the specimen. The determination of the immersed mass and surface-dry mass of each specimen will be completed before the next specimen is submerged for its immersed mass determination.
5.5 Place the specimen in a large, flat-bottom drying pan of known weight. Place the pan and specimen in a \(290 \pm 10\) °F oven. Leave the specimen in the oven until it can be easily separated to the point where particles of the fine aggregate portion are not larger than 1/4 inch. During separation of material, ensure that no material is lost. Place the separated specimen in the \(290 \pm 10\) °F oven and dry to constant mass. (Constant mass shall be determined as follows: Dry the sample for a minimum of 1 hour at \(290 \pm 10\) °F. Record the weight of the sample to the nearest 0.1 gram. Continue drying and weighing until the weight does not change more than 0.1 gram at drying intervals of a minimum of 30 minutes.)

5.6 Cool the pan and specimen to room temperature at 77 \(\pm\) 9 °F. Weigh the pan and specimen, subtract the mass of the pan, and record the dry mass of the specimen to the nearest 0.1 gram as "A".

**Note:** For Method C, the drying of specimens to constant weight may be accomplished in a microwave oven, as described in Arizona Test Method 719.

5.7 Calculate the bulk specific gravity, bulk density, and percent water absorption by volume in accordance with Section 7.

---

**VACUUM METHOD**

6. **PROCEDURE**

6.1 This procedure can be used for testing specimens, which are to be saved, and which contain a substantial amount of moisture. Specimens obtained by methods that introduce moisture, such as wet coring or sawing, can generally be tested the same day by this method.

6.2 Turn on the vacuum drying device. Follow the manufacturer’s recommendations for warm up and self-test procedures.

6.3 Run the vacuum drying device without any specimens. The pressure reading on the display should indicate a known dry point value as recommended by the manufacturer. If the indicated dry point is not achieved, refer to the manufacturer’s trouble shooting instructions.

6.4 Ensure the specimen is at room temperature (77 \(\pm\) 9 °F).
6.5 Remove any standing water from the surface of the specimen by using a paper towel or an absorptive cloth.

6.6 Dry the cold trap and the specimen chamber. Place the specimen inside the vacuum chamber.

**Note:** Two 4-inch diameter specimens, that are less than 3-inches in thickness can be placed side-by-side in the chamber during a single drying cycle. Larger specimens shall be placed in the chamber individually.

**Note:** Water and/or ice may buildup in the moisture trap during a drying cycle. Wipe off any free standing water in the moisture trap between drying specimens. This will expedite specimen drying.

6.7 Place the lid on the vacuum chamber and press the lid down to ensure secure contact between the lid and the chamber. Press the appropriate key on the vacuum drying device to begin the drying process.

6.8 The vacuum drying device will automatically stop when the specimen is dry. The unit shall be calibrated to sense the “dry specimen condition”. The “dry specimen condition” shall be determined from the calibrated pressure at which no water remains in the chamber. The pressure is monitored throughout the drying process to ensure that the “dry specimen condition” pressure is achieved in the device.

6.9 Remove the specimen from the vacuum drying device. Weigh the specimen and record the dry mass of the specimen to the nearest 0.1 gram as "A".

6.10 Immerse the specimen in water at $77 \pm 2 ^\circ F$ for $4 \pm 1$ minutes and record the immersed mass to the nearest 0.1 gram as "C". Care shall be exercised to ensure that no trapped air bubbles exist under the specimen.

6.11 Remove the specimen from the water, quickly damp dry the specimen surface by blotting with a damp towel, and as quickly as possible determine and record the surface-dry mass to the nearest 0.1 gram as "B". Any water that seeps from the specimen during the surface-dry weighing operation is considered as part of the specimen. The determination of the immersed mass and surface-dry mass of each specimen will be completed before the next specimen is submerged for its immersed mass determination.
Note: If desired, the sequence of testing operations may be changed to expedite the test results. For example, first the immersed mass, "C", can be taken, then the surface-dry mass, "B", and finally the dry mass, "A".

6.12 Calculate the bulk specific gravity, bulk density, and percent water absorption by volume in accordance with Section 7.

7. **CALCULATION**

7.1 Calculate and record the bulk specific gravity of the specimen to the nearest 0.001 as follows:

\[
\text{Bulk Specific Gravity} = \frac{A}{B - C}
\]

Where:  
\(A\) = mass of dry specimen  
\(B\) = mass of surface-dry specimen  
\(C\) = mass of immersed specimen

7.2 Calculate and record the bulk density of the specimen to the nearest 0.1 lb/cu ft by multiplying the bulk specific gravity by 62.3 lbs/cu ft.

7.3 For laboratory molded specimens of 1/2-inch or 3/4-inch asphaltic concrete mixes, the range of bulk density results for three replicate specimens shall not differ by more than 2.5 lbs/cu ft. For laboratory molded specimens of asphaltic concrete Base Mixes, the range of bulk density results for three replicate specimens shall not differ by more than 3.0 lbs/cu ft. If the respective requirement is not met, the entire set of specimens shall be discarded and a new set of specimens shall be prepared and tested.

7.4 Calculate and record the percent water absorbed by the specimen to the nearest 0.01 percent (on volume basis) as follows:

\[
\text{Percent Water Absorption by Volume} = \left(\frac{B - A}{B - C}\right) \times 100
\]
8. REPORT

8.1 The method that was used.

8.2 The bulk specific gravity to the nearest 0.001.

8.3 The bulk density to the nearest 0.1 lb/cu ft.

8.4 The absorption to the nearest 0.01 percent.
PREPARING AND SPLITTING FIELD SAMPLES
OF BITUMINOUS MIXTURES FOR TESTING

(An Arizona Method)

1. SCOPE

1.1 This procedure describes the preparation and splitting of field samples of bituminous mixtures for testing.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.3 Oven capable of maintaining a temperature of 290 ± 10 °F.

2.4 Closed-end samplers as illustrated in Figure 1 (or similar), constructed of 16 to 18 gauge sheet metal, having a height of 3 inches, a minimum length of 13 inches, and widths of approximately 2-1/2 inches, 3 inches, or 3-1/2 inches.

2.5 A concrete trowel or hand float. If desired, a straightedge of sufficient length to span the final diameter of the circular mass may be used.

2.6 Small scoop, spatulas, and suitable size containers.
3. PREPARATION OF SAMPLE

3.1 Samples may be stored for indefinite periods of time at temperatures not exceeding 140 °F.

3.2 The material shall be easily workable and pliable when splitting. If necessary, the sample may be heated at 290 ± 10 °F for a maximum of 3 hours. The 3-hour time period begins when the oven reaches the specified temperature.

3.3 If necessary, the material shall be reduced in size to provide a workable amount of material from which to obtain all required samples by thoroughly mixing and quartering, splitting with a mechanical (clam-shell) splitter, or using a four-way splitter such as a “Quartermaster”. When utilizing a mechanical (clam-shell) splitter, the width of the individual chute openings shall be approximately 1-1/2 to 2-1/2 times larger than the largest particles in the sample to be split. Generally it will only be necessary to reduce the material one time by quartering or splitting. A light coat of vegetable cooking spray may be sprayed on the equipment to help shed the asphaltic concrete. In no case shall diesel fuel or similar solvent be used. Individual samples for testing shall not be obtained by quartering or splitting, but rather as described in Section 4.
3.4 Place the hot material on a tarp or a sheet of heavy paper large enough to manipulate the sample. In a rolling motion thoroughly mix the material. Leave the mound in a circular shape after rolling is completed.

3.5 Spread the material into a circular mass. Spreading may be accomplished either by leveling the mound of material with a concrete trowel or hand float; or by placing a straightedge of sufficient length to span the final diameter of the circular mass over the center of the material and rotating it until the desired height is obtained. Whichever method is utilized, the operator shall assure that the material is evenly distributed with as little segregation as possible. The thickness of the circular mass shall not exceed 3 inches. Generally a depth of 1-1/2 to 2 inches will provide acceptable size samples.

3.6 At any time during obtaining test samples, the material may be reheated for a maximum of 1 hour at a temperature not exceeding 300 °F to restore pliability.

4. **OBTAINING TEST SAMPLES**

4.1 The required samples for testing, with the exception of moisture content, shall be obtained as described in Subsections 4.2 through 4.6 below. For obtaining test samples for moisture content, see Subsection 4.7 below. The samples may be obtained in any sequence as long as the sample for moisture content is taken immediately before or after the sample for determination of asphalt content. The width of sampler to be used is dependent upon the size of aggregate in the sample and/or the amount of material needed.

4.2 The samples shall be obtained by placing the closed end of the sampler as near the center of the mass as possible with the open end of the sampler extending beyond the edge of the circular mass (see Figure 2). Force the sampler down to the bottom of the pile and remove the contents that are captured by sliding the sampler out of the pile, and placing the contents into a tared container.

4.3 Obtain additional material, as necessary, by repeating the procedure in Subsection 4.2, at a different location in the pile so that a cut does not overlap a previous cut (see Figure 3).

4.4 If small amounts of material are needed, slide the sampler out and to the side of the pile. Lift the sampler up and turn it perpendicular to the material. Force the sampler down through the full width of the material, starting at the closed end portion of the material. (Figure 4 provides an illustration of this procedure.) If necessary, additional material may be obtained by taking multiple cuts. Utilize the entire portion(s) taken and do not attempt to obtain an exact weight.
4.5 If excess material is obtained, the sample shall be returned to a place not disturbing the rest of the circular mass. If the remaining mass is large enough, and is undisturbed, obtain another sample for the test, if necessary utilizing a smaller width sampler. Alternatively, the sample may be returned to the circular mass and the material re-rolled and spread to a thinner depth.

4.6 If the remaining mass is not large or uniform enough to obtain required samples, re-roll and spread the material in the same manner described in Subsection 3.5.

4.7 Test samples for determining moisture content may be obtained by use of the sampler as described in Subsections 4.2 through 4.6 above, or by taking several small portions with a small scoop at random locations throughout the mass.

FIGURE 2
MAXIMUM THEORETICAL SPECIFIC GRAVITY AND DENSITY OF FIELD PRODUCED BITUMINOUS MIXTURES (RICE TEST)

(A Modification of AASHTO T 209)

1. SCOPE

1.1 This method of test is intended for determining the maximum specific gravity and density of uncompacted bituminous mixtures that have been field produced.

Note: Two methods are provided for determining the maximum specific gravity. The method given in Section 6 is for determining results without fan drying the samples. Section 7 describes the procedure which is used when fan drying is necessary. For the first four samples taken at the beginning of production on a project the maximum specific gravity shall be determined in accordance with Section 6 and also shall be fan dried and maximum specific gravity determined in accordance with Section 7. If the difference in resultant air voids, when determined as described in Arizona Test Method 424 is greater than 0.2% subsequent samples will be subjected to fan drying. During the course of the project comparisons should be made on approximate 10 sample intervals to determine need for fan drying. In case of dispute, fan drying shall be used.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.
2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Balance - A balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.1 gram.

2.3 Container - A heavy walled Erlenmeyer flask having a capacity of at least 1500 mL and strong enough to withstand a partial vacuum; the cover shall consist of a rubber stopper with a tight hose connection. A small piece of No. 200 wire mesh covering the hose opening shall be used to minimize the possibility of loss of fine material.

Note: If a procedure which subjects multiple flasks to a vacuum simultaneously is used, the vacuum gauge shall be placed beyond the last bottle to insure that all the bottles are being subjected to the same amount of vacuum.

2.4 Flat glass plate large enough to cover mouth of the flask.

2.5 Vacuum pump for evacuating air from the container.

2.6 All water used in this procedure shall be distilled or de-mineralized water.

3. CALIBRATION OF FLASK

3.1 Record the weight of the flask and flat glass plate separately to the nearest 0.1 gram on the work card. Using water at a temperature of 77 ± 1 °F, fill the flask with water to approximately one inch below the top of the flask. Using a long narrow rod, remove air bubbles adhering to the walls of the flask. Confirm that the temperature of the water is at 77 ± 1 °F. Fill to the top and slide the flat glass plate over the mouth of the flask. Verify that no air is trapped under the flat glass plate. Dry the outside of the flask and glass plate and weight to the nearest 0.1 gram. Subtract the weight of the glass plate and record the weight of the “flask and water” as "B".
4. PREPARATION OF SAMPLES

4.1 Obtain 3 representative 1050 ± 50 gram samples of the material, as described in Arizona Test Method 416.

Note: If necessary, heat the sample for not more than one hour at a maximum temperature of 285 °F ONLY until it is pliable enough to allow separation of the coated aggregate.

4.2 Spread each sample on a sheet of heavy paper or in a large flat bottom pan. Before the samples are completely cooled, separate the particles of the mixture, taking care not to fracture the coarse aggregate particles, so that the particles of the fine aggregate portion are not larger than 1/4 inch. Allow the samples to cool to room temperature.

5. PROCEDURE

5.1 Place the sample in the flask and determine the weight to the nearest 0.1 gram. Subtract the weight of the flask and record the "weight of sample in air" as "Wmm".

5.2 Add sufficient water to cover the sample. The water shall be at a temperature of approximately 77 °F and shall have been treated with a wetting agent.

Note: Aerosol OT in a concentration of 0.01%, or one mL of 10% solution per 1000 mL of water, has been found to be a suitable wetting agent and shall be used to facilitate the release of entrapped air.

5.3 Remove entrapped air by subjecting the contents of the flask to a partial vacuum with a minimum of 20 inches of mercury (gauge) for 15 ± 2 minutes, agitating the contents of the flask four times at evenly spaced intervals throughout this period.

CAUTION: Do not agitate the sample too frequently or vigorously, as that can cause stripping of the asphalt film from some particles, resulting in erroneous specific gravities.

5.4 After the evacuation period, fill the flask with water to approximately one inch below the top of the flask. Gently stir the sample with a long narrow rod in such
a way to release any trapped air bubbles, avoiding breakage of the aggregates. Using the long narrow rod, carefully remove any air bubbles adhering to the walls of the flask. Fill completely to the top and confirm that the temperature is at 77 ± 2 °F. Slide the pre-weighed flat glass plate over the mouth of the flask. Verify that no air is trapped under the flat glass plate. Dry the outside of the flask and glass plate and weigh immediately to the nearest 0.1 gram. Subtract the weight of the glass plate and record the weight of the "flask + water + sample" as "C".

6. **CALCULATIONS**

6.1 The Volume of Voidless Mix, "Vvm", in mL, and Maximum Specific Gravity, “Gmm” is determined for each sample by the following:

\[ Vvm = Wmm + B - C \]

\[ Gmm = \frac{Wmm}{Vvm} \]

Where:  
- Wmm = Wt. of Sample in Air  
- B = Wt. of Flask + Water  
- C = Wt. of Flask + Water + Sample

6.2 Compare the three individual values for maximum specific gravity. If the range of the three is within 0.024, all are used to determine the average maximum specific gravity as shown in Subsection 6.4. If the range is greater than 0.024, the average of two may be used if they are within a range of 0.012. If values are not achieved within the above criteria, the samples shall be discarded and a set of three new samples shall be tested. If material is not available, results should be used cautiously in the analysis of the bituminous mix. If results are used for specification compliance, additional material must be obtained for retesting.

6.3 The average maximum specific gravity of the bituminous mix is determined for the samples with acceptable maximum specific gravity values, and recorded to the nearest 0.001.

6.4 To determine the maximum density, the average maximum specific gravity is multiplied by 62.3 lbs./cu. ft.
PROCEDURE FOR FAN DRYING SAMPLES

7.1 The entire contents of the flask shall be poured into a nest of sieves consisting of a No. 40 and a No. 200 screen.

Note: If stripping has occurred, as evidenced by discoloration of water in the flask, significant loss of Minus No. 200 material may be expected. Provisions for the recovery and addition of this material to the Plus No. 200 material shall be made.

7.2 Allow the mix to drain through the sieves until excess moisture is removed from the mix. Separate the sieves and place both sieves in a tared pan. Place the pan in front of a fan (see Note below) and dry the material retained on the No. 40 and No. 200 sieves until the material can be easily removed from the sieves. Remove the material from the sieves and spread it in the pan. Determine and record the initial weight of the pan and the material.

Note: The air through the fan shall be at room temperature and no heat shall be used to dry the material.

7.3 Continue fan drying the material, determining and recording the weight of the pan and the material at 15 minute intervals. When the weight loss is 0.5 gram or less for a 15 minute interval, the mix is considered to be surface dry. Record the surface dry weight as "Wsd". Intermittent stirring of the sample is required during the drying period. Conglomerations of the mix shall be broken by hand. Care must be taken to prevent loss of particles of the mixture.

Note: If the "Wsd" weight for any of the three samples is less than its corresponding "Wmm" weight, the samples shall be discarded and a set of three new samples shall be tested. If material is not available, the maximum specific gravity shall be determined utilizing the "Wmm" weight and results should be used cautiously in the analysis of the bituminous mix. If results are used for specification compliance, additional material must be obtained for retesting.
7.4 To calculate the fan dried "Vvm" and maximum specific gravity, "Gmm", of each sample, the surface dry weight, "Wsd", is substituted for "Wmm" in the equation given for "Vvm" in Subsection 6.1, as shown in the example below.

\[ V_{vm} = W_{sd} + B - C \]

\[ G_{mm} = \frac{W_{mm}}{V_{vm}} \]

Where:
- \( W_{mm} \) = Wt. of Sample in Air
- \( W_{sd} \) = Wt of Fan Dried Sample
- \( B \) = Wt. of Flask + Water
- \( C \) = Wt. of Flask + Water + Sample

8. EXAMPLE

8.1 An example of the completed form is shown in Figure 1 and an example of the blank form is shown in Figure 2.
If samples were fan dried, the maximum density is determined utilizing the "Wsd" weight as shown below:

<table>
<thead>
<tr>
<th>Flask Number or I.D.</th>
<th>&quot;WB&quot;</th>
<th>&quot;Wmm&quot;</th>
<th>&quot;B&quot;</th>
<th>&quot;C&quot;</th>
<th>&quot;Vvm&quot;</th>
<th>&quot;Gmm&quot;</th>
<th>&quot;Wsd&quot;</th>
<th>&quot;Vvm&quot;</th>
<th>&quot;Gmm&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Wt. of Flask</td>
<td>Wt. of Sample in Air</td>
<td>Wt. of Flask + Water</td>
<td>Wt. of Flask + Sample + Water</td>
<td>Volume of Voidless Mix</td>
<td>Maximum Specific Gravity</td>
<td>Maximum Density (lbs./cu. ft.)</td>
<td>Volume of Voidless Mix</td>
<td>Maximum Specific Gravity</td>
</tr>
<tr>
<td>1</td>
<td>998.7</td>
<td>1046.9</td>
<td>3215.7</td>
<td>3848.7</td>
<td>431.9</td>
<td>2.466</td>
<td>1067.7</td>
<td>434.7</td>
<td>2.450</td>
</tr>
<tr>
<td>2</td>
<td>977.6</td>
<td>1076.5</td>
<td>3178.7</td>
<td>3819.2</td>
<td>436.0</td>
<td>2.469</td>
<td>1080.5</td>
<td>440.0</td>
<td>2.447</td>
</tr>
<tr>
<td>3</td>
<td>994.3</td>
<td>1097.4</td>
<td>3294.1</td>
<td>3925.1</td>
<td>436.4</td>
<td>2.446</td>
<td>1071.2</td>
<td>440.3</td>
<td>2.425</td>
</tr>
<tr>
<td>AVERAGE</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2.468</td>
<td>153.8</td>
<td></td>
<td>2.449</td>
</tr>
</tbody>
</table>

**Remarks:** Flask #3 eliminated from the average due to Specific Gravity being outside specified 0.024 allowable range.

**Specific Gravity Range:** Range of 3: 0.023 - 0.003

### Maximum Specific Gravity (Rice) Fan Dry Weigh backs

<table>
<thead>
<tr>
<th>Flask Number or I.D.</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tare weight of Pan</td>
<td>453.7</td>
<td>502.4</td>
<td>499.6</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1536.3</td>
<td>1597.8</td>
<td>1585.7</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1529.2</td>
<td>1590.7</td>
<td>1578.6</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1525.0</td>
<td>1584.7</td>
<td>1572.6</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1527.4</td>
<td>1584.0</td>
<td>1571.0</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1526.9</td>
<td>1583.4</td>
<td>1571.2</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1521.4</td>
<td>1582.9</td>
<td>1570.8</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1521.4</td>
<td>1582.9</td>
<td>1570.8</td>
</tr>
<tr>
<td>Weight of Pan and Sample</td>
<td>1521.4</td>
<td>1582.9</td>
<td>1570.8</td>
</tr>
<tr>
<td>Surface Dry Weight (Wsd)</td>
<td>1067.7</td>
<td>1080.5</td>
<td>1071.2</td>
</tr>
</tbody>
</table>

### Air Voids Calculations

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>A C Mix</td>
<td>1</td>
<td>145.5</td>
<td>5.4 %</td>
</tr>
<tr>
<td>Bulk Density</td>
<td>153.8</td>
<td>5.4 %</td>
<td></td>
</tr>
<tr>
<td>Maximum Density From Rice Test</td>
<td>153.8</td>
<td>5.4 %</td>
<td></td>
</tr>
<tr>
<td>Difference in Air Voids = [Air Voids (Sample Not Fan Dried)] - [Air Voids (Sample Fan Dried)] =</td>
<td>0.8 %</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
If samples were fan dried, the maximum density is determined utilizing the "Wsd" weight as shown below:

<table>
<thead>
<tr>
<th>Flask Number or I.D.</th>
<th>&quot;Wf&quot;</th>
<th>&quot;Wmm&quot;</th>
<th>&quot;B&quot;</th>
<th>&quot;C&quot;</th>
<th>&quot;Vmm&quot;</th>
<th>&quot;Gmm&quot;</th>
<th>&quot;Wsd&quot;</th>
<th>Vmm</th>
<th>&quot;Gmm&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt. of Flask</td>
<td>Wf</td>
<td>Wt. of Sample in Air</td>
<td>Wf s - Wf</td>
<td>Wt. of Flask + Sample + Water</td>
<td>Wa - Wp</td>
<td>Wm m + B - C</td>
<td>Gm m (lbs/cu. ft.)</td>
<td>(See Below)</td>
<td>Wsd + B - C</td>
</tr>
<tr>
<td>AVERAGE</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Remarks:

Specific Gravity Range: Range of 3: Range of 2:

Non Fan Dried

Fan Dried

Maximum Specific Gravity (Rice) Fan Dry Weigh back

Air Voids Calculations

<table>
<thead>
<tr>
<th>Flask Number or I.D.</th>
<th>Tare weight of Pan</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Weight of Pan and Sample</th>
<th>Surface Dry Weight (Wsd)</th>
</tr>
</thead>
</table>

Air Voids Calculations

Non Fan Dried

Fan Dried

Difference in Air Voids = ([Air Voids (Sample Not Fan Dried)] - [Air Voids (Sample Fan Dried)])

FIGURE 2
BITUMINOUS MATERIAL CONTENT OF ASPHALTIC CONCRETE MIXTURES BY THE NUCLEAR METHOD

(An Arizona Method)

SCOPE

1. (a) This procedure describes a method for determining the percent bituminous material content of asphaltic concrete mixtures, by testing a sample with a device that utilizes neutron thermalization techniques.

(b) Modifications to this procedure for Asphaltic Concrete Friction Course, Asphaltic Concrete (Asphalt-Rubber), and Asphaltic Concrete Friction Course (Asphalt-Rubber) are given in Section 7.

(c) This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

(d) Safety procedures for operation, transport, and storage of nuclear gauges shall be in accordance with the manufacturer's recommendations and the applicable regulations of the Arizona Radiation Regulatory Agency (ARRA).

(e) If recommended by the manufacturer, equipment operators should wear an approved form of radiation dosimetry film badge capable of monitoring the occupational radiation exposure.

(f) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(g) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with
applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

**APPARATUS**

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

(a) Nuclear asphalt content gauge and manufacturer's instruction manual. While exact details of the manufacture of the apparatus may vary, the system shall consist of the following items and meet the criteria for variability given below. The equipment shall be so constructed as to be licensable in accordance with applicable health and safety regulations.

1) Neutron Source - An encapsulated and sealed radioactive source such as "Americium-241: Beryllium", or "Californium-252".

2) Detectors - Any type of thermal neutron detector, such as helium-3 or boron tri-fluoride.

3) Read-Out Instrument, such as a scaler or a direct-reading digital device calibrated in percent asphalt cement.

4) A minimum of four standard stainless steel sample pans [approximately 230 mm long by 185 mm wide by 100 mm deep (9" x 7-1/4" x 4"").

5) The variability of the gauge shall be no greater than ±0.05 percent asphalt for a four-minute count. The variability shall be determined utilizing a three point calibration for a conventional asphaltic concrete mix with a design asphalt content of approximately five percent. The variability of the gauge is determined from the slope of the calibration curve and the standard deviation of the measured counts of the calibration sample at the mix design percent asphalt. If the gauge fails to meet the variability requirement shown above for a four-minute count, the variability may be determined utilizing either 8-minute or 16-minute counts as necessary. If the variability is determined using an increased count time, that count time is the minimum that is to be used for testing subsequent field samples [See paragraphs 6(g) and 6(i)]. In lieu of an increased count time, a new set of calibration samples may be prepared and tested with greater care. If the gauge fails to meet the variability requirement after increased count times or preparation of additional calibration samples, the gauge must be returned to the manufacturer for necessary repair and recalibration. Variability is
calculated as shown below. The variability of the gauge should be checked at minimum 12 month intervals. Documentation of the variability determination shall be kept on file.

\[ V = \frac{S.D.}{S} \]

Where:  
\( V \) = apparatus variability, in percent asphalt.  
\( S \) = slope of the calibration curve, in counts (y-axis) per percent asphalt cement (x-axis). [See note below.]  
\( S.D. \) = standard deviation, in counts. The standard deviation is calculated from 20 individual four-minute readings (8-minute or 16-minute counts if used) taken on the calibration sample at the mix design percent asphalt content.

NOTE: The slope of the calibration curve, as calculated and given by the gauge may not be the actual numerical value for the slope, and in addition, the slope may not be given in counts (y-axis) per percent asphalt (x-axis). For instance, the Troxler Nuclear Asphalt Content Gauge (Model 3241-C) gives a numerical value for slope which is actually 1000 times the actual slope, and in addition, the value given by the Troxler gauge is given for percent asphalt (y-axis) per counts (x-axis). The following is an example of the slope of the calibration curve as given by the Troxler gauge and the determination of the corrected slope value to be used in the Apparatus Variability equation given above:

Troxler gauge reported slope = 5.220075  
[Slope based on percent asphalt (y-axis) per counts (x-axis).]

Determination of corrected slope:

\[ \frac{5.220075}{1000} = 0.005220075 \text{ (percent asphalt/counts)} \]
Reciprocal of slope to obtain slope based on counts (y-axis) per percent asphalt (x-axis):

\[
\frac{1}{0.005220075} = 191.568 \text{ (counts/percent asphalt)}
\]

In this case, the value of 191.568 is the slope of the calibration curve and is the value which is to be used for “S” in the equation for Apparatus Variability given above.

(b) Scale(s) or balance(s) capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be as shown below:

1) The scale or balance to be used for preparing calibration samples shall have a readability and sensitivity of at least 1 gram.

2) The scale or balance to be used for testing field samples shall have a readability and sensitivity of at least 5 grams.

(c) Oven(s), capable of heating and maintaining temperatures to 177 °C (350 °F). Two ovens may be required.

(d) Steel straightedge, approximately 300 mm (12 inches) in length.

(e) Either a piece of plywood of at least 20 mm (3/4 inch) nominal thickness, or a metal plate of at least 6 mm (1/4 inch) nominal thickness, having an area slightly larger than the top of the sample pans.

(f) Mixing apparatus - Mechanical mixing is recommended; 19 liter (20 quart) capacity mixer is required. (Hand mixing may be performed if desired.)

(g) Miscellaneous pans, bowls, spoons, and spatulas.

(h) Thermometer with temperature range 10 to 260 °C (50 to 500 °F).

(i) Hot plate(s) capable of heating and maintaining temperatures to 177 °C (350 °F).
(j) Flat bottom metal scoop at least 50 mm wide by 150 mm long (2 inches x 6 inches).

PRECAUTIONS

3. (a) Since nuclear equipment measures the total amount of hydrogen in the sample, this procedure is sensitive to changes in moisture content. It must be remembered that both bituminous material and water contain hydrogen.

(b) Keep any other source of neutron radiation at least 10 meters (30 feet) from the equipment.

(c) Do not place the equipment where the amount of hydrogenous material may change during the calibration or testing procedures. Hydrogenous materials are those containing hydrogen, for example, water or plastic materials.

(d) Moving the gauge to a new location may have an effect on the results. Therefore, a new background count and/or calibration is required if the gauge is moved. If the original calibration specimens are retested, they shall be heated in an oven at 121 to 149 °C (250 to 300 °F) for at least one hour to drive out any condensation or absorbed moisture and to bring them to testing temperature.

SAMPLING

4. (a) For preparing calibration samples, aggregate samples are obtained in accordance with Arizona Test Method 105. Samples shall be adequately dried, if necessary, to a free-flowing condition in the portion passing the 4.75 mm (No. 4) sieve. A maximum of approximately 45 kg (100 pounds) of aggregate, representative of the mix design gradation, will normally be required.

(b) When testing field samples of Asphaltic Concrete, a sample of the freshly produced mix is obtained in accordance with Arizona Test Method 104.

CALIBRATION

5. (a) A calibration must be performed for each asphaltic concrete mixture that is to be used on the project. Test results on asphaltic concrete mixtures may vary with changes in the type and gradation of aggregate, (percentage and source of...
mineral admixture, if used), and percentage and source of bituminous material. Accordingly, a calibration curve must be developed for each mix design. The calibration curve shall be established with three or more points. The calibration procedure consists of preparing and testing one dry aggregate blank sample, and a minimum of three asphaltic concrete calibration samples at varying bituminous material contents.

(b) Before a calibration is performed, a 16-minute background count shall be taken and recorded.

(c) For the dry aggregate blank sample, weigh up one approximate 10 kg (22 pound) aggregate sample representative of the mix design aggregate gradation. Thoroughly blend the aggregate and dry to constant weight at 121 to 149 °C (250 to 300 °F), and then place in a tared sample pan in at least two approximately equal depth layers. After placing each layer in the pan, gently spade the aggregate with the scoop or wide blade-like tool. The coarse and fine material shall be uniformly distributed. Too much manipulation of the material can result in segregation of the material, causing the fines to migrate to the bottom. Fill the pan slightly above the top rim. Place the straightedge firmly across the rim and, using a sawing motion, strike off the surface of the sample so that it is flush with the rim. Gaps between the straightedge and the sample shall be filled with fine aggregate and the sample leveled. Do not further compact the sample. Determine and record the weight of the dry aggregate blank sample to the nearest 5 grams.

(d) Place the dry aggregate blank sample in an oven at 121 to 149 °C (250 to 300 °F). A count will be determined on this sample, as described in paragraph 5(l), at the completion of the testing of the asphaltic concrete calibration samples.

(e) For the asphaltic concrete calibration samples, weigh up a minimum of three aggregate samples representative of the mix design aggregate gradation. These aggregate samples shall weigh at least 500 grams more than the dry aggregate blank sample weight determined in paragraph 5(c). If mineral admixture is used, the appropriate type and amount shall be added to the aggregate and thoroughly blended. Dry the samples to constant weight at the mixing temperature required as shown in paragraph (g) below.

(f) Bituminous material of the same source, grade, and type as that which will be used in production of the asphalt concrete mixture shall be used in the calibration samples. Normally three samples are prepared; one at the design bituminous material content, one at 1.0% above, and one at 1.0% below. The percent of bituminous material is based on the weight of total mix. The weight of bituminous material to be added for each percent of bituminous material is determined by the following:
(g) All bowls, sample pans, and mixing tools shall be heated to approximately 149 °C (300 °F). It is recommended that a "butter mix" be utilized to condition the mixing equipment. If a "butter mix" is utilized, the mixing equipment shall be scraped consistently clean with a spatula after each batch. As an alternate to a "butter mix", the mixing equipment shall be scraped as clean as possible with a spatula after each batch. All samples shall be mixed at the same mixing temperature, ± 6 °C (10 °F). During mixing, a hot plate shall be placed under the mixing bowl to reduce loss of heat while mixing. The sample shall be sufficiently mixed to ensure thorough coating. Unless otherwise indicated on the mix design, the temperature of the bituminous material and aggregate (and mineral admixture when used) at the time mixing begins shall be in accordance with the following:

\[
\text{Weight of Bituminous Material} = \left(\frac{\text{Weight of Aggregate (and mineral admixture when used)}}{\text{Percent of Aggregate}}\right) \times \left(\frac{\text{Percent of Bituminous Material}}{100 - \text{Percent of Bituminous Material}}\right)
\]

(h) At the time of beginning the fabrication of each asphaltic concrete calibration sample, the mixture shall be 121 to 149 °C (250 to 300 °F). The mix with the lowest bituminous material content shall be prepared first. Fill the tared sample pan in approximately equal depth layers (two layers minimum), uniformly distributing the sample in the pan. After placing each layer in the pan, gently spade the mix with a spatula, wide blade-like tool, or scoop and lightly press the mix down. Add mix until the top of the material is mounded approximately 25 to 38 mm (1 to 1-1/2 inches) above the top of the pan. Using a spatula or trowel, round the top of the mixture so that it is at the interior edges of the pan but not on the top rim of the pan. Use the metal plate or plywood to consolidate the asphaltic concrete mixture. If necessary, this can be done by placing the pan on a clean floor, placing the metal plate or plywood on top of the sample pan and standing on the plate. If desired, a piece of waxed paper or other suitable material may be placed on the top of the mix to avoid sticking. If necessary, fill any voids and consolidate the mix with the metal plate of plywood until the pan is
completely full and the mix is even with the top of the pan. Determine and record the weight of this asphaltic concrete calibration sample to the nearest gram. This weight is identified as the asphaltic concrete calibration sample weight and is used (± 5 grams) for the remaining calibration samples.

(i) Place the sample in the gauge and take a 16-minute count. Record the measured count.

(j) Repeat steps in paragraphs 5(h) and 5(i) to fabricate and test the remaining calibration samples. These samples shall be fabricated in the same manner as the first calibration sample and shall be within ± 5 grams of the asphaltic concrete calibration sample weight. If samples are not able to be tested immediately after fabrication, they shall be placed in an oven at 121 to 149 °C (250 to 300 °F) until they can be tested.

(k) Prepare a calibration curve in accordance with the manufacturer's calibration instructions. On some gauges this curve may be generated internal to the gauge. To be considered acceptable, a calibration curve should have a correlation coefficient greater than or equal to 0.995.

\[
\text{Correlation Coefficient} = \sqrt{1 - \frac{\sum (y - y')^2}{\sum (y - \bar{y})^2}}
\]

Where: 
- \( y \) = Actual percent bituminous material value for each sample.
- \( y' \) = Calculated percent bituminous material value for each sample, from curve.
- \( \bar{y} \) = Mean of the actual percent bituminous material values for all samples.

(l) Remove the dry aggregate blank sample from the oven. Place in the gauge take a minimum four-minute count. This count is recorded and may be used to determine if significant changes occur in the aggregate during asphaltic concrete production. [See paragraph 6(i).]

(m) Once a calibration has been performed for the asphaltic concrete mixture, other gauges from the same manufacturer may use a cross calibration for testing of the same asphaltic concrete mixture. Cross calibrations shall be performed in accordance with the gauge manufacturer's recommendations, and shall have been done within the preceding 12 months. In cases of dispute, the gauge derived calibration shall be the referee method. A new cross calibration is required after any gauge repair.
PROCEDURE (FIELD TEST SAMPLE)

6. (a) A 16-minute background count shall be taken and recorded each day before any test samples are run.

(b) Obtain a representative sample for determining the bituminous material content. Generally 8 kg (18 pounds) should be adequate.

(c) If desired, the entire sample for determining the bituminous material content may be dried to constant weight in an oven at 143 ± 6 °C (290 ± 10 °F) in lieu of obtaining a moisture content sample as described in paragraph (d) or (h) below. If a moisture sample is obtained, the percent moisture is determined in accordance with Arizona Test Method 406. The percent moisture is subtracted from the gauge percent bituminous material to obtain the corrected (actual) percent bituminous material.

(d) If the entire sample for determining the bituminous material content has not been dried to constant weight as described in paragraph (c) above, or the moisture content sample is not to be taken at the completion of testing of the bituminous material content sample as described in paragraph (h) below, a moisture content sample is taken immediately after the fabrication of the bituminous material content test sample as described in paragraph (e) below.

(e) If the field sample is at fabrication temperature at the time of obtaining the bituminous material content test sample, the bituminous material content test sample is fabricated immediately. If the field sample is not at fabrication temperature, enough material for both the bituminous material content test sample, and a moisture content sample when necessary, shall be obtained and placed in a single pan and brought to fabrication temperature.

(f) At the time of beginning the fabrication of the bituminous material test sample, the mixture shall be at a temperature of 121 to 149 °C (250 to 300 °F). Fill the tared sample pan in approximately equal depth layers, using the same number of layers as was used in fabrication of the calibration samples. Uniformly distribute each layer of the sample in the pan. After placing each layer in the pan, gently spade the mix with a spatula, wide blade-like tool, or scoop and lightly press the mix down. When the pan is nearly full, weigh the sample. Add enough mix until the sample weight is within ± 10 grams of the asphaltic concrete calibration sample weight as determined in paragraph 5(h). If necessary, the mix may be pressed down using the metal plate or plywood during addition of material. Using a spatula or trowel, round the top of the mixture so that it is at the interior edges of the pan but not on the top rim of the pan. Use the metal plate or plywood to consolidate the asphaltic concrete mixture until it is even with the top rim of the pan. If necessary, this can be done by placing the pan on a
clean floor, placing the metal plate or plywood on top of the sample pan and standing on the plate. If desired, a piece of waxed paper or other suitable material may be placed on the top of the mix to avoid sticking. Reweigh the sample, and assure the weight is within $\pm 10$ grams of the asphaltic concrete calibration sample weight. Record the weight of the test sample to at least the nearest 5 grams.

(g) Immediately after fabrication place the sample in the gauge. Take and record a minimum four-minute count, and determine the gauge percent bituminous material. If an increased count time (8-minute or 16-minute) was used to determine the variability of the gauge [paragraph 2(a)(5)], that count time is the minimum that is to be used.

(h) If the bituminous material content sample has not been dried to constant weight prior to testing, or if the moisture content sample has not been taken at the time of fabrication of the bituminous material content test sample, a representative 1000 ± 50 gram moisture content sample shall be immediately taken from the bituminous material content test sample upon determination of the bituminous material percent in the gauge.

NOTE: Generally the samples taken for bituminous material content and moisture content are obtained from a larger field sample from which material for other tests, e.g., Marshall and Rice is also split out. In case there is a need for additional testing, it may be desirable to save material from the bituminous material content test sample along with excess material from the field sample.

(i) If desired during asphaltic concrete production, a dry aggregate blank sample may be prepared and tested to ensure that changes in aggregate do not occur unnoticed. Testing may be performed at any time that a change in the aggregate is suspected. This sample shall be dried to constant weight and fabricated in the manner described in paragraph 5(c). The weight of this sample shall be within $\pm 10$ grams of the weight of the dry aggregate blank sample as determined in paragraph 5(c). Place sample in the gauge and take and record a minimum four-minute count. If an increased count time (8-minute or 16-minute) was used to determine the variability of the gauge [paragraph 2(a)(5)], that count time is the minimum that is to be used. If a significant change is noted in this count [greater than $\pm 1.0\%$ of the calibration blank count determined in paragraph 5(i)], a new calibration may be warranted. The project supervisor should be notified immediately of a possible change in aggregate, pit conditions, and/or moisture retained in the aggregate.
MODIFICATIONS FOR ASPHALTIC CONCRETE FRICTION COURSE, ASPHALTIC CONCRETE (ASPHALT-RUBBER), AND ASPHALTIC CONCRETE FRICTION COURSE (ASPHALT-RUBBER)

7. The following modifications apply for Asphaltic Concrete Friction Course, Asphaltic Concrete (Asphalt-Rubber), and Asphaltic Concrete Friction Course (Asphalt-Rubber).

(a) In paragraph 3(d), the last sentence is changed to read: If the original calibration specimens are retested, they shall be heated in an oven at 104 ± 11 °C (220 ± 20 °F) for Asphaltic Concrete Friction Course, or at 149 ± 14 °C (300 ± 25 °F) for Asphaltic Concrete (Asphalt-Rubber) or Asphaltic Concrete Friction Course (Asphalt-Rubber), for at least one hour to drive out any condensation or absorbed moisture and to bring them to testing temperature.

(b) In the calibration procedure given in Section 5, the following changes are made:

1) In paragraph 5(g), a “butter mix” shall be used for preparing calibration samples.

2) In paragraph 5(h):

   a) At the time of beginning the fabrication of calibration samples for Asphaltic Concrete Friction Course, the temperature of the mixture shall be 104 ± 11 °C (220 ± 20 °F).

   b) At the time of beginning the fabrication of calibration samples for Asphaltic Concrete (Asphalt-Rubber) or Asphaltic Concrete Friction Course (Asphalt-Rubber), the temperature of the mixture shall be 149 ± 14 °C (300 ± 25 °F).

   c) For Asphaltic Concrete Friction Course or Asphaltic Concrete Friction Course (Asphalt-Rubber), instead of adding mix until the top of the material is mounded approximately 25 to 38 mm (1 to 1-1/2 inches) above the top of the pan, mix shall be added so that the top of the mix is mounded slightly above the top of the pan.

3) In paragraph 5(j), the last sentence is changed to read: If samples are not able to be tested immediately after fabrication, they shall be placed in an oven at 104 ± 11 °C (220 ± 20 °F) for Asphaltic Concrete Friction Course, or at 149 ± 14 °C (300 ± 25 °F), for Asphaltic Concrete (Asphalt-Rubber) or Asphaltic Concrete Friction Course (Asphalt-Rubber).
(c) Paragraph 4 (b) is changed to read: When sampling Asphaltic Concrete Friction Course or Asphaltic Concrete Friction Course (Asphalt-Rubber), an adequate amount of material shall be taken from the truck at the mixing plant and placed into a 19 liter (5 gallon) bucket, or other suitable container, which has been conditioned with a "butter mix". The sample shall be taken at random locations, approximately 300 mm (12 inches) below the surface, within 5 minutes from the time the loading of the truck is completed. Representative samples of Asphaltic Concrete (Asphalt-Rubber) may be obtained either from the truck at the mixing plant, the windrow, or in accordance with Arizona Test Method 104.

(d) In the procedure for testing field samples (Section 6), the following changes are made:

1) At the beginning of the fabrication of field samples of Asphaltic Concrete Friction Course, the temperature of the mixture shall be 104 ± 11 °C (220 ± 20 °F).

2) At the beginning of the fabrication of field samples of Asphaltic Concrete (Asphalt-Rubber) or Asphaltic Concrete Friction Course (Asphalt-Rubber), the temperature of the mixture shall be 149 ± 14 °C (300 ± 25 °F).

CALIBRATION REPORT

8. The calibration report shall contain, as a minimum, the following information:

(a) Make, model, and serial number of the asphalt content gauge.

(b) ADOT project number.

(c) Type of mix.

(d) Calibration number.

(e) Name of test operator.

(f) Identification (type and source) of bituminous material and aggregate materials, (and mineral admixture when used).

(g) Calibration date and background count.

(h) Correlation coefficient of the calibration curve (Fit Coeff.).
(i) Weight and measured count of the dry aggregate blank sample prepared and tested during the calibration procedure.

(j) Mix design percent bituminous material content.

(k) Percent of mineral admixture if used.

(l) Percentages of bituminous material in calibration samples.

(m) Weight of asphaltic concrete calibration samples.

(n) Measured count of each asphaltic concrete calibration sample.

FIELD SAMPLE TEST REPORT

9. The field sample test report shall, as a minimum, contain the following information:

(a) ADOT project number.

(b) Type of mix.

(c) Calibration number.

(d) Mix design percent bituminous material content.

(e) Date and location of field test sample.

(f) Test number and lot number, if applicable.

(g) Field sample test date and name of test operator.

(h) Background count.

(i) Weight of test sample.

(j) Count time (minutes) of test.

(k) Measured count of test sample.

(l) Gauge measured percent bituminous material.
(m) Percent moisture.

(n) Corrected (actual) percent bituminous material.

(o) When a dry aggregate blank sample is prepared and tested, the following information is also reported:

1) Weight and measured count of the dry aggregate blank sample prepared and tested during the calibration procedure.

2) Weight and measured count of dry aggregate blank sample prepared and tested during production.

3) Percentage change in counts from the calibration dry aggregate blank sample count.
COMPACTION AND TESTING OF
BITUMINOUS MIXTURES UTILIZING 152.4 mm
(SIX INCH) MARSHALL APPARATUS

(A Modification of ASTM D 5581)

SCOPE

1. (a) This method covers the procedure for compacting and testing bituminous mixtures utilizing 152.4 mm (six inch) Marshall apparatus.

   (b) This procedure is used for bituminous mixtures with a mix design gradation target of less than 85% passing the 19.0 mm (3/4 inch) sieve.

   (c) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

   (d) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (e) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:
(a) Compaction Mold Assembly [152.4 mm (six inch) diameter] - Cylindrical mold, baseplate, and extension collar; constructed of steel and conforming to the requirements of Figure 1. (Three compaction mold assemblies are normally utilized.)

(b) Specimen Extruding Device - Extrusion jack or press for extruding specimens from molds.

(c) Compaction Hammer.

1) A mechanical compaction hammer having a flat, circular tamping face with a nominal diameter of 149.4 mm (5-7/8 inches), and a 10.21 ± 0.01 kg (22.50 ± 0.02 pound) sliding weight with a free fall of 457.2 ± 2.5 mm (18.0 ± 0.1 inches).

2) For calibration purposes, the density of a 152.4 mm (six inch) diameter specimen of a 19 mm (3/4") AC mix compacted with the 152.4 mm (six inch) mechanical hammer utilizing approximately 112 blows per side shall not vary by more than 16.0 kg/m³ (1.0 lb./cu. ft.) from the density of a 101.6 mm (four inch) diameter specimen of the same mix compacted with a compaction hammer meeting the requirements of Arizona Test Method 410. The number of blows may be varied from 112 in order to obtain calibration. If changes or adjustments to the compaction hammer or pedestal are made, a recalibration may be warranted.

(d) Compaction pedestal - The compaction pedestal shall consist of a 203.2 mm by 203.2 mm by 457.2 mm (8" x 8" x 18") wooden post capped with a 304.8 mm by 304.8 mm by 25.4 mm (12" x 12" x1") steel plate. The steel cap shall be firmly fastened to the post. The wooden post shall have a dry weight of 673 to 769 kg/m³ (42 to 48 lbs./cu. ft.) and shall rest squarely on, and be firmly secured to, a solid concrete slab. The pedestal assembly shall be installed so that the post is plumb and the cap is level.

(e) Specimen Mold Holder - Mounted on the compaction pedestal so as to center the compaction mold over the center of the post. It shall hold the compaction mold, collar, and base plate securely in position during compaction of specimen.

(f) Oven for heating bituminous mixtures and specimen mold assemblies at required temperature.

(g) Hot plate for heating compaction hammer, spoon, and spatula.
(h) A flat spatula with blade approximately 25 mm (1 inch) wide and at least 150 mm (6 inches) long, stiff enough to penetrate the entire bituminous mixture.

(i) Thermometers for determining temperatures of bituminous mixtures. Armored glass thermometers or dial-type with metal stem with a range of 10 to 200 °C (50 to 400 °F) are recommended. Thermometers shall have increments of not greater than 2.8 °C (5 °F).

(j) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least one gram.

(k) Large spoon for placing mixtures in specimen molds.

(l) If Marshall stability and flow are to be determined, the following additional apparatus is required:

1) Breaking head consisting of upper and lower cylindrical segments or test heads. The lower segment shall be mounted on a base having two perpendicular guide rods or posts extending upward. Guide sleeves in the upper segment shall be in such a position as to direct the two segments together without appreciable binding or loose motion the guide rods. When a 152.4 ± 3.1 mm (6.000 ± 0.120 inch) diameter by 100 mm (4 inch) thick metal block is placed between the two segments, the breaking head shall conform to the requirements of Figure 2.

2) Water bath of sufficient depth to provide for the complete immersion of specimens and thermostatically controlled so as to maintain the bath at 60 ± 1 °C (140° ± 1.8 °F). The tank shall have a perforated false bottom or be equipped with a shelf for supporting specimens 50.8 mm (2 inches) above the bottom of the bath.

3) Marshall stability and flow testing apparatus, with operating instruction manual. The apparatus shall be capable of applying a load with a constant rate of travel of 50.8 ± 2.5 mm (2.0 ± 0.1 inches) per minute.

4) Height gauge capable of measuring the height of specimens to the nearest 0.1 mm (0.001 inch).
PROCEDURE

3 (a) A set of three 4000 gram (approximate) representative samples of bituminous mixture shall be obtained. The weight utilized will be such that compacted specimens for Marshall stability and flow testing meet the height requirement of 88.9 to 101.6 mm (3.500 to 4.000 inches).

(b) Before placing in the mold, the mixture and a mold assembly (baseplate, mold and collar) shall be at approximately 143 °C (290 °F). The face of the compaction hammer shall be thoroughly cleaned and heated on a hot plate set at approximately 143 °C (290 °F). The temperature of the laboratory during compaction of the specimens shall be between 20 and 30 °C (68 and 86 °F).

(c) Place a 152.4 mm (6 inch) paper disc in the bottom of the mold before the mixture is introduced. Place approximately one-half of the batch in the mold with a heated spoon. Spade the mixture vigorously, penetrating the entire mixture, with the heated spatula 15 times around perimeter and 10 times over the interior. Place the second half of the batch in the mold and repeat the spading procedure. Smooth the surface of the mix to a slightly rounded shape.

(d) Unless otherwise indicated on the mix design, the compaction temperature range (except for recycle mixes and bituminous mixtures containing asphalt-rubber) shall be as shown in Table 1 below for the specified type of bituminous material. For asphalt-rubber mixes, the compaction temperature range shall be 163 ± 6 °C (325 ± 10 °F). For recycle mixes, the compaction temperature range shall be 141 ± 6 °C (285 ± 10 °F).

<table>
<thead>
<tr>
<th>Bituminous Material</th>
<th>Compaction Temperature Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC 10, AC 20</td>
<td>138 ± 6 °C (280 ± 10 °F)</td>
</tr>
<tr>
<td>AC 30, AC 40</td>
<td>141 ± 6 °C (285 ± 10 °F)</td>
</tr>
<tr>
<td>PG 58-XX, PG 64-XX</td>
<td>138 ± 6 °C (280 ± 10 °F)</td>
</tr>
<tr>
<td>PG 70-XX, PG 76-XX</td>
<td>141 ± 6 °C (285 ± 10 °F)</td>
</tr>
<tr>
<td>Polymer Modified Asphalts</td>
<td>143 ± 6 °C (290 ± 10 °F)</td>
</tr>
</tbody>
</table>

(e) If necessary, the mixture and mold shall be returned to an oven at the required temperature for the minimum time necessary to achieve the required compaction temperature.
(f) Place a 152.4 mm (six inch) paper disc on top of material, place the mold assembly on the compaction pedestal in the mold holder, and apply the calibrated number of blows (approximately 112) with the mechanical compaction hammer. Compaction shall be performed at a minimum rate of 40 blows per minute. The compaction hammer shall apply only one blow with each fall, that is, there shall not be a rebound impact. Remove the base plate and collar, and reverse and reassemble the mold. Apply the calibrated number of compaction blows to the face of the reversed specimen.

(g) Remove collar, baseplate, and paper discs, and allow specimen to cool. Cooling may be accomplished at room temperature, in a 25 °C (77 °F) air bath, or if more rapid cooling is desired the mold and specimen may be placed in front of a fan until cool.

(h) Extrude the specimen from the mold. Care shall be taken in extruding the specimen from the mold, so as not to develop tensile stresses in the specimen or tear the sides of the specimen.

SPECIMEN TESTING

4. Specimens prepared as above shall be tested by some or all of the following procedures, as applicable due to testing requirements and availability of apparatus.

(a) If Marshall stability and flow are to be determined, measure height of specimens to the nearest 0.1 mm (0.001 inch). Prior to measurement of height, excess material shall be brushed from the edges of the specimens. Compacted specimens shall be 88.9 to 101.6 mm (3.500 to 4.000 inches) in height. If this criteria is not met, the entire set of specimens shall be discarded and a new set prepared after necessary adjustments in sample weight have been made.

(b) Determine the specific gravity of the specimens in accordance with Arizona Test Method 415, Method A. (Assume specimen is at constant weight after cooling.)

(c) Determine the bulk density of each of the specimens, by multiplying the respective specific gravity by 998 kg/m³ (62.3 lbs./cu. ft.). Record the individual bulk densities to the nearest 1 kg/m³ (0.1 lb./cu. ft.). The densities of the three specimens shall not differ by more than 48 kg/m³ (3.0 lbs./cu. ft.). If this density
requirement is not met, the entire set of specimens shall be discarded and a new set of specimens prepared.

(d) Determine the average specific gravity of the specimens and record to the nearest 0.001. Calculate the average bulk density of the specimens, by multiplying the average specific gravity by 998 kg/m³ (62.3 lbs./cu. ft.). Record the average bulk density to the nearest 1 kg/m³ (0.1 lb./cu. ft.).

(e) If the stability and flow are to be determined, the steps in paragraphs (f) through (l) below are followed, utilizing apparatus in accordance with the operating instructions for that apparatus.

(f) Bring the specimens to 60 ± 1 °C (140° ± 1.8 °F) by immersing in the water bath 45 to 60 minutes. Prior to testing, it shall be assured that the inside of the test heads are clean, and that the guide rods are clean and lubricated so that the upper test head slides freely over them.

(g) The breaking head temperature shall be maintained between 21 to 38 °C (70 to 100 °F), using a water bath when required. Remove the specimen from the water bath, quickly towel dry specimen and place in the lower segment of the breaking head. Place the upper segment of the breaking head on the specimen, and place the complete assembly in position on the testing machine.

(h) Apply the load to the specimen with a constant rate of 50.8 ± 2.5 mm (2.0 ± 0.1 inches) per minute until the maximum load is reached and the load decreases. The maximum load is defined as the last point in the load/time curve before the load decreases. The elapsed time for the test from removal of the test specimen from water bath to maximum load determination shall not exceed 30 seconds.

(i) Record the stability of each specimen to the nearest 50 newtons (10 pounds force), and the flow to the nearest 0.1 mm (0.01 inch).

(j) Correct the stability obtained for each specimen, for the height of the specimen, by the table in Figure 3. Record the corrected stability to the nearest 50 newtons (10 pounds force).

(k) Determine and record the average corrected stability to the nearest 50 newtons (10 pounds force), and the average flow to the nearest 0.1 mm (0.01 inch).

(l) The average stability shall be corrected to correspond to a 101.6 mm (four inch) specimen by dividing the average test result by 2.0.
All dimensions are nominal, except where tolerances are indicated.

152.4 mm (Six Inch) Compaction Mold, Extension Collar, and Baseplate

FIGURE 1
A  50 mm (2 in.)
B  22 mm (0.87 in.) minimum
C  66.7 ± 1.3 mm (2.625 ± 0.050 in.)
D  19.0 ± 0.5 mm (0.750 ± 0.020 in.)
E  152.4 ± 3.1 mm (6.000 ± 0.120 in.)
F  13.5 ± 0.3 mm (0.53 ± 0.01 in.)
G  45°
H  152.4 (6 in.) nominal
I  108 mm (4.25 in.)

Breaking Head

FIGURE 2
<table>
<thead>
<tr>
<th>Height of Specimen (millimeters)</th>
<th>Height of Specimen (inches)</th>
<th>Correlation Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>88.9 - 89.1</td>
<td>3.500 - 3.511</td>
<td>1.12</td>
</tr>
<tr>
<td>89.2 - 89.6</td>
<td>3.512 - 3.532</td>
<td>1.11</td>
</tr>
<tr>
<td>89.7 - 90.2</td>
<td>3.533 - 3.553</td>
<td>1.10</td>
</tr>
<tr>
<td>90.3 - 90.7</td>
<td>3.554 - 3.574</td>
<td>1.09</td>
</tr>
<tr>
<td>90.8 - 91.2</td>
<td>3.575 - 3.594</td>
<td>1.08</td>
</tr>
<tr>
<td>91.3 - 91.7</td>
<td>3.595 - 3.615</td>
<td>1.07</td>
</tr>
<tr>
<td>91.8 - 92.3</td>
<td>3.616 - 3.636</td>
<td>1.06</td>
</tr>
<tr>
<td>92.4 - 92.8</td>
<td>3.637 - 3.657</td>
<td>1.05</td>
</tr>
<tr>
<td>92.9 - 93.3</td>
<td>3.658 - 3.678</td>
<td>1.04</td>
</tr>
<tr>
<td>93.4 - 93.9</td>
<td>3.679 - 3.699</td>
<td>1.03</td>
</tr>
<tr>
<td>94.0 - 94.4</td>
<td>3.700 - 3.719</td>
<td>1.02</td>
</tr>
<tr>
<td>94.5 - 94.9</td>
<td>3.720 - 3.739</td>
<td>1.01</td>
</tr>
<tr>
<td>95.0 - 95.5</td>
<td>3.740 - 3.760</td>
<td>1.00</td>
</tr>
<tr>
<td>95.6 - 96.0</td>
<td>3.761 - 3.782</td>
<td>0.99</td>
</tr>
<tr>
<td>96.1 - 96.5</td>
<td>3.783 - 3.803</td>
<td>0.98</td>
</tr>
<tr>
<td>96.6 - 97.2</td>
<td>3.804 - 3.831</td>
<td>0.97</td>
</tr>
<tr>
<td>97.3 - 97.9</td>
<td>3.832 - 3.858</td>
<td>0.96</td>
</tr>
<tr>
<td>98.0 - 98.6</td>
<td>3.859 - 3.886</td>
<td>0.95</td>
</tr>
<tr>
<td>98.7 - 99.2</td>
<td>3.887 - 3.907</td>
<td>0.94</td>
</tr>
<tr>
<td>99.3 - 99.7</td>
<td>3.908 - 3.928</td>
<td>0.93</td>
</tr>
<tr>
<td>99.8 - 100.4</td>
<td>3.929 - 3.956</td>
<td>0.92</td>
</tr>
<tr>
<td>100.5 - 101.1</td>
<td>3.957 - 3.983</td>
<td>0.91</td>
</tr>
<tr>
<td>101.2 - 101.6</td>
<td>3.984 - 4.000</td>
<td>0.90</td>
</tr>
</tbody>
</table>

* The measured stability of a specimen multiplied by the correlation ratio for the height of the specimen equals the corrected stability for a 95.25 mm (3-1/2 inch) specimen.

FIGURE 3
1. SCOPE

1.1 This procedure is used to determine the air voids in compacted bituminous mixtures. It is applicable for specimens which are either laboratory compacted or field compacted (for example, cores).

1.2 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. CALCULATION

2.1 For specimens which are either Marshall laboratory compacted or field compacted (e.g., cores), the percent air voids shall be calculated using the bulk density of the compacted bituminous mixture (Arizona Test Method 415) and maximum density of the mixture from the Rice Test (Arizona Test Method 417).

2.1.1 The percent air voids are calculated by the following equation:

\[
\text{Percent Air Voids} = \left( 1 - \frac{\text{Bulk Density}}{\text{Maximum Density}} \right) \times 100
\]

2.1.1.1 An example of the calculations is given in Figure 1.

2.1.1.2 A blank form for performing the calculations is given in Figure 3.

2.2 For specimens which are gyratory laboratory compacted, the percent air voids shall be calculated using the average relative density of the compacted bituminous mixture at \(N_{\text{design}}\) (AASHTO T 312).
2.2.1 The percent air voids are calculated by the following equation:

\[
\text{Percent Air Voids} = (100) - \text{(Average Relative Density, } \% \text{ G}_{\text{mm}, \text{ at } N_{\text{design}}})
\]

2.2.1.1 An example of the calculations is given in Figure 2.

2.2.1.2 A blank form for performing the calculations is given in Figure 4.

3. REPORT

3.1 The percent air voids shall be reported to the nearest 0.1%.
CALCULATION OF AIR VOIDS FOR MARSHALL LABORATORY COMPACTED SPECIMENS OR FIELD COMPACTED SPECIMENS

Specimens Compacted by: Hand ☑, Mechanical ☑, 4 inch ☑, 6 inch ☐; Core ☐;

<table>
<thead>
<tr>
<th>Specimen I.D.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specimen Height</td>
<td>2.516</td>
<td>2.515</td>
<td>2.519</td>
<td></td>
</tr>
</tbody>
</table>

Bulk Specific Gravity, Bulk Density, and Absorption (Arizona Test Method 415: Method A ☑, Method C ☐, or Vacuum Method ☐)

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>1.65</th>
<th>1.68</th>
<th>1.674</th>
</tr>
</thead>
<tbody>
<tr>
<td>A = Mass in grams of specimen in Air</td>
<td>1155.9</td>
<td>1155.4</td>
<td>1158.2</td>
<td></td>
</tr>
<tr>
<td>B = Mass in grams of SSD specimen in Air</td>
<td>1156.9</td>
<td>1156.3</td>
<td>1156.5</td>
<td></td>
</tr>
<tr>
<td>C = Mass in grams of specimen in Water</td>
<td>647.9</td>
<td>649.6</td>
<td>651.8</td>
<td></td>
</tr>
<tr>
<td>$G_{mb} =$ Bulk Specific Gravity = $A/(B – C)$</td>
<td>2.271</td>
<td>2.280</td>
<td>2.283</td>
<td>2.278</td>
</tr>
<tr>
<td>% Absorption = $[(B – A)/(B – C)] \times 100$</td>
<td>0.20</td>
<td>0.18</td>
<td>0.20</td>
<td></td>
</tr>
<tr>
<td>Bulk Density = $(G_{mb} \times 62.3 \text{ lbs./cu. ft.})$</td>
<td>141.5</td>
<td>142.0</td>
<td>142.2</td>
<td></td>
</tr>
<tr>
<td>Range of Bulk Density values (lbs./cu. ft.)</td>
<td>0.7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average Bulk Density = (Average $G_{mb}$ $\times$ 62.3 lbs./cu. ft)</td>
<td></td>
<td></td>
<td></td>
<td>141.9</td>
</tr>
<tr>
<td>Maximum Density (lbs./cu. ft.) [from Rice Test]</td>
<td></td>
<td></td>
<td></td>
<td>149.4</td>
</tr>
</tbody>
</table>

Notes:
- The individual specimen heights are reported to the nearest 0.001 inch.
- The individual specimen masses are reported to the nearest 0.1 gram.
- The individual bulk specific gravities are reported to the nearest 0.001.
- The average bulk specific gravity is calculated, and reported to the nearest 0.001, using the individual bulk specific gravities which have been reported to the nearest 0.001.
- The individual bulk densities are reported to the nearest 0.1 lb./cu. ft.
- The average bulk density is reported to the nearest 0.1 lb./cu. ft.
- The maximum density [from Rice Test] is reported to the nearest 0.1 lb./cu. ft.

Percent Air Voids $= \left[ 1 - \frac{\text{Average Bulk Density}}{\text{Maximum Density from Rice Test}} \right] \times 100 = \left[ 1 - \frac{141.9}{149.4} \right] \times 100 = 5.0\%$

EXAMPLE AIR VOIDS CALCULATION FOR MARSHALL LABORATORY COMPACTED SPECIMENS

FIGURE 1
## CALCULATION OF AIR VOIDS FOR GYRATORY LABORATORY COMPACTED SPECIMENS

<table>
<thead>
<tr>
<th>Specimen I.D.</th>
<th>1</th>
<th>2</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>( h_{\text{ini}} )  = Height, in mm, of specimen at ( N_{\text{ini}} ) (8 gyrations)</td>
<td>128.7</td>
<td>129.3</td>
<td></td>
</tr>
<tr>
<td>( h_{\text{des}} )  = Height, in mm, of specimen at ( N_{\text{des}} ) (100 gyrations)</td>
<td>117.0</td>
<td>117.4</td>
<td></td>
</tr>
<tr>
<td>( h_{\text{max}} )  = Height, in mm, of specimen at ( N_{\text{max}} ) (160 gyrations)</td>
<td>115.6</td>
<td>116.0</td>
<td></td>
</tr>
</tbody>
</table>

### Bulk Specific Gravity and Absorption (Arizona Test Method 415: Method A, Method C, or Vacuum Method)

<table>
<thead>
<tr>
<th></th>
<th>Specimen</th>
<th>Specimen</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>( A ) = Mass, in grams, of specimen at ( N_{\text{max}} ) in Air</td>
<td>4747.4</td>
<td>4744.6</td>
<td></td>
</tr>
<tr>
<td>( B ) = Mass, in grams, of SSD specimen at ( N_{\text{max}} ) in Air</td>
<td>4759.4</td>
<td>4756.0</td>
<td></td>
</tr>
<tr>
<td>( C ) = Mass, in grams, of specimen at ( N_{\text{max}} ) in Water</td>
<td>2752.7</td>
<td>2751.2</td>
<td></td>
</tr>
<tr>
<td>( G_{\text{mb}} ) = Bulk Specific Gravity of specimen at ( N_{\text{max}} )</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( % \text{ Absorption} )</td>
<td>((B - A)/(B - C)) x 100</td>
<td>0.60</td>
<td>0.57</td>
</tr>
<tr>
<td>( G_{\text{mm}} ) = Maximum Specific Gravity [from Rice Test]</td>
<td>2.366</td>
<td>2.367</td>
<td></td>
</tr>
<tr>
<td>*Relative Density, % ( G_{\text{mm}} ), of specimen at ( N_{\text{ini}} )</td>
<td>86.8</td>
<td>86.7</td>
<td>86.8</td>
</tr>
<tr>
<td>*Relative Density, % ( G_{\text{mm}} ), of specimen at ( N_{\text{des}} )</td>
<td>95.5</td>
<td>95.5</td>
<td>95.5</td>
</tr>
<tr>
<td>*Relative Density, % ( G_{\text{mm}} ), of specimen at ( N_{\text{max}} )</td>
<td>96.6</td>
<td>96.7</td>
<td>96.7</td>
</tr>
</tbody>
</table>

### Notes:
- The Individual specimen heights are reported to the nearest 0.1 mm.
- The Individual specimen masses are reported to the nearest 0.1 gram.
- The Individual bulk specific gravities are reported to the nearest 0.001.
- The maximum specific gravity [from Rice Test] is reported to the nearest 0.001.
- The individual relative densities are reported to the nearest 0.1 percent.
- The average relative density for each set of specimens (at \( N_{\text{ini}}, N_{\text{des}}, \) and \( N_{\text{max}} \)) is calculated, and reported to the nearest 0.1 percent, using the corresponding individual relative densities which have been reported to the nearest 0.1 percent.
- Three specimens are used when referee testing is performed.

### Percent Air Voids

\[
\text{Percent Air Voids} = (100) - \text{Average Relative Density, } \% G_{\text{mm}}, \text{ at } N_{\text{des}}
\]

\[
= (100) - (95.5) = 4.5\%
\]

**EXAMPLE AIR VOIDS CALCULATION FOR GYRATORY LABORATORY COMPACTED SPECIMENS**

**FIGURE 2**
### CALCULATION OF AIR VOIDS FOR MARSHALL LABORATORY COMPACTED SPECIMENS OR FIELD COMPACTED SPECIMENS

<table>
<thead>
<tr>
<th>Specimens Compacted by:</th>
<th>Hand</th>
<th>Mechanical</th>
<th>4 inch</th>
<th>6 inch</th>
<th>Core</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specimen I.D.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Specimen Height</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Bulk Specific Gravity, Bulk Density, and Absorption (Arizona Test Method 415: Method A ☑️, Method C ☐️, or Vacuum Method ☐️)

\[
A = \text{Mass in grams of specimen in Air}
\]

\[
B = \text{Mass in grams of SSD specimen in Air}
\]

\[
C = \text{Mass in grams of specimen in Water}
\]

\[
G_{mb} = \text{Bulk Specific Gravity} = \frac{A}{B - C}
\]

\[
\% \text{ Absorption} = \frac{[(B - A)/(B - C)] \times 100}{100}
\]

\[
\text{Bulk Density} = (G_{mb} \times 62.3 \text{ lbs./cu. ft.})
\]

\[
\text{Range of Bulk Density values (lbs./cu. ft.)}
\]

\[
\text{Average Bulk Density} = (\text{Average } G_{mb} \times 62.3 \text{ lbs./cu. ft})
\]

\[
\text{Maximum Density (lbs./cu. ft.) [from Rice Test]}
\]

**Notes:**

- The individual specimen heights are reported to the nearest 0.001 inch.
- The individual specimen masses are reported to the nearest 0.1 gram.
- The individual bulk specific gravities are reported to the nearest 0.001.
- The average bulk specific gravity is calculated, and reported to the nearest 0.001, using the individual bulk specific gravities which have been reported to the nearest 0.001.
- The individual bulk densities are reported to the nearest 0.1 lb./cu. ft.
- The average bulk density is reported to the nearest 0.1 lb./cu. ft.
- The maximum density [from Rice Test] is reported to the nearest 0.1 lb./cu. ft.

\[
\text{Percent Air Voids} = \left[1 - \frac{\text{Average Bulk Density}}{\text{Maximum Density from Rice Test}}\right] \times 100 = \left[1 - \left(\frac{(\_\_\_)}{(\_\_\_)}\right)\right] \times 100 = \_\_\_%
\]

**FIGURE 3**
### CALCULATION OF AIR VOIDS FOR GYRATORY LABORATORY COMPACTED SPECIMENS

<table>
<thead>
<tr>
<th>Specimen I.D.</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>( h_{\text{ini}} ) = Height, in mm, of specimen at ( N_{\text{ini}} ) (8 gyrations)</td>
<td></td>
</tr>
<tr>
<td>( h_{\text{des}} ) = Height, in mm, of specimen at ( N_{\text{des}} ) (100 gyrations)</td>
<td></td>
</tr>
<tr>
<td>( h_{\text{max}} ) = Height, in mm, of specimen at ( N_{\text{max}} ) (160 gyrations)</td>
<td></td>
</tr>
</tbody>
</table>

Bulk Specific Gravity and Absorption (Arizona Test Method 415:
Method A \( \square \), Method C \( \square \), or Vacuum Method \( \square \))

\( A = \) Mass, in grams, of specimen at \( N_{\text{max}} \) in Air

\( B = \) Mass, in grams, of SSD specimen at \( N_{\text{max}} \) in Air

\( C = \) Mass, in grams, of specimen at \( N_{\text{max}} \) in Water

\[ G_{\text{mb}} = \text{Bulk Specific Gravity of specimen at } N_{\text{max}} \]

\[ \% \text{ Absorption} = \left( \frac{B - A}{B - C} \right) \times 100 \]

\( G_{\text{mm}} = \text{Maximum Specific Gravity [from Rice Test]} \)

*Relative Density, %\( G_{\text{mm}} \), of specimen at \( N_{\text{ini}} \)

*Relative Density, %\( G_{\text{mm}} \), of specimen at \( N_{\text{des}} \)

*Relative Density, %\( G_{\text{mm}} \), of specimen at \( N_{\text{max}} \)

\[ \% G_{\text{mmx}} = \frac{G_{\text{mb}} \times h_{\text{max}}}{G_{\text{mm}} \times h_{x}} \times 100 \]

Where:

\( % G_{\text{mmx}} = \) Relative Density, %\( G_{\text{mm}} \), of specimen at \( N_{\text{ini}} \), \( N_{\text{des}} \), or \( N_{\text{max}} \)

\( G_{\text{mb}} = \) Bulk Specific Gravity of specimen at \( N_{\text{max}} \)

\( h_{\text{max}} = \) Height, in mm, of specimen at \( N_{\text{max}} \)

\( G_{\text{mm}} = \) Maximum Specific Gravity [from Rice Test]

\( h_{x} = \) Height of specimen, in mm, at \( N_{\text{ini}} \), \( N_{\text{des}} \), or \( N_{\text{max}} \)

Notes:

The Individual specimen heights are reported to the nearest 0.1 mm.

The Individual specimen masses are reported to the nearest 0.1 gram.

The Individual bulk specific gravities are reported to the nearest 0.001.

The maximum specific gravity [from Rice Test] is reported to the nearest 0.001.

The individual relative densities are reported to the nearest 0.1 percent.

The average relative density for each set of specimens (at \( N_{\text{ini}} \), \( N_{\text{des}} \), and \( N_{\text{max}} \)) is calculated, and reported to the nearest 0.1 percent, using the corresponding individual relative densities which have been reported to the nearest 0.1 percent.

Three specimens are used when referee testing is performed.

\[ \% \text{ Air Voids} = (100) - (\text{Average Relative Density, } % G_{\text{mm}}, \text{ at } N_{\text{des}}) \]

\[ = (100) - (\text{_______}) = \text{_______}% \]
ASPHALT BINDER CONTENT
OF ASPHALTIC CONCRETE MIXTURES
BY THE IGNITION FURNACE METHOD

(A Modification of AASHTO T 308)

1. SCOPE

1.1 This procedure describes the method for determining the percent asphalt binder content of asphaltic concrete mixtures, by use of an ignition furnace. The aggregate remaining after ignition can be used for sieve analysis, as indicated in Section 6.

1.1.1 This procedure does not address the use of reclaimed asphalt pavement (RAP) in asphaltic concrete mixtures. See Arizona Test Method 428 when testing is to be performed on asphaltic concrete mixtures containing RAP.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 A listing of subsequent Sections and Figures in this procedure is given below:

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<td>15</td>
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APPARATUS

2. REQUIREMENTS FOR THE FREQUENCY OF EQUIPMENT CALIBRATION AND VERIFICATION

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 Ignition Furnace - a forced-air ignition furnace that heats the sample by the convection method. The furnace must be capable of heating to temperatures up to 538 °C (1000 °F), and able to maintain a given temperature at ± 5 °C (± 9 °F). The furnace shall have an internal weighing system thermally isolated from the furnace chamber and accurate to 0.1 gram. The balance shall be capable of weighing a 3500 gram sample in addition to the sample baskets. A data collection system shall also be included so that the sample mass loss can be automatically determined to an accuracy of 0.1 gram and displayed during a test. The furnace shall provide a printout that includes, as a minimum, the initial sample mass, sample mass loss, 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 one minute intervals. A filter capable of reducing emissions to an acceptable level shall also be incorporated into the furnace. The furnace shall be vented into a hood or to the outside and be set up properly so that there are no noticeable odors escaping into the laboratory. The furnace shall have a fan with the capability to pull air through the furnace to expedite the test and to reduce escape of smoke into the laboratory. The furnace shall be equipped so that the door cannot be opened until testing is complete.

2.1.2 Stainless Steel Perforated Baskets - the baskets shall be an appropriate size that allow the samples to be a thickness which allows air to flow up through and around the sample particles. The sample shall be completely enclosed with screen mesh, perforated stainless steel plate, or other suitable material. Screen mesh or other suitable material with openings of No. 8 has been found to perform well.

2.1.3 Stainless Steel Catch Pan - of sufficient size to hold the sample baskets so that aggregate particles and melting asphalt binder falling through the screen mesh are caught.

2.1.4 Oven(s) - capable of heating to temperatures up to 350 °F, and able to maintain a given temperature at the tolerances specified herein.
2.1.5  Scale(s) or balance(s) - capable of measuring the maximum mass to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.1.6  Safety Equipment - safety glasses or face shield, high temperature gloves, and long sleeve jacket. Additionally, a heat resistant surface capable of withstanding 1200 °F and a protective cage capable of surrounding the sample baskets shall be provided.

2.1.7  Miscellaneous Equipment - a pan larger than the sample basket(s) for transferring samples after ignition, spatulas, bowls, spoons, and wire brushes.

2.1.8  Mixing apparatus - Mechanical mixing is recommended; 20 quart capacity mixer is required. (Hand mixing may be performed if desired.)

2.1.9  Thermometer - with a temperature range of 50 to 500 °F.

2.1.10  Hot plate - capable of heating to temperatures up to 350 °F, and able to maintain a given temperature at ± 5 °F.

2.1.11  For performing sieve analysis, apparatus as specified in Arizona Test Method 201.

3.   SAMPLING

3.1  For preparing calibration samples, obtain representative samples of aggregates in accordance with Arizona Test Method 105. Samples shall be sufficiently large to provide enough material for calibration testing. The samples shall be adequately dried, if necessary, to a free-flowing condition in the portion passing the 4.75 mm (No. 4) sieve.

3.2  For testing field samples of asphaltic concrete, obtain a representative sample of the freshly produced mix in accordance with Arizona Test Method 104. Obtain representative test samples, in accordance with the appropriate sections of Arizona Test Method 416, for the determination of moisture content (if required) and asphalt binder content.

3.2.1  The size of the asphalt binder content test sample shall be within ± 150 grams of the sample size used for calibration and must also conform to the mass requirement shown in Table 1. When the mass of the test sample exceeds the capacity of the equipment used, the test sample shall
be divided into suitable increments, tested, and the results appropriately combined.

<table>
<thead>
<tr>
<th>TABLE 1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Size of Test Sample</strong></td>
</tr>
<tr>
<td><strong>Nominal Maximum Aggregate Size (See Note)</strong></td>
</tr>
<tr>
<td>1-1/2 in.</td>
</tr>
<tr>
<td>1 in.</td>
</tr>
<tr>
<td>3/4 in.</td>
</tr>
<tr>
<td>1/2 in.</td>
</tr>
<tr>
<td>3/8 in.</td>
</tr>
<tr>
<td>No. 4</td>
</tr>
</tbody>
</table>

Note: The smallest sieve opening through which the entire amount of material, by specification, is permitted to pass.

4. **CALIBRATION**

4.1 Asphalt binder content results may be affected by the type of aggregate and binder in the mixture. A calibration factor for the asphalt binder content must be established for each mix design. Certain aggregate types may result in an unusually high calibration factor and erroneous gradation results due to aggregate breakdown. Such mixes should be calibrated and tested at a lower temperature as described in Subsection 4.14.

4.2 Dry the aggregate samples to constant mass at 290 ± 10 °F. Allow the material to cool.

4.3 Screen the aggregate stockpile samples and separate into individual sizes for No. 8 and larger, and minus No. 8 material.

**Note:** In lieu of screening the aggregate sample for each individual stockpile, a bulk-batched sample may be used as described in Subsection 4.3.1.

4.3.1 Using the aggregate stockpile percentages shown in the mix design composite, the material from the individual stockpiles may be bulk-batched in a single sample of an adequate amount of material necessary to prepare the required calibration samples. Screen the bulk-batched material and separate into individual sizes for No. 8 and larger, and minus No. 8 material.
4.4 Using the individual sizes of aggregate for No. 8 and larger, and minus No. 8 material, as obtained either by screening the material from the individual aggregate stockpiles or by screening the the bulk-batched sample, weigh up four aggregate samples representative of the mix design gradation without mineral admixture. These samples will be used for a gradation check, two calibration samples, and a butter mix. The appropriate type and quantity of mineral admixture (by weight of aggregate) shall be added to the aggregate, and thoroughly blended. The weight of the gradation check sample shall conform to the requirements of Table 1. The weight of the two calibration samples and the butter mix shall be such that when the required amount of asphalt binder is added, they conform to the requirements of Table 1.

4.5 Using the aggregate and mineral admixture sample prepared for the gradation check, perform a gradation analysis according to Section 6 to determine the actual gradation. The gradation shall be representative of the mix design gradation with mineral admixture. If the gradation is not representative of the mix design, four new aggregate and mineral admixture samples shall be prepared.

4.6 Using the remaining three aggregate and mineral admixture samples, two calibration samples and a butter mix are prepared as described below at the design asphalt binder content. The asphalt binder grade and type shall be the same as will be used in the asphalt concrete mixture to be tested during production. Heat the samples to the laboratory mixing temperature prescribed in the mix design. (See Note following Subsection 4.7). Allow the samples to cool. Weigh and determine the mass of each sample to the nearest 0.1 gram. If mass is lost during the heating of the samples, do not add make-up material, as this will change the gradation of the samples. The percent asphalt binder content is based on the mass of total mix. For each sample, the weight of asphalt binder to be used is determined by the following:

\[
\text{Weight of Asphalt Binder} = \left( \frac{\text{Weight of Aggregate and Mineral Admixture}}{100 - \text{Percent of Asphalt Binder}} \right) \times \text{Percent of Asphalt Binder}
\]

4.7 All bowls, sample pans, and mixing tools shall be preheated to approximately the laboratory mixing temperature prescribed in the mix design. At the time mixing of the samples begins, the temperature of the asphalt binder, aggregate, and mineral admixture shall be in accordance with the prescribed laboratory mixing temperature ± 5 °F. Each individual
sample shall be thoroughly mixed. All samples shall be mixed at the same mixing temperature ± 5 °F.

**Note:** If the mix design laboratory mixing temperature is not specified, a temperature of 300 ± 5 °F shall be used for mixes which do not use asphalt-rubber, and 325 ± 5 °F for asphalt-rubber mixes.

4.8 Preheat the ignition furnace to 538 ± 5 °C (1000 ± 9 °F), or as modified in Subsection 4.14. Do not preheat the sample basket.

4.9 Weigh and record the mass of the basket assembly to the nearest 0.1 gram.

4.10 The freshly mixed samples may be placed directly in the sample basket assembly. If the samples are allowed to cool, they must be reheated in a 290 ± 10 °F oven for 25 minutes.

4.11 Test samples in accordance with Subsections 5.6 through 5.14.

4.12 If the difference between the measured asphalt binder content of the two samples exceeds 0.07, repeat the test using two additional samples, and from the four results discard the high and the low values.

4.13 Subtract the actual asphalt binder content for each of the two samples from their respective measured asphalt binder content. The asphalt binder content calibration factor is the average of the two resultant values expressed in percent by mass of the asphalt mixture.

4.14 If the asphalt binder content calibration factor exceeds ± 1.0 percent, lower the test temperature to 482 ± 5 °C (900 ± 9 °F) and repeat the test to determine a new calibration factor. If the calibration factor continues to exceed ± 1.0 percent, lower the test temperature to 427 ± 5 °C (800 ± 9 °F) and repeat the test to determine a new calibration factor. Use the calibration factor obtained at 427 ± 5 °C (800 ± 9 °F) even if it exceeds ± 1.0 percent.

4.15 Perform a gradation analysis on the residual aggregate as indicated in Section 6. Subtract the actual percent passing the No. 200 sieve for each sample from the measured percent passing the No. 200 sieve (as determined in Subsection 4.5). Determine the average of the two values. If the resultant average value is greater than ± 0.50, an aggregate gradation correction factor (equal to the resultant average value) for the
passing No. 200 material may be applied to the production field sample test results.

5. PROCEDURE

5.1 The moisture content of the asphaltic concrete shall be determined in accordance with Arizona Test Method 406. The moisture content sample shall be obtained at the same time and subjected to the same treatment prior to testing as the asphalt binder content test sample. As an alternate to performing the moisture determination, the test sample may be dried to a constant mass in an oven at $290 \pm 10 \, ^\circ\text{F}$.

5.2 Preheat the ignition furnace to $538 \pm 5 \, ^\circ\text{C}$ ($1000 \pm 9 \, ^\circ\text{F}$), or to the alternate temperature determined during the calibration (Subsection 4.14). Do not preheat the sample basket. Record the furnace temperature set point prior to the initiation of the test.

5.3 Record the asphalt binder content calibration factor, determined in accordance with Subsections 4.12 through 4.14, for the specific mix to be tested.

5.4 Weigh and record the mass of the basket assembly to the nearest 0.1 gram.

5.5 Obtain the asphalt binder content test sample in accordance with Subsection 3.2, ensuring that the size of the test sample is within $150$ grams of the sample size used for calibration and that the test sample conforms to the requirements shown in Table 1.

5.6 Evenly distribute the test sample over the center of the sample basket(s) and level the material. Use a spatula or trowel to pull material approximately one inch away from the outside edge of basket(s).

5.7 Weigh and record the mass of the sample and basket assembly to the nearest 0.1 gram.

5.8 Calculate and record the initial mass of the sample to the nearest 0.1 gram.

5.9 Set the ignition furnace controller print mode to give a printout of the test data required in Subsection 2.1.1. Input the initial mass of the sample into the ignition furnace controller. Verify that the correct mass has been entered.
5.10 Open the furnace door and place the sample and basket assembly so that it is centered in the chamber. After assuring that the sample basket assembly is not in contact with any wall, close the door. Initiate the test by pressing the start button. This will lock the furnace door and start testing.

5.11 Allow the test to continue until the stable light and audible stable indicator indicates the test is complete. The test is deemed complete when the measured mass loss does not exceed 0.01 percent of the sample mass for three consecutive one minute intervals. If required, press the stop button. This will unlock the furnace door and cause the printer to print out the test results.

5.12 Open the furnace door and remove the sample and basket assembly. Allow the sample to cool 30 ± 5 minutes in the basket assembly. Weigh and record the mass of the sample and basket assembly after ignition to the nearest 0.1 gram.

5.13 Calculate and record the mass of sample after ignition to the nearest 0.1 gram.

5.14 Calculate and record the corrected asphalt binder content of the sample, to the nearest 0.01%, as follows:

\[
\%AC = \left[ \frac{W_i - W_A}{W_i} \times 100 \right] - C_F - %M
\]

Where:
- \%AC = measured (corrected) asphalt binder content in percent by mass of the sample
- \(W_i\) = mass of the sample prior to ignition
- \(W_A\) = mass of the sample after ignition
- \(C_F\) = asphalt binder content calibration factor, percent by mass of the sample
- \%M = percent moisture in the sample

Note: During calibration, \(C_F\) and \%M are zero.

5.14.1 If an ignition furnace correction (tank stab correction) is made, the \%AC determined in Subsection 5.14 is adjusted by that correction.

5.15 Attach the original printed ticket to the back of the card.

5.16 Empty the contents of the baskets into a flat pan. Use a small wire sieve brush to ensure that any residual fines are removed from the
baskets. Take care not to lose any material, as this will affect gradation results.

5.17 If needed, perform a gradation analysis of the residual aggregate according to Section 6.

6. SIEVE ANALYSIS OF AGGREGATE

6.1 If required, the aggregate shall be subjected to sieve analysis as described below. The coarse sieving shall be performed in accordance with Subsection 6.2, and the fine sieving in accordance with Subsection 6.3. The quantity of material on a given sieve at the completion of sieving shall not exceed the amount shown in Table 2.

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Maximum Mass Allowed (grams/sq. in.)</th>
<th>8 inch Diameter Sieve</th>
<th>12 inch Diameter Sieve</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-1/2”</td>
<td>25</td>
<td>---</td>
<td>2827</td>
</tr>
<tr>
<td>1”</td>
<td>18</td>
<td>---</td>
<td>2036</td>
</tr>
<tr>
<td>3/4”</td>
<td>14</td>
<td>---</td>
<td>1583</td>
</tr>
<tr>
<td>1/2”</td>
<td>10</td>
<td>---</td>
<td>1131</td>
</tr>
<tr>
<td>3/8”</td>
<td>8</td>
<td>---</td>
<td>905</td>
</tr>
<tr>
<td>1/4”</td>
<td>6</td>
<td>---</td>
<td>679</td>
</tr>
<tr>
<td>No. 4</td>
<td>5</td>
<td>---</td>
<td>565</td>
</tr>
<tr>
<td>No. 8 and smaller</td>
<td>4</td>
<td>201</td>
<td>452</td>
</tr>
</tbody>
</table>

6.2 The coarse sieving of the aggregate shall be performed as follows:

6.2.1 Weigh and record the mass of the sample to be sieved to the nearest gram. Place sample on the top sieve of a nest of 12 inch sieves. The nest of sieves shall consist of sieves starting with the smallest size sieve that 100% of the material will pass, down through and including the No. 8 sieve and pan. Place lid on nested sieves and screen the material by either mechanical or hand shaking, until not more than 0.5 percent by mass of sample passes any sieve during one minute.
6.2.2 Weigh and record separately, to the nearest gram, the mass of the material retained on the individual sieves and in the pan. The material retained in the pan is recorded as the minus No. 8 material.

6.2.3 Do not discard any of the sieved material until the sum of the individual masses is compared to the mass of the sample prior to sieving. If the difference between the two masses is less than or equal to 1.0% of the mass of the sample prior to sieving, an adjustment in mass shall be made on the sieve which has the largest mass retained, except no adjustment shall be made on the minus No. 8 material. If the difference is greater than 1.0%, the sample shall be recombined, resieved, and carefully reweighed.

6.2.4 Determine the coarse sieve factor by dividing 100 by the total mass sieved. Record the factor to six decimal places.

6.2.5 The percent passing for each sieve in the coarse sieve analysis is determined by multiplying the mass retained on that sieve by the coarse sieve factor, and subtracting the result from the unrounded % passing for the next larger sieve. Values for "mass retained multiplied by the coarse sieve factor" and "percent passing each sieve" shall be determined and used in the calculations to six decimal places. The percent passing value for each sieve is recorded in the sieve analysis to the nearest percent.

6.2.6 As a check on the coarse sieve analysis, multiply the mass of minus No. 8 material by the coarse sieve factor. The result of this calculation, rounded to the nearest percent, should be the same as the value for percent passing the No. 8 sieve determined in Subsection 6.2.5 above.

6.2.7 The material passing the No. 8 sieve is split, if necessary, to obtain a minimum 500 gram sample for fine sieving; however, the sample size may be less than 500 grams if a minimum of 500 grams is not obtained from coarse sieving. If less than 800 grams passes the No. 8 sieve, the entire amount shall be subjected to fine sieving. The mass of the sample for fine sieving is recorded to the nearest gram as mass of pass No. 8 split.

6.3 The elutriation and fine sieving of the pass No. 8 material shall be performed as follows:

6.3.1 Subject sample to elutriation through a No. 200 screen either by hand or mechanical washing.
6.3.2 Dry sample to constant mass, allow to cool, then weigh and record the dry mass to the nearest gram.

6.3.3 Place sample on the top sieve of a nest of fine sieves. The nest of sieves shall consist of sieves starting with the No. 10 sieve, down through and including the No. 200 sieve and pan. Place lid on nested sieves and screen the material by either mechanical or hand shaking, until not more than 0.5 percent by mass of sample passes any sieve during one minute.

6.3.4 Weigh and record separately, to the nearest gram, the mass of material retained on the individual sieves and in the pan.

6.3.5 Do not discard any of the sieved material until the sum of the individual masses is compared to the mass of the sample prior to sieving. If the difference between the two masses is less than or equal to 1.0% of the mass of the sample prior to sieving, an adjustment in mass shall be made on the sieve which has the largest mass retained, except no adjustment shall be made on the minus No. 200 material. If the difference is greater than 1.0%, the sample shall be recombined, resieved, and carefully reweighed.

6.3.6 Determine and record elutriation to nearest gram by determining the difference between the dry mass and the mass of the pass No. 8 split.

6.3.7 Determine a factor for calculating the fine sieve analysis by dividing the percent passing the No. 8 sieve (recorded to the nearest percent) by the mass of pass No. 8 split. Record the factor to six decimal places. If all the pass No. 8 material from coarse sieving was subjected to elutriation and fine sieving, a fine sieve factor is not determined. Rather, the coarse sieve factor is utilized and the calculation of the percent passing each sieve is continuous through the entire sieve analysis.

6.3.8 The percent passing for each sieve in the fine sieve analysis is determined by multiplying the mass retained on that sieve by the fine sieve factor, and subtracting the result from the unrounded % passing the next larger sieve, with the exception of the percent passing the No. 8 which has previously been recorded to the nearest percent. Values for "mass retained multiplied by the fine sieve factor" and "percent passing each sieve" shall be determined and used in the calculations to six decimal places. The percent passing value for each sieve is recorded in the sieve analysis to the nearest percent, except the percent passing the No. 200 sieve is recorded to the nearest 0.1 percent.
6.3.9  As a check on the fine sieve analysis, the mass of material passing the No. 200 sieve is added to the elutriation mass, and this total is multiplied by the fine sieve factor. The result of this calculation, rounded to the nearest 0.1 percent, should be the same as the value for the percent passing the No. 200 sieve determined in Subsection 6.3.8 above.

6.3.10  If an aggregate gradation correction factor is utilized, the percent passing the No. 200 sieve shall be adjusted by subtracting the correction factor determined in Subsection 4.15.

6.4  Other methods may be used that differ from that specified in Subsections 6.2 and 6.3 above to determine % passing each sieve, so long as the method utilized has been proven to give equivalent results. However, any procedure which includes recording percent retained values prior to completing the calculation of all percent passing values is not allowed.

7. REPORT AND EXAMPLE

7.1  Report test information on the Asphaltic Concrete Tabulation – Ignition Furnace laboratory card. An example for the testing performed on a field sample is shown in Figure 1. Only the portion of the laboratory card relevant to the ignition furnace test is used for the example. A blank Asphaltic Concrete Tabulation – Ignition Furnace laboratory card is shown in Figure 2.

7.1.1  Mass of basket assembly.

7.1.2  Mass of sample and basket assembly.

7.1.3  Calculated initial mass of the sample.

7.1.4  Mass of sample and basket assembly after ignition.

7.1.5  Calculated mass of sample after ignition.

7.1.6  Asphalt binder content calibration factor.

7.1.7  Percent moisture from moisture test, if one was performed.

7.1.8  Corrected percent asphalt binder content.

7.1.9  Elapsed time of test.
7.1.10 Name of the operator.
7.1.11 Sample test date.
7.1.12 Design percent asphalt binder content.
7.2.13 Ignition furnace set temperature.
7.2.14 If determined, the sieve analysis of the residual aggregate (corrected for passing the No. 200 sieve if applicable).
### FIGURE 1

**EXAMPLE**

<table>
<thead>
<tr>
<th>WEIGHTS RETAINED</th>
<th>% RET</th>
<th>% PASS</th>
<th>SPECS</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
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<td>1</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>3/4</td>
<td>8</td>
<td>85</td>
<td></td>
</tr>
<tr>
<td>1/4</td>
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</tr>
<tr>
<td>1/8</td>
<td>1</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>2.00</td>
<td></td>
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</tbody>
</table>

**WEIGHTING of PASS #1 Split**

<table>
<thead>
<tr>
<th>WEIGHTS RETAINED</th>
<th>% RET</th>
<th>% PASS</th>
<th>SPECS</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>5.0</td>
<td>40</td>
<td></td>
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<tr>
<td>16</td>
<td>3.4</td>
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<tr>
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<td>40</td>
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</tr>
<tr>
<td>50</td>
<td>1.9</td>
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</tr>
<tr>
<td>100</td>
<td>0.3</td>
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</tr>
<tr>
<td>200</td>
<td>0.2</td>
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</tr>
<tr>
<td>400</td>
<td>0.0</td>
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<td></td>
</tr>
<tr>
<td>Total</td>
<td>4.51</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**BULK SPECIFICATIONS**

- Marshall = 7.5
- Gyratory = 5.8
- Standard Penetration = 50
- Shear Wave Velocity = 0.05

**COMPACTION**

- Sample Max. Sp. Gr. (Gmm) = 3.02
- Sample Max. Density [Gmm x 82.5] = 0.01
- Air Voids = 5.8%

**MARSHALL**

- Average Bulk Density (Gmm) = 3.02
- Average Bulk Density [Gmm x 82.5] = 0.01
- Air Voids = 5.8%

**GYRATORY**

- Average Relative Density % Gmm at 100 = 50.2
- Air Voids = 5.8%

**RICE**

- Sample Max. Sp. Gr. (Gmm) = 3.02
- Sample Max. Density [Gmm x 82.5] = 0.01
- Air Voids = 5.8%

**REMARKS**

- 4:16 Special Mix
### Figure 2

#### Asphaltic Concrete Tabulation – Ignition Furnace

<table>
<thead>
<tr>
<th>Use Capital Letters</th>
<th>Lab Number</th>
<th>Org Number</th>
<th>Matl</th>
<th>Type</th>
<th>Purpose</th>
<th>Test Lab</th>
<th>Size</th>
<th>Size %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test No.</td>
<td>Lot Or Suffix</td>
<td>Sampled By</td>
<td>Mo</td>
<td>Day</td>
<td>Year</td>
<td>Time</td>
<td>Military Time</td>
<td>Station</td>
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<td>Sampled From</td>
<td>Lift No.</td>
<td>Hwy</td>
<td>F Milepost, Input Decimal</td>
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<td></td>
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<tr>
<td>Original Source</td>
<td>Project Engineer / Supervisor</td>
<td>Project Number</td>
<td>TRACS Number</td>
<td>Remarks</td>
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</table>

#### Use Capital Letters

<table>
<thead>
<tr>
<th>Coarse Sieve Total</th>
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</thead>
</table>

#### Ignition Furnace

<table>
<thead>
<tr>
<th><strong>ARIZ. 427A</strong></th>
<th><strong>ARIZ. 428A</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>a. Wet Mass of Moisture Sample</td>
<td></td>
</tr>
<tr>
<td>b. Dry Mass of Moisture Sample</td>
<td></td>
</tr>
<tr>
<td>c. Moisture Content ((%)) = ((\text{b} - \text{a})) (\times 100)</td>
<td></td>
</tr>
<tr>
<td>d. Mass of Racket Assembly</td>
<td></td>
</tr>
<tr>
<td>e. Mass of Sample and Racket Assembly</td>
<td></td>
</tr>
<tr>
<td>f. Initial Mass of Sample ((\text{b} - \text{a}))</td>
<td></td>
</tr>
<tr>
<td>g. Ignition Furnace Set Temperature</td>
<td></td>
</tr>
<tr>
<td>h. Mass of Sample and Racket Assembly After Ignition</td>
<td></td>
</tr>
<tr>
<td>i. Mass of Sample After Ignition ((\text{b} - \text{a}))</td>
<td></td>
</tr>
<tr>
<td>j. Uncorrected Asphalt Binder Content ((%)) = (\frac{\text{g}}{\text{h}}) (\times 100)</td>
<td></td>
</tr>
<tr>
<td>k. Asphalt Binder Content ((%))</td>
<td></td>
</tr>
<tr>
<td>l. Ignition Furnace Operation (Tank Stall Correction)</td>
<td></td>
</tr>
<tr>
<td>m. Corrected Asphalt Binder Content ((%)) = (\frac{\text{j}}{\text{k}}) (\times \text{l})</td>
<td></td>
</tr>
<tr>
<td>n. Design Asphalt Binder Content</td>
<td></td>
</tr>
<tr>
<td>o. Exposure Time of Test (minutes)</td>
<td></td>
</tr>
</tbody>
</table>

#### Compaction

- **Marshall**
  - Marshall = M
  - Gyratory = G
  - Core = C

- **Rice**
  - Sample Max. G. O. S. (Gross) | pcf
  - Sample Max. Density (Gross) (Gross) \(\times 10^3\) | pcf

- **Marshall**
  - Average Bulk G. O. S. (Gross) | pcf
  - Average Bulk Density (Gross) \(\times 10^3\) | pcf
  - Air Voids = | %
  - Stability | Bs
  - Flow | 0.1 |

- **Gyratory**
  - Average Relative Density (\% Gross) or Moldsign | %
  - Average Relative Density (\% Gross) or Moldsign \(\times 10^3\) | %

#### White

- Fl. Ll.
- Blue

---

**Received Date:** 44-6372/R03/13  
**Test Operator and Date:**  
**See Back Also:**  
**Figure 2**
ASPHALT BINDER CONTENT OF ASPHALTIC CONCRETE MIXTURES CONTAINING RECLAIMED ASPHALT PAVEMENT (RAP) BY THE IGNITION FURNACE METHOD

(A Modification of AASHTO T 308)

1. SCOPE

1.1 This procedure describes the method for determining the percent asphalt binder content of asphaltic concrete mixtures containing reclaimed asphalt pavement (RAP), by use of an ignition furnace. The aggregate remaining after ignition can be used for sieve analysis, as indicated in Section 6.

1.1.1 The gradation, moisture content, and binder content of the RAP material are determined as described in Appendix A of this test method. The determination of the RAP aggregate gradation is also discussed in Appendix A. The procedure for determining the RAP binder content correction factor is described in Appendix B.

1.1.2 This procedure addresses the use of reclaimed asphalt pavement (RAP) in asphaltic concrete mixtures. See Arizona Test Method 427 when testing is to be performed on asphaltic concrete mixtures which do not contain RAP.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 A listing of subsequent Sections and Figures in this procedure is given below:
2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 Ignition Furnace - a forced-air ignition furnace that heats the sample by the convection method. The furnace must be capable of heating to temperatures up to 538 °C (1000 °F), and able to maintain a given temperature at ± 5 °C (± 9 °F). The furnace shall have an internal weighing system thermally isolated from the furnace chamber and accurate to 0.1 gram. The balance shall be capable of weighing a 3500 gram sample in addition to the sample baskets. A data collection system shall also be included so that the sample mass loss can be automatically determined to an accuracy of 0.1 gram and displayed during a test. The furnace shall provide a printout that includes, as a minimum, the initial sample mass, sample mass loss, 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 one minute intervals. A filter capable of reducing
emissions to an acceptable level shall also be incorporated into the furnace. The furnace shall be vented into a hood or to the outside and be set up properly so that there are no noticeable odors escaping into the laboratory. The furnace shall have a fan with the capability to pull air through the furnace to expedite the test and to reduce escape of smoke into the laboratory. The furnace shall be equipped so that the door cannot be opened until testing is complete.

2.1.2 Stainless Steel Perforated Baskets - the baskets shall be an appropriate size that allow the samples to be a thickness which allows air to flow up through and around the sample particles. The sample shall be completely enclosed with screen mesh, perforated stainless steel plate, or other suitable material. Screen mesh or other suitable material with openings of No. 8 has been found to perform well.

2.1.3 Stainless Steel Catch Pan - of sufficient size to hold the sample baskets so that aggregate particles and melting asphalt binder falling through the screen mesh are caught.

2.1.4 Oven(s) - capable of heating to temperatures up to 350 °F, and able to maintain a given temperature at the tolerances specified herein.

2.1.5 Scale(s) or balance(s) - capable of measuring the maximum mass to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.1.6 Safety Equipment - safety glasses or face shield, high temperature gloves, and long sleeve jacket. Additionally, a heat resistant surface capable of withstanding 1200 °F and a protective cage capable of surrounding the sample baskets shall be provided.

2.1.7 Miscellaneous Equipment - a pan larger than the sample basket(s) for transferring samples after ignition, spatulas, bowls, spoons, and wire brushes.

2.1.8 Mixing apparatus - Mechanical mixing is recommended; 20 quart capacity mixer is required. (Hand mixing may be performed if desired.)

2.1.9 Thermometer - with a temperature range of 50 to 500 °F.

2.1.10 Hot plate - capable of heating to temperatures up to 350 °F, and able to maintain a given temperature at ± 5 °F.
2.1.11 For performing sieve analysis, apparatus as specified in Arizona Test Method 201.

3. **SAMPLING**

3.1 For preparing calibration samples, obtain representative samples of virgin aggregate and RAP material in accordance with Arizona Test Method 105. Samples of virgin aggregate and RAP material shall be sufficiently large to provide enough material for calibration testing.

3.2 For testing field samples of asphaltic concrete, obtain a representative sample of the freshly produced mix in accordance with Arizona Test Method 104. Obtain representative test samples, in accordance with the appropriate sections of Arizona Test Method 416, for the determination of moisture content (if required) and asphalt binder content.

3.2.1 The size of the asphalt binder content test sample shall be within ±150 grams of the sample size used for calibration and must also conform to the mass requirement shown in Table 1. When the mass of the test sample exceeds the capacity of the equipment used, the test sample shall be divided into suitable increments, tested, and the results appropriately combined.

4. **CALIBRATION**

4.1 Asphalt binder content results may be affected by the type of aggregate and binder in the mixture. A calibration factor for the asphalt binder content must be established for each mix design. Certain aggregate types may result in an unusually high calibration factor and erroneous gradation results due to aggregate breakdown. Such mixes should be calibrated and tested at a lower temperature as described in Subsection 4.13.

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>Size of Test Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nominal Maximum Aggregate Size (See Note)</strong></td>
<td><strong>Mass of Sample, grams</strong></td>
</tr>
<tr>
<td>1-1/2 in.</td>
<td>4000 – 4500</td>
</tr>
<tr>
<td>1 in.</td>
<td>3000 – 3500</td>
</tr>
<tr>
<td>3/4 in.</td>
<td>2000 – 2500</td>
</tr>
<tr>
<td>1/2 in.</td>
<td>1500 – 2000</td>
</tr>
<tr>
<td>3/8 in.</td>
<td>1200 – 1700</td>
</tr>
<tr>
<td>No. 4</td>
<td>1200 – 1700</td>
</tr>
</tbody>
</table>

Note: The smallest sieve opening through which the entire amount of material, by specification, is permitted to pass.
4.2 Spread the RAP material from each individual RAP stockpile to be used for the calibration in separate pans at 1 to 1-1/2 inches in depth and dry overnight at \(140 \pm 10 \, ^\circ\text{F}\), and continue drying until the mass, after successive one-half hour periods of drying, indicates no change in mass. Allow the material to cool. Screen the RAP material from the individual stockpiles and separate into individual sizes for No. 8 and larger sieves, and minus No. 8 material. The RAP material shall be screened for 5 minutes \(\pm 15\) seconds per Arizona Test Method 240 to prevent excessive breakdown of the RAP agglomerations.

4.3 Dry the individual virgin aggregate stockpile samples to constant mass at \(290 \pm 10 \, ^\circ\text{F}\). Allow the material to cool. Screen the virgin aggregate from the individual stockpiles and separate into individual sizes for No. 8 and larger sieves, and minus No. 8 material.

**Note:** In lieu of drying and screening the virgin aggregate samples for each individual stockpile, a bulk-batched sample may be used as described in Subsection 4.3.1.

4.3.1 Using the adjusted stockpile percentages shown in Mix Design Composite #2 (Virgin Aggregate Composite), the material from the individual stockpiles may be bulk-batched into a single sample of an adequate amount of material necessary to prepare the required calibration samples. Dry the bulk-batched sample to constant mass at \(290 \pm 10 \, ^\circ\text{F}\). Allow the material to cool. Screen the bulk-batched material and separate into individual sizes for No. 8 and larger sieves, and minus No. 8 material.

4.4 The RAP aggregate [extracted aggregate (AASHTO T 164 - Method A) from the RAP material] shall be dried to constant mass at \(290 \pm 10 \, ^\circ\text{F}\). Allow the material to cool. Screen the RAP aggregate material from the individual stockpiles and separate into individual sizes for No. 8 and larger sieves, and minus No. 8 material.

**Note:** The batching of samples described below in Subsection 4.5 and Subsection 4.6 is based on the mix design weigh-up information, as shown in the example in Figure 1.

4.5 A gradation check sample is batched using:

- The virgin mineral aggregate component (individual stockpile material or bulk-batched material), which has been screened into individual sizes for No. 8 and larger sieves, and minus No. 8 material.
- The RAP aggregate component, which has been screened into individual sizes for No. 8 and larger sieves, and minus No. 8 material.

- Mineral admixture.

4.5.1 Batch the virgin aggregate portion to Mix Design Composite #2 (Virgin Aggregate Composite). If the individual virgin aggregate stockpiles were screened, batch to the individual accumulative weights required for each individual stockpile, as shown in the example in Figure 2. If bulk-batched virgin aggregate material is used, batch to the accumulative weights required for the total batch weight as shown in the example in Figure 2.

4.5.1.1 The weight of the virgin aggregate portion is determined as follows:

$$W_{va} = \left( \frac{100 - T_{pra}}{100} \right) \times (W_{ta})$$

Where:

- $W_{va} =$ Weight of virgin aggregate
- $T_{pra} =$ Total percent of RAP aggregate in mix design
- $W_{ta} =$ Weight of total aggregate desired (combined virgin aggregate and RAP aggregate)

Example:

$$W_{va} = \left( \frac{100 - (15 + 10)}{100} \right) \times (2200) = 1650 \text{ grams}$$

4.5.2 Batch the RAP aggregate portion to Mix Design Composite #1 (Virgin Aggregate and RAP Aggregate Composite), utilizing the accumulative weights required for each individual RAP aggregate material, as shown in the example in Figure 2.

4.5.2.1 The weight of RAP aggregate portion required for each RAP stockpile is determined as follows:

$$W_{ra} = \left( \frac{P_{ra}}{100} \right) \times (W_{ta})$$
Where:

\[ W_{ra} = \frac{15}{100} \times (2200) = 330.0 \text{ grams} \]

Example (for coarse RAP aggregate):

\[ W_{ra} = \frac{10}{100} \times (2200) = 220.0 \text{ grams} \]

4.5.3 Add the required type and amount of mineral admixture to the virgin aggregate and RAP aggregate and blend thoroughly.

4.5.3.1 The weight of mineral admixture is determined as follows:

\[ W_{ad} = \frac{P_{ad}}{100} \times (W_{ta}) \]

Where:

\[ W_{ad} = \text{Weight of mineral admixture} \]
\[ P_{ad} = \text{Percent mineral admixture} \]
\[ W_{ta} = \text{Weight of total aggregate desired (combined virgin aggregate and RAP aggregate)} \]

Example:

\[ W_{ad} = \frac{1.0}{100} \times (2200) = 22 \text{ grams} \]

4.5.4 The weight of the combined virgin aggregate, RAP aggregate, and mineral admixture for the gradation check sample shall conform to the requirements of Table 1.
4.5.5 Perform a gradation analysis on the gradation check sample in accordance with Section 6 to determine the actual gradation.

4.6 Two calibration (burn) samples and also a “butter mix” sample are batched using:

- The virgin mineral aggregate component (individual stockpile material or bulk-batched material), which has been screened into individual sizes for No. 8 and larger sieves, and minus No. 8 material.

- The RAP material (dry screened RAP) component, which has been screened into individual sizes for No. 8 and larger sieves, and minus No. 8 material.

- Mineral admixture.

4.6.1 Batch the virgin aggregate portion to Mix Design Composite #2 (Virgin Aggregate Composite). If the individual virgin aggregate stockpiles were screened, batch to the individual accumulative weights required for each individual stockpile, as shown in the example in Figure 3. If bulk-batched virgin aggregate material is used, batch to the accumulative weights required for the total batch weight as shown in as shown in the example in Figure 3.

4.6.1.1 The weight of the virgin aggregate portion is determined as shown in Subsection 4.5.1.1.

4.6.2 Add the required type and amount of mineral admixture to each of the batched virgin aggregate samples and blend thoroughly.

4.6.2.1 The weight of the mineral admixture is determined as shown in Subsection 4.5.3.1.

4.6.3 Form a shallow crater in the center of each of the blended virgin aggregate/mineral admixture samples.

4.6.4 Batch the dry screened RAP portion to Mix Design Composite #3 (Virgin Aggregate and Dry Screened RAP Composite), utilizing the accumulative weights required for each individual dry screened RAP material, as shown in the example in Figure 3.

4.6.4.1 The weight of the dry screened RAP portion from each RAP stockpile is determined as follows:
Where:

\[ W_{\text{rap}} = \left( \frac{W_{\text{ra}}}{1 - \frac{P_{\text{br}}}{100}} \right) \]

Example (for fine RAP stockpile):

\[ W_{\text{rap}} = \left( \frac{330.0}{1 - \frac{5.82}{100}} \right) = 350.4 \text{ grams} \]

Example (for coarse RAP stockpile):

\[ W_{\text{rap}} = \left( \frac{220.0}{1 - \frac{3.43}{100}} \right) = 227.8 \text{ grams} \]

4.6.5 Place the dry screened RAP material in the crater formed in each of the blended virgin aggregate/mineral admixture samples.

4.6.6 The weight of the combined virgin aggregate, dry screened RAP, and mineral admixture for each calibration sample and the “butter mix” sample shall be such that when the required amount of virgin asphalt binder is added, they conform to the requirements of Table 1.

Note: It is recommended that an initial calibration sample be prepared and tested in accordance with the requirements of Subsection 4.6.7 prior to the preparation of additional samples. Doing so may help avoid the waste of the limited amount of dry screened RAP material.

4.6.7 Prior to the addition of the virgin binder, record the weight of the combined virgin aggregate, dry screened RAP, and mineral admixture for each calibration sample to the nearest 0.1 gram as the “Weight of Initial Charge Before Drying”. Dry each charge to constant mass at 290 ± 10 °F, allow to cool, and record the weight to the nearest 0.1 gram as the “Weight of Initial Charge After Drying”. If the loss from drying is greater than
25 grams, discard the charge and prepare a new charge using the same procedure, except the dry screened RAP shall be initially dried overnight at 160° ± 3 °F, and drying continued at 160° ± 3 °F until the mass, after successive one-half hour periods of drying, indicates no change in mass. Do not add make-up material to account for any loss from drying, as doing so will change the gradation of the samples.

Note: The extended drying at the increased temperature (from 140 ± 10 °F to 160° ± 3 °F) specified above is to ensure that all moisture is removed from the dry screened RAP material.

4.6.8 Two calibration samples and the butter mix are prepared as described below at the design total asphalt binder content. The asphalt binder grade and type shall be the same as will be used in the asphalt concrete mixture to be tested during production. The percent total asphalt binder content is based on the mass of the total mix. For each sample, the weight of virgin asphalt binder to be added is determined using the calculations in Subsections 4.5.1.1, 4.5.3.1, and 4.5.6.1 in conjunction with Subsections 4.6.8.1, 4.6.8.2, and 4.6.8.3.

4.6.8.1 The weight of binder contributed by the RAP material from each RAP stockpile is determined as follows:

\[ W_{rb} = \left( \frac{P_{br}}{100} \right) \times (W_{rap}) \]

Where:

- \( W_{rb} \) = Weight of binder contributed by each RAP stockpile
- \( P_{br} \) = Percent binder content of each RAP stockpile
- \( W_{rap} \) = Weight of RAP from each RAP stockpile

Example (for fine RAP stockpile):

\[ W_{rb} = \left( \frac{5.82}{100} \right) \times (350.4) = 20.39 \text{ grams} \]

Example (for coarse RAP stockpile):

\[ W_{rb} = \left( \frac{3.43}{100} \right) \times (227.8) = 7.81 \text{ grams} \]
4.6.8.2  The weight of total binder is determined as follows:

\[
W_{tb} = \left( \frac{W_{va} + T_{wra} + W_{ad}}{100 - P_{tb}} \right) \times (P_{tb})
\]

Where:
- \(W_{tb}\) = Weight of total binder
- \(W_{va}\) = Weight of virgin aggregate
- \(T_{wra}\) = Total weight of RAP aggregate
- \(W_{ad}\) = Weight of mineral admixture
- \(P_{tb}\) = Percent total binder (mix design percent binder content)

Example:

\[
W_{tb} = \left( \frac{1650 + (330 + 220) + 22}{100 - 5.4} \right) \times (5.4) = 126.84 \text{ grams}
\]

4.6.8.3  The weight of virgin asphalt binder to be added is determined as follows:

\[
W_{vb} = (W_{tb}) - (W_{trb})
\]

Where:
- \(W_{vb}\) = Weight of virgin asphalt binder to be added
- \(W_{tb}\) = Weight of total binder
- \(W_{trb}\) = Total weight of binder contributed by RAP stockpiles

Example:

\[
W_{vb} = (126.84) - (20.39 + 7.81) = 98.6 \text{ grams}
\]

**Note:** If desired, the percent of the total binder which is contributed by RAP can be calculated as follows:

\[
\begin{bmatrix}
\text{Percent of Total Binder Contributed by RAP}
\end{bmatrix} = \left( \frac{W_{trb}}{W_{tb}} \right) \times (100)
\]
Where:

\[ W_{trb} = \text{Total weight of binder contributed by RAP stockpiles} \]
\[ W_{tb} = \text{Weight of total binder required} \]

Example:

\[ \left[ \frac{\text{Percent of Total Binder Contributed by RAP}}{100} \right] = \left( \frac{22.23}{126.84} \right) \times (100) = 22.23\% \]

4.6.8.4 All bowls, sample pans, and mixing tools shall be preheated to approximately the laboratory mixing temperature prescribed in the mix design. At the time mixing of the samples begins, the temperature of the asphalt binder, virgin aggregate, RAP, and mineral admixture shall be in accordance with the prescribed laboratory mixing temperature \( \pm 5 \) °F. Each individual sample shall be thoroughly mixed. All samples shall be mixed at the same mixing temperature \( \pm 5 \) °F.

**Note:** If the mix design laboratory mixing temperature is not specified, a temperature of 300 \( \pm 5 \) °F shall be used.

4.7 Preheat the ignition furnace to 538 \( \pm 5 \) °C (1000 \( \pm 9 \) °F), or as modified in Subsection 4.13. Do not preheat the sample basket.

4.8 Weigh and record the mass of the basket assembly to the nearest 0.1 gram.

4.9 The freshly mixed samples may be placed directly in the sample basket assembly. If the samples are allowed to cool, they must be reheated in a 290 \( \pm 10 \) °F oven for 25 minutes.

4.10 Test samples in accordance with Subsections 5.6 through 5.14.

4.11 If the difference between the measured asphalt binder content of the two samples exceeds 0.07, repeat the test using two additional samples, and from the four results discard the high and the low values.

4.12 Subtract the actual asphalt binder content for each of the two samples from their respective measured asphalt binder content. The asphalt binder content calibration factor is the average of the two resultant values expressed in percent by mass of the asphalt mixture.
4.13 If the asphalt binder content calibration factor exceeds ± 1.25 percent, lower the test temperature to 482 ± 5 °C (900 ± 9 °F) and repeat the test to determine a new calibration factor. If the calibration factor continues to exceed ± 1.25 percent, lower the test temperature to 427 ± 5 °C (800 ± 9 °F) and repeat the test to determine a new calibration factor. Use the calibration factor obtained at 427 ± 5 °C (800 ± 9 °F) even if it exceeds ± 1.25 percent.

4.14 Perform a gradation analysis on the residual aggregate as indicated in Section 6. Subtract the actual percent passing the No. 200 sieve for each sample from the measured percent passing the No. 200 sieve (as determined in Subsection 4.5.5). Determine the average of the two values. If the resultant average value is greater than ± 0.50, an aggregate gradation correction factor (equal to the resultant average value) for the passing No. 200 material may be applied to the production field sample test results.

5. PROCEDURE

5.1 The moisture content of the asphaltic concrete shall be determined in accordance with Arizona Test Method 406. The moisture content sample shall be obtained at the same time and subjected to the same treatment prior to testing as the asphalt binder content test sample. As an alternate to performing the moisture determination, the test sample may be dried to a constant mass in an oven at 290 ± 10 °F.

5.2 Preheat the ignition furnace to 538 ± 5 °C (1000 ± 9 °F), or to the alternate temperature determined during the calibration (Subsection 4.13). Do not preheat the sample basket. Record the furnace temperature set point prior to the initiation of the test.

5.3 Record the asphalt binder content calibration factor, determined in accordance with Subsections 4.11 through 4.13, for the specific mix to be tested.

5.4 Weigh and record the mass of the basket assembly to the nearest 0.1 gram.

5.5 Obtain the asphalt binder content test sample in accordance with Subsection 3.2, ensuring that the size of the test sample is within ± 150 grams of the sample size used for calibration and that the test sample conforms to the requirements shown in Table 1.
5.6 Evenly distribute the test sample over the center of the sample basket(s) and level the material. Use a spatula or trowel to pull material approximately one inch away from the outside edge of basket(s).

5.7 Weigh and record the mass of the sample and basket assembly to the nearest 0.1 gram.

5.8 Calculate and record the initial mass of the sample to the nearest 0.1 gram.

5.9 Set the ignition furnace controller print mode to give a printout of the test data required in Subsection 2.1.1. Input the initial mass of the sample into the ignition furnace controller. Verify that the correct mass has been entered.

5.10 Open the furnace door and place the sample and basket assembly so that it is centered in the chamber. After assuring that the sample basket assembly is not in contact with any wall, close the door. Initiate the test by pressing the start button. This will lock the furnace door and start testing.

5.11 Allow the test to continue until the stable light and audible stable indicator indicates the test is complete. The test is deemed complete when the measured mass loss does not exceed 0.01 percent of the sample mass for three consecutive one minute intervals. If required, press the stop button. This will unlock the furnace door and cause the printer to print out the test results.

5.12 Open the furnace door and remove the sample and basket assembly. Allow the sample to cool 30 ± 5 minutes in the basket assembly. Weigh and record the mass of the sample and basket assembly after ignition to the nearest 0.1 gram.

5.13 Calculate and record the mass of sample after ignition to the nearest 0.1 gram.

5.14 Calculate and record the corrected asphalt binder content of the sample, to the nearest 0.01%, as follows:

\[
\%AC = \left[ \frac{W_i - W_A}{W_i} \times 100 \right] - C_F - \%M
\]
Where: $\%AC = \text{measured (corrected) asphalt binder content in percent by mass of the sample}$

$W_i = \text{mass of the sample prior to ignition}$

$W_A = \text{mass of the sample after ignition}$

$C_F = \text{asphalt binder content calibration factor, percent by mass of the sample}$

$\%M = \text{percent moisture in the sample}$

**Note:** During calibration, $C_F$ and $\%M$ are zero.

5.14.1 If an ignition furnace correction (tank stab correction) is made, the $\%AC$ determined in Subsection 5.14 is adjusted by that correction.

5.15 Attach the original printed ticket to the back of the card.

5.16 Empty the contents of the baskets into a flat pan. Use a small wire sieve brush to ensure that any residual fines are removed from the baskets. Take care not to lose any material, as this will affect gradation results.

5.17 If needed, perform a gradation analysis of the residual aggregate according to Section 6.

6. **SIEVE ANALYSIS OF AGGREGATE**

6.1 If required, the aggregate shall be subjected to sieve analysis as described below. The coarse sieving shall be performed in accordance with Subsection 6.2, and the fine sieving in accordance with Subsection 6.3. The quantity of material on a given sieve at the completion of sieving shall not exceed the amount shown Table 2.

6.2 The coarse sieving of the aggregate shall be performed as follows:

6.2.1 Weigh and record the mass of the sample to be sieved to the nearest gram. Place sample on the top sieve of a nest of 12 inch sieves. The nest of sieves shall consist of sieves starting with the smallest size sieve that 100% of the material will pass, down through and including the No. 8 sieve and pan. Place lid on nested sieves and screen the material by either mechanical or hand shaking, until not more than 0.5 percent by mass of sample passes any sieve during one minute.

6.2.2 Weigh and record separately, to the nearest gram, the mass of the material retained on the individual sieves and in the pan. The material retained in the pan is recorded as the minus No. 8 material.
### TABLE 2

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Maximum Mass Allowed (grams/sq. in.)</th>
<th>Maximum Mass Allowed (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>8 inch Diameter Sieve</td>
<td>12 inch Diameter Sieve</td>
</tr>
<tr>
<td>1-1/2&quot;</td>
<td>25</td>
<td>---</td>
</tr>
<tr>
<td>1&quot;</td>
<td>18</td>
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</tr>
<tr>
<td>3/4&quot;</td>
<td>14</td>
<td>---</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>10</td>
<td>---</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>8</td>
<td>---</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>6</td>
<td>---</td>
</tr>
<tr>
<td>No. 4</td>
<td>5</td>
<td>---</td>
</tr>
<tr>
<td>No. 8 and smaller</td>
<td>4</td>
<td>201</td>
</tr>
</tbody>
</table>

6.2.3 Do not discard any of the sieved material until the sum of the individual masses is compared to the mass of the sample prior to sieving. If the difference between the two masses is less than or equal to 1.0% of the mass of the sample prior to sieving, an adjustment in mass shall be made on the sieve which has the largest mass retained, except no adjustment shall be made on the minus No. 8 material. If the difference is greater than 1.0%, the sample shall be recombined, resieved, and carefully reweighed.

6.2.4 Determine the coarse sieve factor by dividing 100 by the total mass sieved. Record the factor to six decimal places.

6.2.5 The percent passing for each sieve in the coarse sieve analysis is determined by multiplying the mass retained on that sieve by the coarse sieve factor, and subtracting the result from the unrounded % passing for the next larger sieve. Values for "mass retained multiplied by the coarse sieve factor" and "percent passing each sieve" shall be determined and used in the calculations to six decimal places. The percent passing value for each sieve is recorded in the sieve analysis to the nearest percent.

6.2.6 As a check on the coarse sieve analysis, multiply the mass of minus No. 8 material by the coarse sieve factor. The result of this calculation, rounded to the nearest percent, should be the same as the value for percent passing the No. 8 sieve determined in Subsection 6.2.5 above.
6.2.7 The material passing the No. 8 sieve is split, if necessary, to obtain a minimum 500 gram sample for fine sieving; however, the sample size may be less than 500 grams if a minimum of 500 grams is not obtained from coarse sieving. If less than 800 grams passes the No. 8 sieve, the entire amount shall be subjected to fine sieving. The mass of the sample for fine sieving is recorded to the nearest gram as mass of pass No. 8 split.

6.3 The elutriation and fine sieving of the pass No. 8 material shall be performed as follows:

6.3.1 Subject sample to elutriation through a No. 200 screen either by hand or mechanical washing.

6.3.2 Dry sample to constant mass, allow to cool, then weigh and record the dry mass to the nearest gram.

6.3.3 Place sample on the top sieve of a nest of fine sieves. The nest of sieves shall consist of sieves starting with the No. 10 sieve, down through and including the No. 200 sieve and pan. Place lid on nested sieves and screen the material by either mechanical or hand shaking, until not more than 0.5 percent by mass of sample passes any sieve during one minute.

6.3.4 Weigh and record separately, to the nearest gram, the mass of material retained on the individual sieves and in the pan.

6.3.5 Do not discard any of the sieved material until the sum of the individual masses is compared to the mass of the sample prior to sieving. If the difference between the two masses is less than or equal to 1.0% of the mass of the sample prior to sieving, an adjustment in mass shall be made on the sieve which has the largest mass retained, except no adjustment shall be made on the minus No. 200 material. If the difference is greater than 1.0%, the sample shall be recombined, resieved, and carefully reweighed.

6.3.6 Determine and record elutriation to nearest gram by determining the difference between the dry mass and the mass of the pass No. 8 split.

6.3.7 Determine a factor for calculating the fine sieve analysis by dividing the percent passing the No. 8 sieve (recorded to the nearest percent) by the mass of pass No. 8 split. Record the factor to six decimal places. If all the pass No. 8 material from coarse sieving was subjected to elutriation and fine sieving, a fine sieve factor is not determined. Rather, the coarse
sieve factor is utilized and the calculation of the percent passing each sieve is continuous through the entire sieve analysis.

6.3.8 The percent passing for each sieve in the fine sieve analysis is determined by multiplying the mass retained on that sieve by the fine sieve factor, and subtracting the result from the unrounded % passing the next larger sieve, with the exception of the percent passing the No. 8 which has previously been recorded to the nearest percent. Values for "mass retained multiplied by the fine sieve factor" and "percent passing each sieve" shall be determined and used in the calculations to six decimal places. The percent passing value for each sieve in the sieve analysis is recorded to the nearest percent, except the percent passing the No. 200 sieve is recorded to the nearest 0.1 percent.

6.3.9 As a check on the fine sieve analysis, the mass of material passing the No. 200 sieve is added to the elutriation mass, and this total is multiplied by the fine sieve factor. The result of this calculation, rounded to the nearest 0.1 percent, should be the same as the value for the percent passing the No. 200 sieve determined in Subsection 6.3.8 above.

6.3.10 If an aggregate gradation correction factor is utilized, the percent passing the No. 200 sieve shall be adjusted by subtracting the correction factor determined in Subsection 4.14.

6.4 Other methods may be used that differ from that specified in Subsections 6.2 and 6.3 above to determine % passing each sieve, so long as the method utilized has been proven to give equivalent results. However, any procedure which includes recording percent retained values prior to completing the calculation of all percent passing values is not allowed.

7. REPORT AND EXAMPLE

7.1 Report test information on the Asphaltic Concrete Tabulation – Ignition Furnace laboratory card. An example for the testing performed on a field sample is shown in Figure 4. Only the portion of the laboratory card relevant to the ignition furnace test is used for the example. A blank Asphaltic Concrete Tabulation – Ignition Furnace laboratory card is shown in Figure 5.

7.1.1 Mass of basket assembly.

7.1.2 Mass of sample and basket assembly.

7.1.3 Calculated initial mass of the sample.
7.1.4 Mass of sample and basket assembly after ignition.
7.1.5 Calculated mass of sample after ignition.
7.1.6 Asphalt binder content calibration factor.
7.1.7 Percent moisture from moisture test, if one was performed.
7.1.8 Corrected percent asphalt binder content.
7.1.9 Elapsed time of test.
7.1.10 Name of the operator.
7.1.11 Sample test date.
7.1.12 Design percent asphalt binder content.
7.1.13 Ignition furnace set temperature.
7.1.14 If determined, the sieve analysis of the residual aggregate (corrected for passing the No. 200 sieve if applicable).
Ignition Furnace Calibration
Mix Design Weigh-Up Information

Weight of Total Virgin Aggregate Required if Bulk Batching is Used (grams)
(Furnace Calibration, Aggregate Properties Testing, etc.)

10000

Total Weight of Aggregate Charge (grams)
(Virgin + RAP Aggregate needed for each Gradation Check and Burn)

2000

Total Binder Content (% by wt of total mix)

5.4

Virgin Binder Content (% by wt of total mix)

4.2

RAP Binder Content (% by wt of total mix)

1.2

Total Virgin Aggregate (% by weight of total aggregate)

75.0

Total RAP Aggregate (% by weight of total aggregate)

25.0

Total

100.0

Admixture (% by weight of total aggregate)

1.0

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>WCF</th>
<th>CF</th>
<th>3/8</th>
<th>3/4</th>
<th>Total</th>
<th>Virgin Composite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent Used (w/o Admix)*</td>
<td>35.0</td>
<td>13.0</td>
<td>8.0</td>
<td>19.0</td>
<td>75.0</td>
<td>100.0</td>
</tr>
<tr>
<td>Adjusted Percent Used</td>
<td>46.67</td>
<td>17.33</td>
<td>10.67</td>
<td>25.33</td>
<td>100.0</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Virgin Aggregate Stockpiles (Composite #2)</th>
<th>Adjusted Virgin Composite</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.5&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>1&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>#4</td>
<td>97.1 98.8 22.5 1.6 1.6</td>
<td>65.24</td>
</tr>
<tr>
<td>#8</td>
<td>60.8 70.1 1.3 1.5 1.5</td>
<td>41.04</td>
</tr>
</tbody>
</table>

* If unadjusted percent virgin aggregate stockpile use is not shown in Composite #2, see Composite #1.

Total Weight Required from each Stockpile if Bulk Batching Virgin Aggregates

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Dry Screened RAP (Composite #3)</th>
<th>RAP Aggregate (Composite #1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieve Analysis (Percent Passing)</td>
<td>Fine</td>
<td>Coarse</td>
</tr>
<tr>
<td>Percent Used (w/o Admix)</td>
<td>15.0</td>
<td>10.0</td>
</tr>
<tr>
<td>Adjusted Percent Used</td>
<td>60.0</td>
<td>40.0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Virgin Aggregate Stockpiles (Composite #2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieve Size</td>
<td>1.5&quot;</td>
</tr>
<tr>
<td>-----------------------</td>
<td>-------</td>
</tr>
<tr>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>100.0</td>
<td>100.0</td>
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<tr>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>100.0</td>
<td>100.0</td>
</tr>
</tbody>
</table>

NOTE: Overall, Composite #2 (RAP Aggregate) will be finer than Composite #3 (Dry Screened RAP).

RAP Ma'l Binder Content (%)

5.82 3.43

FIGURE 1
# Ignition Furnace Calibration

## Weigh-Up Requirements for Gradation Check

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Virgin Aggregate Stockpiles (Composite #2)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>WCF</td>
<td>CF</td>
</tr>
<tr>
<td>Percent Used (w/o Admix)</td>
<td>35.0</td>
<td>13.0</td>
</tr>
<tr>
<td>Adjusted Percent Used</td>
<td>45.67</td>
<td>17.33</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Accumulative Weights Required (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.5&quot; 0.0 0.0 0.0 0.0 0.0</td>
</tr>
<tr>
<td>1&quot; 0.0 0.0 0.0 0.0 0.0</td>
</tr>
<tr>
<td>3/4&quot; 0.0 0.0 0.0 75.2 75.2</td>
</tr>
<tr>
<td>1/2&quot; 0.0 0.0 0.0 361.2 361.2</td>
</tr>
<tr>
<td>3/8&quot; 0.0 0.0 0.0 101.3 101.3</td>
</tr>
<tr>
<td>1/4&quot; 0.0 0.0 72.2 411.3 483.5</td>
</tr>
<tr>
<td>#4 22.3 3.4 136.4 411.3 573.5</td>
</tr>
<tr>
<td>#8 301.8 86.6 173.7 411.7 972.8</td>
</tr>
<tr>
<td>#8 770.0 286.0 176.0 415.0 1650.0</td>
</tr>
</tbody>
</table>

**NOTE:** Use individual stockpile accumulative weights if Virgin Aggregate is not Bulk Batched. Use total accumulative weights if Virgin Aggregate is Bulk Batched prior to sieving.

### Dry Screened RAP (Composite #3)

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Fine</th>
<th>Coarse</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent Used (w/o Admix)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adjusted Percent Used</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Accumulative Weights Required (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.5&quot; N/A N/A</td>
</tr>
<tr>
<td>1.25&quot; N/A N/A</td>
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<td>1&quot; N/A N/A</td>
</tr>
<tr>
<td>3/4&quot; N/A N/A</td>
</tr>
<tr>
<td>1/2&quot; N/A N/A</td>
</tr>
<tr>
<td>3/8&quot; N/A N/A</td>
</tr>
<tr>
<td>1/4&quot; N/A N/A</td>
</tr>
<tr>
<td>#4 N/A N/A</td>
</tr>
<tr>
<td>#8 N/A N/A</td>
</tr>
<tr>
<td>#8 N/A N/A</td>
</tr>
</tbody>
</table>

**RAP Mat'l Binder Content (%)**

<table>
<thead>
<tr>
<th>Fine</th>
<th>Coarse</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.82</td>
<td>3.43</td>
</tr>
</tbody>
</table>

### RAP Aggregate (Composite #1)

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Fine</th>
<th>Coarse</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent Used (w/o Admix)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adjusted Percent Used</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Accumulative Weights Required (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fine 15.0</td>
</tr>
<tr>
<td>Coarse 60.0</td>
</tr>
</tbody>
</table>

**Gradation Check Material Requirements**

| Total Virgin Aggregate Weight Required (grams) | 1650.0 |
| Total RAP Aggregate Weight Required (grams) | 250.0  |
| Admixture Weight Required (grams) | 22.0   |
| Final Charge Weight (grams) | 2222.0 |

**EXAMPLE**

**FIGURE 2**
# Ignition Furnace Calibration

## Weigh-Up Requirements for Burn 1 & 2

### Virgin Aggregate Stockpiles (Composite #2)

<table>
<thead>
<tr>
<th></th>
<th>WCF</th>
<th>CF</th>
<th>3/8</th>
<th>3/4</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjusted Percent Used</td>
<td>46.67</td>
<td>17.33</td>
<td>10.87</td>
<td>25.33</td>
<td>100.00</td>
</tr>
<tr>
<td>Sieve Size</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>1.5&quot;</td>
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<td>0.0</td>
</tr>
<tr>
<td>3/4&quot;</td>
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<td>0.0</td>
<td>75.2</td>
<td>75.2</td>
</tr>
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<td>361.2</td>
<td>361.2</td>
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<tr>
<td>3/8&quot;</td>
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<td>0.0</td>
<td>0.0</td>
<td>401.3</td>
<td>401.3</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>0.0</td>
<td>0.0</td>
<td>72.2</td>
<td>411.3</td>
<td>483.5</td>
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<tr>
<td>#4</td>
<td>22.3</td>
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<td>136.4</td>
<td>411.3</td>
<td>573.5</td>
</tr>
<tr>
<td>#6</td>
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<td>85.5</td>
<td>173.7</td>
<td>411.7</td>
<td>972.6</td>
</tr>
<tr>
<td>#8</td>
<td>770.0</td>
<td>286.0</td>
<td>176.0</td>
<td>418.0</td>
<td>1650.0</td>
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</tbody>
</table>

**NOTE:** Use individual stockpile accumulative weights if Virgin Aggregate is Batched. Use total accumulative weights if Virgin Aggregate is Batched prior to sieving.

### Dry Screened RAP (Composite #3)

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Fine</th>
<th>Coarse</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjusted Percent Used</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sieve Size</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.5&quot;</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>1.25&quot;</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>1&quot;</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>0.0</td>
<td>12.8</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>0.0</td>
<td>97.7</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>0.0</td>
<td>196.5</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>111.8</td>
<td>209.6</td>
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<td>159.4</td>
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<td>222.8</td>
</tr>
<tr>
<td>#8</td>
<td>350.4</td>
<td>227.8</td>
</tr>
</tbody>
</table>

### RAP Aggregate (Composite #1)

<table>
<thead>
<tr>
<th>Stockpile Description</th>
<th>Fine</th>
<th>Coarse</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjusted Percent Used</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sieve Size</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.5&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>1.25&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>1&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>#4</td>
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<td>N/A</td>
</tr>
<tr>
<td>#8</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

### RAP Mat'l Binder Content (%)

- 5.82
- 3.43

Total Wt. of RAP Mat'l (grams)

| 350.4 | 227.8 |

### Weight of RAP Binder (grams)

| 20.39 | 7.81  |

**EXAMPLE**

### Burn 1 & 2 Material Requirements

| Total Virgin Aggregate Weight Required (grams) | 1650.0 |
| Total RAP Material Weight Required (grams)   | 578.2  |
| Admixture Weight Required (grams)            | 22.0   |
| Virgin Binder Weight Required (grams)        | 98.8   |
| Final Charge Weight (grams)                  | 2348.8 |

**FIGURE 3**
**FIGURE 4**

### Example

#### Coarse Factor

<table>
<thead>
<tr>
<th>WEIGHTS RETAINED</th>
<th>% RET</th>
<th>% PASS</th>
</tr>
</thead>
<tbody>
<tr>
<td>3&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
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<td></td>
<td></td>
</tr>
<tr>
<td>2&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 1/2&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>91</td>
<td>4</td>
</tr>
<tr>
<td>1/2&quot;</td>
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<td>82</td>
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<tr>
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<td>61</td>
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<tr>
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<td>42</td>
</tr>
<tr>
<td>Total</td>
<td>22.8</td>
<td>100</td>
</tr>
</tbody>
</table>

#### Fine Factor

<table>
<thead>
<tr>
<th>WEIGHTS RETAINED</th>
<th>% RET</th>
<th>SPECs</th>
</tr>
</thead>
<tbody>
<tr>
<td>#10</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>#16</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>#30</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>#40</td>
<td>3</td>
<td>14</td>
</tr>
<tr>
<td>#50</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>#100</td>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>#200</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>#400</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>4.6</td>
<td></td>
</tr>
</tbody>
</table>

**Marshall**

- Marshall + M
- Gyratory = G
- Core = C

**Compaction**

- Sample Max. Dens. (Gmm) = 92.3
- Sample Max. Dens. (Gmm) = 92.3
- Average Bulk Dens. (Gmm) = 82.3
- Air Voids =

**Gyratory**

- Average Relative Density (Gmm) =
- Air Voids =

- Test Operator and Date:
  - M. Twins 05-03-13
  - T. Sawyer 05-03-13

**Remarks**

416 3 1/4 Special Mix (with Rap)
**ARIZONA DEPARTMENT OF TRANSPORTATION**  
**ASPHALTIC CONCRETE TABULATION – IGNITION FURNACE**

<table>
<thead>
<tr>
<th>USE CAPITAL LETTERS</th>
<th>LAB NUMBER</th>
<th>ORG NUMBER</th>
<th>MATL</th>
<th>TYPE</th>
<th>PURPOSE</th>
<th>TEST LAB</th>
<th>SIZE</th>
<th>SIZE %</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEST NO.</td>
<td>LOT OR SUFFIX</td>
<td>SAMPLED BY</td>
<td>MO</td>
<td>DAY</td>
<td>YEAR</td>
<td>TIME</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SAMPLED FROM</td>
<td>LIFT NO.</td>
<td>RDWY</td>
<td>STATION</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ORIGINAL SOURCE</td>
<td>PROJECT ENGINEER / SUPERVISOR</td>
<td>PROJECT NUMBER</td>
<td>TRACS NUMBER</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**FIGURE 5**

### WEIGHTS RETAINED

<table>
<thead>
<tr>
<th>Size</th>
<th>% RET</th>
<th>% PASS</th>
<th>SPECS</th>
</tr>
</thead>
<tbody>
<tr>
<td>$3'$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 1/2'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 1/2'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/4'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/2'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/8'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/4'</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- #8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**RINE FACTOR** = \( \frac{\text{Wt. of Pass #8 Split}}{\text{Pass % of #8 Split}} \)

**INTEGRATED IGNITION FURNACE**

ARIZ. 427T  
ARIZ. 426T

<table>
<thead>
<tr>
<th>SPEC</th>
<th>ARIZ. 427T</th>
<th>ARIZ. 426T</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### COMPACTION

**MARSHALL**

- Average Bulk Density (100 x 100) [pcf]
- Average Bulk Density (2,500 x 80.38) [pcf]

<table>
<thead>
<tr>
<th>SPEC</th>
<th>MARSHALL</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td></td>
</tr>
</tbody>
</table>

### RICE

- Soil Max. (Sample Max. Density)
- Soil Min. (Sample Min. Density)

<table>
<thead>
<tr>
<th>SPEC</th>
<th>RICE</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td></td>
</tr>
</tbody>
</table>

### GYRATORY

- Average Relative Density (％) at 2% Nosing
- Average Relative Density (％) at 4% Nosing

<table>
<thead>
<tr>
<th>SPEC</th>
<th>GYRATORY</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td></td>
</tr>
</tbody>
</table>

### WHITE

- Test Operator and Date

**FIGURE 5**

<table>
<thead>
<tr>
<th>RECEIVED DATE</th>
<th>TEST OPERATOR AND DATE</th>
<th>SUPERVISOR AND DATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>44-9372 R03/13</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
APPENDIX A

DETERMINATION OF GRADATION, MOISTURE CONTENT, AND BINDER CONTENT OF THE RAP MATERIAL

A.1 Obtain a representative sample of the RAP material in accordance with Arizona Test Method 105. When multiple RAP stockpiles are used, a separate representative sample shall be obtained from each stockpile.

A.1.1 The sample shall be split to provide a sufficient amount of material for gradation testing, moisture content testing, and binder content testing.

A.2 The entire split sample of RAP material from each stockpile is dried at 140 °F and the percent moisture content determined as described below.

**Note:** A higher drying temperature is not appropriate because it will soften the binder causing the RAP material to break into smaller particles and adhere to the drying pan.

A.2.1 The weight of the RAP material from each stockpile is determined and recorded to the nearest 0.1 gram.

A.2.2 The material is dried to constant weight at 140 °F ± 10 °F.

**Note:** Drying to constant weight at 140 °F will typically take overnight.

A.2.3 After drying to constant weight at 140 °F ± 10 °F, cover the material and allow to cool 30 ± 10 minutes at room temperature. The weight of the RAP material is then determined and recorded to the nearest 0.1 gram.

A.2.4 The percent moisture content of the RAP material from each stockpile is determined and recorded to the nearest 0.01 percent by the following:

\[
\text{Percent Moisture Content} = \left( \frac{\text{Weight of Material Prior to Drying}}{\text{Weight of Material After Drying}} \right) - \left( \frac{\text{Weight of Material After Drying}}{\text{Weight of Material After Drying}} \right) \times 100
\]
A.3  After drying and determining the moisture content at 140 °F, the RAP material shall be tested for gradation, moisture content (at 290 °F), and binder content.

A.3.1  The gradation of the RAP material from each stockpile shall be determined as described below.

A.3.1.1 Split out a representative sample of the RAP material from each stockpile which conforms to the size specified in Table 3.

<table>
<thead>
<tr>
<th>Maximum Size of Particle (See Note)</th>
<th>Minimum Weight of Sample, grams</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/4 in. and larger</td>
<td>5000</td>
</tr>
<tr>
<td>1/2 in.</td>
<td>2000</td>
</tr>
<tr>
<td>3/8 in.</td>
<td>1000</td>
</tr>
</tbody>
</table>

Note: The smallest sieve opening through which the entire amount of material will pass.

A.3.1.2  Dry sieve the material in accordance with Arizona Test Method 240, with the exception that the No. 8 sieve shall be used as the smallest sieve. (Arizona Test Method 240 limits the time for shaking the sample to 5 minutes ± 15 seconds to control breakdown of the particles of RAP material into smaller size fractions.) The gradation of the RAP material from each stockpile is then determined in accordance with Arizona Test Method 248, Alternate #2.

A.3.2  The percent moisture content of the RAP material from each stockpile shall be determined by drying at 290 °F as described below.

Note: The sample for determining the moisture content at 290 °F shall be obtained at the same time and subjected to the same treatment prior to testing as the sample obtained for determining the RAP binder content.

A.3.2.1 Split out a representative 1000 ± 50 gram sample of the RAP material from each stockpile. The weight of each sample is determined and recorded to the nearest 0.1 gram.

A.3.2.2 Each sample is dried at 290 ± 10 °F to constant weight. Constant weight is defined as the weight at which further drying does not alter the weight more than 0.1 gram at intervals of a minimum of 30 minutes.
A.3.2.3  After drying to constant weight at 290 ± 10 °F, cover the sample and allow to cool 30 ± 10 minutes at room temperature. The weight of the sample is then determined and recorded to the nearest 0.1 gram.

A.3.2.4  The percent moisture content of the RAP material from each stockpile is determined and recorded to the nearest 0.01 percent by the following:

\[
\text{Percent Moisture Content} = \frac{\left( \frac{\text{Weight of Material Prior to Drying}}{\text{Weight of Material After Drying}} \right)}{\left( \frac{\text{Weight of Material Prior to Drying}}{\text{Weight of Material After Drying}} \right)} \times 100
\]

A.3.3  The total percent moisture content of the RAP material from each stockpile is determined by adding the percent moisture content by drying at 140 °F (Subsection A.2) to the percent moisture content by drying at 290 °F (Subsection A.3.2).

A.3.4  The binder content of the RAP material from each stockpile shall be determined as described below.

A.3.4.1  Split out a representative sample of the RAP material from each stockpile which conforms to the size specified in Table 4.

<table>
<thead>
<tr>
<th>Nominal Maximum RAP Aggregate Size (See Notes)</th>
<th>Mass of Sample, grams</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 in.</td>
<td>2500 – 3000</td>
</tr>
<tr>
<td>3/4 in.</td>
<td>2000 – 2500</td>
</tr>
<tr>
<td>1/2 in.</td>
<td>1500 – 2000</td>
</tr>
<tr>
<td>3/8 in.</td>
<td>1200 – 1700</td>
</tr>
<tr>
<td>No. 4</td>
<td>1200 – 1700</td>
</tr>
</tbody>
</table>

Note: The nominal maximum RAP aggregate size is defined as: One size larger than the first sieve that retains more than 10 percent RAP aggregate.

Note: To determine the nominal maximum RAP aggregate size, the RAP aggregate gradation of each RAP stockpile, as shown in the mix design or as determined from previous testing, may provide information.
A.3.4.2  Preheat the ignition furnace to 538 ± 5 °C (1000 ± 9 °F). Do not preheat the sample basket.

A.3.4.3  Weigh and record the mass of the basket assembly to the nearest 0.1 gram.

A.3.4.4  Perform ignition furnace testing on the RAP material from each stockpile in accordance with Subsections 5.6 through 5.13.

A.3.4.5  Calculate and record the binder content of the material from each RAP stockpile, to the nearest 0.01%, as follows:

\[
\%AC = \left( \frac{W_i - W_A}{W_i} \right) \times 100 - %M
\]

Where:
- \( \%AC \) = measured RAP binder content in percent by mass of the sample
- \( W_i \) = mass of the sample prior to ignition
- \( W_A \) = mass of the sample after ignition
- \( %M \) = percent moisture in the sample (determined by drying at 290 °F, per Subsection A.3.2)

A.3.4.5.1  Retain the original printed ticket and save with other test documentation.

A.3.4.6  Each measured binder content test result, as determined in Subsection A.3.4.5, is adjusted by the appropriate RAP binder content correction factor (See Appendix B). This adjustment is made by adding the RAP binder correction factor to each measured binder content. The corrected RAP binder content is recorded to the nearest 0.01%.

A.4  If needed, determine the RAP aggregate gradation by performing a sieve analysis of the residual aggregate in accordance with Section 6.

**Note:** Subsection 6.3.10 does not apply when determining the RAP aggregate gradation.

A.5  Report test information on the "RAP Material Tabulation - Ignition Furnace" laboratory card. An example is shown in Figure 6. A blank "RAP Material Tabulation - Ignition Furnace" laboratory card is shown in Figure 7.
**FIGURE 6**

### **RAP Material Moisture Content @ 140°F**

- Wet Mass of RAP Material: 101.855 g
- Dry Mass of RAP Material: 101.348 g
- Moisture Content @ 140°F: 0.501%

**RAP Material Gradation (ARIZ 240 / ARIZ 248, Alt. #2)**

<table>
<thead>
<tr>
<th>WEIGHTS RETAINED</th>
<th>% RET</th>
<th>% PASS</th>
</tr>
</thead>
<tbody>
<tr>
<td>5&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2-1/2&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-1/4&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1&quot;</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>98</td>
<td></td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>91</td>
<td></td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>61</td>
<td></td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>16</td>
<td></td>
</tr>
<tr>
<td>#4</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>#8</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>3</td>
<td></td>
</tr>
</tbody>
</table>

### **Ignition Furnace**

- Wet Mass of Moisture Sample: 98.93 g
- Dry Mass of Moisture Sample: 98.70 g
- Moisture Content @ 290°F: 0.16%

### **RAP Aggregate Gradation**

**WEIGHTS RETAINED**

<table>
<thead>
<tr>
<th>WEIGHTS RETAINED</th>
<th>% RET</th>
<th>% PASS</th>
<th>% PASS DRY</th>
</tr>
</thead>
<tbody>
<tr>
<td>5&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2-1/2&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-1/4&quot;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1&quot;</td>
<td>100</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>98</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>91</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>61</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#4</td>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#8</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Total RAP Material Moisture Content**

- Moisture Content @ 140°F: 0.501%
- Moisture Content @ 290°F: 0.16%
- Total Moisture Content: 0.66%

**RPM RAP Material Tabulation - Ignition Furnace**

**Test Number:** 13 - 0027

**Sampled By:** Betty Boot

**Sampled From:** Stockpile

**Original Source:** Existing Roadway Millings

**Remark:**

**Example**

**FIGURE 6**
ARIZONA DEPARTMENT OF TRANSPORTATION
RAP MATERIAL TABULATION – IGNITION FURNACE
(Arizona Test Method 428 - Appendix A)

<table>
<thead>
<tr>
<th>USE CAPITAL LETTERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>LAB NUMBER</td>
</tr>
<tr>
<td>ORG NUMBER</td>
</tr>
<tr>
<td>MATL</td>
</tr>
<tr>
<td>TYPE</td>
</tr>
<tr>
<td>PURPOSE</td>
</tr>
<tr>
<td>TEST LAB</td>
</tr>
<tr>
<td>SIZE</td>
</tr>
<tr>
<td>SIZE %</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TEST NO</th>
</tr>
</thead>
<tbody>
<tr>
<td>LOT OR SUFFIX</td>
</tr>
<tr>
<td>SAMPLED BY</td>
</tr>
<tr>
<td>MO</td>
</tr>
<tr>
<td>TIME</td>
</tr>
<tr>
<td>MILITARY TIME</td>
</tr>
</tbody>
</table>

SAMPLED FROM
LIF NU. | HWY |
| STATION |

IF MILEPOST INPUT DECIMAL

ORIGINAL SOURCE
PROJECT ENGINEER / SUPERVISOR
PROJECT NUMBER
TRACS NUMBER

REMARKS

---

**RAP Material Moisture Content @ 140°F**

<table>
<thead>
<tr>
<th>p: Wet Mass of RAP Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>q: Dry Mass of RAP Material</td>
</tr>
<tr>
<td>r: Moisture Content @ 140°F</td>
</tr>
<tr>
<td>[p - q = r] x 100</td>
</tr>
</tbody>
</table>

**Ignition Furnace**

| a: Wet Mass of Moisture Sample |
| b: Dry Mass of Moisture Sample |
| c: Moisture Content @ 230°F |
| \[(a - b) \times 100\] |

**RAP Aggregate Gradation**

<table>
<thead>
<tr>
<th>COURSE FACTOR</th>
</tr>
</thead>
<tbody>
<tr>
<td>WEIGHTS RETAINED</td>
</tr>
<tr>
<td>(3)</td>
</tr>
<tr>
<td>(1)</td>
</tr>
<tr>
<td>(3/4)</td>
</tr>
<tr>
<td>(1/2)</td>
</tr>
<tr>
<td>(3/8)</td>
</tr>
<tr>
<td>(1/4)</td>
</tr>
<tr>
<td>(#4)</td>
</tr>
<tr>
<td>(#8)</td>
</tr>
<tr>
<td>(-#8)</td>
</tr>
<tr>
<td>TOTAL</td>
</tr>
</tbody>
</table>

**Total RAP Material Moisture Content**

| r: Moisture Content @ 140°F |
| c: Moisture Content @ 230°F |
| s: Total Moisture Content (r + c) |
| \(\%\) |

**FIGURE 7**
APPENDIX B

DETERMINATION OF THE RAP BINDER CORRECTION FACTOR

B.1 A RAP binder content correction factor is determined for each RAP stockpile used in the asphaltic concrete mixture.

B.2 At the start of asphaltic concrete production, the first two samples of RAP material from each stockpile are split and tested for asphalt binder content; one split is tested by ignition furnace as described in Subsections A.3.4.2 through A.3.4.5.1, and the other split is tested by solvent extraction in accordance with AASHTO T 164.

Note: Generally, 9000 grams of RAP material from each stockpile will be adequate to obtain the split samples for determining the RAP binder content correction factor.

Note: At the discretion of the Engineer, the RAP binder correction factor may be determined prior to the start of asphaltic concrete production provided representative RAP samples are available.

B.3 The average asphalt binder content determined by ignition furnace is recorded to the nearest 0.001%.

B.4 The average asphalt binder content determined by solvent extraction is recorded to the nearest 0.001%.

B.5 The RAP binder content correction factor is determined by subtracting the average ignition furnace result from the average solvent extraction result. The RAP binder content correction factor is recorded to the nearest 0.01%.

B.6 A new RAP binder correction factor may be determined at any time the Engineer believes it is necessary due to a change in material or other circumstances.

B.7 Report the determination of the RAP binder content correction factor on the “RAP Binder Content Correction Factor” laboratory card. An example is shown in Figure 8. A blank “RAP Binder Content Correction Factor” laboratory card is shown in Figure 9.
### RAP Binder Content Correction Factor
(Ignition Furnace vs. Solvent Extraction)

<table>
<thead>
<tr>
<th>Sample #</th>
<th>Ignition Furnace (ARIZ 428)</th>
<th>Solvent Extraction (AASHTO T 184)</th>
<th>RAP Binder Content Correction Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.81</td>
<td>3.39</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>3.96</td>
<td>3.65</td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>3.885</td>
<td>3.520</td>
<td>(-0.37)</td>
</tr>
</tbody>
</table>

Remarks:

**EXAMPLE**

Received Date: 04-24-13  
Test Operator and Date: M. Tunn  
Supervisor and Date: T. Sauer
ARIZONA DEPARTMENT OF TRANSPORTATION
RAP BINDER CONTENT CORRECTION FACTOR
(Arizona Test Method 428 - Appendix B)

Project Number: ________________________________

TRACS Number: __________________________________

RAP Material Type: ____________________________________

Sample # : _______ Sampled By: ___________________________ Sampled From: ________________________________
Date Sampled: ___________________________ Time Sampled: ________________________________

Sample # : _______ Sampled By: ___________________________ Sampled From: ________________________________
Date Sampled: ___________________________ Time Sampled: ________________________________

<table>
<thead>
<tr>
<th>Sample #</th>
<th>RAP Binder Content (%)</th>
<th>RAP Binder Content Correction Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ignition Furnace (ARIZ 428)</td>
<td>Solvent Extraction (AASHTO T 164)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Average Solvent Extraction Value) Minus (Average Ignition Furnace Value)</td>
</tr>
</tbody>
</table>

Average

Remarks:

---

Received Date

Test Operator and Date

Supervisor and Date

44-9380 03/13

FIGURE 9
Field Shear Vane Test for Cold Recycled Asphalt

(Modification of Utah Shear Vane Test)

1. SCOPE

1.1 This method covers a method for performing shear vane testing on cold recycled asphalt, whether cold in-place recycled or cold central plant recycled, to determine the progress of curing after compaction.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

2. APPARATUS

2.1 Sledge Hammer – A minimum of 5 lbs

2.2 Torque Wrench – Capable of recording torque to 0.5 ft-lb increments within the range of 25 to 150 ft-lb

2.3 Shear Vane – See Figures 1 and 2

2.4 15/16 Inch Socket with a 1/2 Drive

2.5 Protective Shear Vane Cover (optional) – A metal cover of any configuration used during the shear vane driving process which facilitates hammering and protects the shear vane during hammering.
3. PROCEDURE

3.1 Select a test location according to project specifications and at least 1 foot from the edge of the cold recycled asphalt and other test sites.

3.2 Using the hammer, drive the shear vane into the cold recycled asphalt, keeping the shear vane vertical, until the washer sits flush on the surface.

Note: The socket or a protective shear vane cover must be on the bolt head before striking with the hammer so that the shear
vane is not damaged. Care must be taken to keep the device as vertical as possible to minimize damage to the cold recycled asphalt before testing.

3.3 Place the torque wrench onto the head of the shear vane bolt. Apply slight downward pressure to ensure that the shear vane does not lift during the test.

3.4 Evenly apply pressure to the torque wrench (90 degrees in 10 seconds) while watching the dial closely. Apply increasing pressure until the cold recycled asphalt is broken loose by the shear vane.

3.5 Record the highest torque value reached during test.

3.6 Determine and record the pavement temperature at two inches below the surface with an infrared thermometer.

3.7 The test will leave a hole in the mat. Repair by pressing loose material into the hole and coating the location with emulsion.

4. REPORT

4.1 Record the following information:

4.1.1 Time and date of the test

4.1.2 Location including stationing and offset

4.1.3 Maximum torque value achieved in ft-lb

4.1.4 Pavement temperature at a depth of two inches
Determining In-Place Flow of Cold Recycled Asphalt Using the Marshall Hammer

1. **SCOPE**

1.1 This procedure provides a quick evaluation of flow characteristics of cold recycled asphalt, whether cold in-place recycled or cold central plant recycled, to determine appropriate hold time for cold recycling mixes before finishing compactive efforts and releasing to traffic.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

2. **APPARATUS**

2.1 Marshall Hammer - A Marshall compaction hammer meeting the dimensional requirements of Arizona Test Method 410.

2.2 Caliper or Tape Measure – Capable of measuring depth and measuring in millimeters.

3. **PROCEDURE**

3.1 This test is to be performed on an initially compacted cold recycled mat.

3.2 Place the Marshall hammer with its head flat on the mat. Do not move the hammer or rock the head.

3.3 Pick up the sliding weight until it reaches the upper stop. Drop the weight. Repeat 50 times. The operator shall hold the handle by one hand so that the axis of the compaction hammer is as nearly perpendicular to the base of the mold assembly as possible while compaction is accomplished. Care shall be taken not to add body weight to the hammer by leaning or pressing down on the hammer. No mechanical device of any kind is to be used to restrict movement of the handle during compaction. Compaction shall be performed at a minimum rate
of 40 blows per minute. The compaction hammer shall apply only one blow with each fall, that is, there shall not be a rebound impact.

3.4 All measurements are from the level of the undisturbed mat.

3.4.1 Measure the depth of the depression made by the hammer, in mm. The depth is determined by the following equation.

Depth of Depression = \( A - B \)

Where:

- \( A \) = Distance between Straightedge and Bottom of Depression
- \( B \) = Distance between Straightedge and Undisturbed Mat

3.4.2 Measure the height of lateral deformation, if any, in mm. The height of lateral deformation, measured from the level of the undisturbed mat, is represented as “\( B \)” in Figure 1.

3.5 Determine the moisture condition of the mat; whether there is water bleeding from the mat, or if it is dry. Visible water on the surface of the mat, which is not from rolling operations, may indicate possible bleeding from the mat.

3.6 Determine temperature of the mat to the nearest 1 degree F.

4. REPORT

4.1 Record the following information:

4.1.1 Time and date of the test

4.1.2 Depth of depression, in mm.

4.1.3 Height of lateral deformation, in mm.
4.1.4 Mat temperature

4.1.5 Location of the test including stationing and offset
SERIES 500
BITUMINOUS MATERIALS
PERCENTAGE OF UNCOATED PARTICLES USING ASPHALT EMULSIONS

(An Arizona Method)

1. SCOPE

1.1 This method describes a procedure for determining percentage of uncoated particles based on the failure of emulsions to coat a specific mixture of Standard Sand and Type III portland cement.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Mixing Bowl or Pan - A mixing bowl or round bottom pan with a surface which is non-reactive with the emulsion (e.g., stainless steel, glass, or plastic) of approximately 3 liter (3 quart) capacity.

2.3 Mixing Tool - A stainless steel spoon approximately ten inches long.

2.4 Thermometer - A thermometer accurate to 1 °F.

2.5 Balance - A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the
readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.6 Stopwatch or other timing device.

2.7 No. 20 sieve conforming to the requirements of ASTM E11.

3. MATERIALS

3.1 Standard Sand (20-30) conforming to ASTM C778.

3.2 Type III Portland cement conforming to ASTM C150 and having a minimum specific surface of 1900 cm²/gram.

4. PROCEDURE

4.1 Thoroughly mix a representative sample of the emulsion to be tested. If any large particles are present, the emulsion shall be strained through a single layer of damp cheesecloth or the No. 20 sieve.

4.2 Screen approximately 500 grams of the Standard Sand through the No. 20 sieve.

4.3 Weigh 461 ± 0.1 grams of the screened sand into the tared mixing bowl or pan.

4.4 Add 4 ± 0.1 grams of the cement to the sand.

4.5 Mix the sand and cement for one minute, using the mixing spoon to obtain a uniform mixture.

4.6 Bring the emulsion to a temperature of 77 ± 3 °F.

4.7 Weigh 35 ± 0.1 grams of the emulsion into the sand-cement mixture.

4.8 Mix vigorously with the spoon for 2-1/2 minutes, using a stirring and kneading motion.

4.9 Empty the contents of the mixing bowl or pan onto absorbent paper so that the mixture covers an area approximately 10 inches in diameter.
4.10 Allow to dry thoroughly (approximately two hours).

4.11 Pour the dried mixture into the No. 20 sieve and shake and vibrate the sieve until all of the uncoated particles have passed through the sieve.

4.12 Pour sand which has passed the No. 20 sieve into a tared pan and weigh to the nearest 0.1 gram.

5. **CALCULATIONS**

5.1 Calculate the percent of uncoated sand using the following formula:

\[ U = \frac{W_p}{465} \times 100 \]

Where:
- \( U \) = Percent of uncoated sand
- \( W_p \) = Weight of Sand passing No. 20 sieve.

6. **REPORT**

6.1 Report percentage of uncoated sand to the nearest one percent.
VACUUM RECOVERY OF ASPHALT EMULSION RESIDUE
(An Arizona Method)

Scope

1. This method describes a low temperature vacuum procedure for recovery of the asphalt residue from asphalt emulsions. It is not suitable for quantitative recovery of solvents from emulsions containing low boiling range distillates.

Apparatus

2. The apparatus shall consist of the following:
   (a) Brass stirring rod.
   (b) 8 oz. ointment can.
   (c) 100 ml. stainless steel beaker.
   (d) No. 20 sieve conforming to AASHTO designation M 92.
   (e) Vacuum recovery apparatus as shown assembled in Fig. 1.

   1) Vacuum source capable of producing an absolute vacuum within the system of approximately 710 mm (28 in.) mercury.

   2) Thermometer — shall have a range of -5° to +200°C (23°F to 392°F). The overall length shall be 600 mm (24 in.) and the distance from the bottom of the bulb to the zero point shall be 300 mm (12 in.)

   3) Stirrer hot plate.

![Diagram of Vacuum Recovery Apparatus](image-url)
4) Teflon covered stirring bar.
5) 1000 ml. and two 500 ml. filtering flasks with
tubulation.
6) 300 mm Allihn condensor.
7) Portable heat gun.
8) Miscellaneous rubber stoppers and
connecting hose.
9) Vacuum release clamp.

(f) Additional vacuum recovery apparatus for
alternate test procedure.
1) 8" waterbath — hemispherical.
2) 500 ml. Separatory Funnel — pear shape.

Material
3. Isopropyl alcohol, technical grade.

Sample Preparation
4. Emulsion shall be thoroughly stirred and
strained through a No. 20 sieve just prior to use.

Test Procedure
5. (a) Pour 75°C ± 25 ml. of the strained emulsion
into a 1000 ml. stainless steel beaker.

(b) Cool the emulsion to approximately 25°C
(77°F).

(c) Add isopropyl alcohol (approximately 25 to
100 ml.) in sufficient quantity to break the emulsion.

(d) Knead lumps with brass stirring rod against
the sides of the beaker to expel the water from the
broken asphalt.

(e) Apply a film of soap around neck of the 1000
ml. vacuum flask to prevent asphalt from sticking to the
flask.

(f) Wet hands thoroughly with distilled water.
Remove the stirring rod on which the broken asphalt
emulsion (now asphalt) is clinging, and with the wet
hands gently squeeze out all entrained water while
washing asphalt under a constant flow of distilled
water. Form asphalt into a rope and drop into the 1000
ml. flask containing the Teflon stirring bar. (Amount of
asphalt to be added should be enough to cover bottom
of flask approx. 1" in depth). Wash soap off of flask and
rinse soap from sample and flask with distilled water
making sure water is drained from flask before it is
placed on hot plate.

(g) Insert the stoppered thermometer (positioned
in the stopper at an angle to prevent contact with
stirring bar) into the flask and set on hot plate at a
medium high heat setting (#4). The bulb of the
thermometer should be 6 mm (¼ inch) from the bottom
of the flask.

(h) Turn on the water to the condenser, start the
stirring bar and open the vacuum line to establish a
vacuum of approximately 710 mm (28 in.) mercury in
the apparatus.

(i) Turn on the portable heat gun to medium
high heat (#6) and use wherever needed to keep
condensation from forming inside the vacuum flask.

(j) Allow temperature to rise until it reaches
approximately 90°C, turn hot plate down to setting #2
and then allow the temperature of the residue to rise
until it reaches 115°C ± 3°C (239°F ± 5°F). Hold at
this temperature for at least 1 minute (if possible).

NOTE: The hot plate temperature will need to be
regulated during heating of residue to allow for
complete removal of water. (This removal is complete
when there is no longer foaming in the residue.) If the
115°C temperature is reached and there is still water
present, the flask should be removed from the hot plate,
allowed to cool a short time, and then heating resumed
at a reduced temperature. As the residue heats up, soap
bubbles will form up into the vacuum line and condense
in the filtering flask. Some emulsions may react
violently between 60°C (140°F) and 105°C (221°F)
and rise up in the flask. Use the vacuum release clamp as
often as needed to control the asphalt. DO NOT let
asphalt escape into the vacuum line.

(k) Turn off vacuum and slowly release vacuum
through apparatus using release clamp. Stop stirrer,
remove thermometer, and pour residue into a 8 oz.
ointment can. Residue is now ready for specified tests.

Alternate Test Procedure
6. This procedure is recommended for use with
emulsions that will not break using isopropyl alcohol.

(a) Pour 350 ml. to 400 ml. of the strained
emulsion into the separatory funnel.

(b) Place the stoppered funnel into the 1000 ml.
flask and carefully set the unit into the preheated water
bath (Setting #6).

(c) With the water circulating through
condenser, open the vacuum line to establish a vacuum
of approximately 710 mm (28 in.) mercury in the
apparatus.
(d) Start the stirring bar at slow speed and slowly add the emulsion at a rate which allows the emulsion to break as it leaves the funnel. (This is indicated by the presence of foaming at the end of the funnel.

CAUTION: If the emulsion is added to the flask too rapidly, it may react violently and rise up in the flask. If this happens quickly return addition of emulsion to proper rate and use the vacuum release clamp to prevent asphalt from entering into the vacuum line.

(e) Continue adding emulsion to flask until all of it is removed from the funnel.

(f) Close the vacuum line and slowly release vacuum from unit. Stop the stirring bar.

(g) Remove the separatory funnel and the water bath. Wipe bottom of flask dry, insert the thermometer as in 5 (g) and follow original method to completion.
ASPHALT REJUVENATING AGENT RESIDUE
INSOLUBLE IN PETROLEUM ETHER

(An Arizona Method)

SCOPE

1. (a) This method of test is intended for determining the percent of insoluble matter in an asphalt rejuvenating agent using petroleum ether.

   (b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

   (a) Erlenmeyer flask, 125 mL capacity, with stopper.

   (b) Drying oven, adequately ventilated, capable of maintaining a temperature of 110 ± 5 °C (230 ± 9 °F).
(c) Desiccator of adequate size to hold several crucibles.

(d) Analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.001 gram.

(e) Filtering apparatus, including:

1. Vacuum pump
2. Filtering flask, 250 mL capacity
3. Filter tube with molded rubber gasket to hold Gooch crucible.
4. Rubber tubing.
5. Glass fiber pad conforming to AASHTO T 44.
6. Gooch crucible, 4 cm high, 3.5 cm top diameter and 2.1 cm bottom diameter, prepared in accordance with AASHTO T 44.

MATERIALS

3. Petroleum either, 30 to 60 °C, reagent grade.

CAUTION: Always work under a hood, avoid breathing petroleum ether fumes, and keep material away from flame. It is highly flammable and toxic.

TEST PROCEDURE

4. (a) Measure approximately 1 gram of asphalt rejuvenating agent residue into the tared Erlenmeyer flask and weigh to the nearest 0.001 gram.

(b) Pour half of a 100 mL portion of petroleum ether into the flask and swirl the contents until all the residue has dissolved.

(c) Add the balance of the 100 mL portion of petroleum ether and swirl the flask to mix the contents thoroughly. Place a stopper in the flask and set aside for 30 minutes.
(d) Place the tared Gooch crucible and filter mat into position on the filtering flask. Wet the mat with a small portion of petroleum ether.

(e) Apply light suction as decantation through crucible is begun. Use suction sparingly because it tends to clog the filter mat. Exercise utmost care in making the filtration to avoid stirring up any precipitate, in order that the filter mat may not be clogged and that the first decantation may be as complete as possible.

(f) Wash the flask with petroleum ether and pour the bulk of the insoluble material on the filter mat after the crucible has drained. Transfer all insoluble material from the flask to the crucible, using a rod with flattened rubber tip (a.k.a. "policeman") if necessary. Rinse the flask and policeman thoroughly with petroleum ether.

(g) Wash the insoluble matter in the crucible until the filtrate is substantially colorless, then apply strong suction to remove the remaining petroleum ether.

(h) Remove crucible from flask and wipe bottom of crucible clean.

(i) Dry the crucible in a 110 ± 5 °C (230 ± 9 °F) oven for 15 to 20 minutes, cool in desiccator for 15 to 20 minutes and weigh to the nearest 0.001 gram.

NOTE: The crucible shall be cooled in the desiccator for the same amount of time that it is dried in the oven.

CALCULATIONS

5. Calculate the percentage of the total sample which is insoluble in petroleum ether. The formula is as follows:

\[
\text{Insoluble Residue, percent} = \frac{A}{B} \times 100
\]

Where: \( A \) = total weight insoluble

\( B \) = total weight of sample

REPORT

6. Report the total insoluble residue to the nearest 0.01 percent.
RAPID DETERMINATION OF ASPHALTENES AND CHEMICAL REACTIVITY OF ASPHALTS

(An Arizona Method)

SCOPE

1. (a) This method describes a procedure for the determination of Asphaltene, (A), content and Chemical Reactivity Ratio, (CRR), of asphalts. It is not intended to supplant the complete asphalt analytical methods, but should be used when only the A and CRR values are needed.

(b) This test method involves hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

(a) Erlenmeyer flasks with glass stoppers, 125 mL capacity.

(b) Analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.0001 gram.

(c) Glass funnels, 75 mm top diameter.

(d) Hot plate, with variable heat control.
(e) Beakers, Griffin type, 250 mL capacity.
(f) Funnel support.
(g) Infra-red heat lamp, variable intensity.

MATERIALS

3. (a) Filter paper, Whatman Glass Microfibre, 12.5 cm, Grade GF/A.
(b) Normal pentane, (nC₅H₁₂), practical grade.
(c) Petroleum Ether, 30 to 60 °C, reagent grade
(d) Concentrated Sulfuric Acid, reagent grade, (H₂SO₄ Assay: 95-98%)
(e) Sodium Hydroxide, pellets, reagent grade.

CAUTION: Pentane and Petroleum Ether are extremely flammable solvents. Smoking or open flames should not be permitted in the area while they are in use. With the exception of weighing, all operations should be carried out in a functioning fume hood. Protective goggles, gloves and laboratory apron shall be worn.

PROCEDURE

4. (a) Weigh approximately 1 gram of the asphalt into a tared 125 mL Erlenmeyer flask. Weigh to the nearest 0.1 mg and record as S₁ (weight of sample and flask).

(b) Add 75 mL of normal pentane to the flask and dissolve the asphalt completely. Frequent shaking during the first 15 minutes will aid in dissolving the asphalt. If it does not dissolve, warm cautiously until the asphalt is in solution.

(c) Allow the solution to settle overnight.

(d) Stir vigorously with a glass stirring rod to initiate precipitation, and allow to settle until clear.
(e) Carefully decant liquid through filter paper into a 250 mL beaker, using the Glass Microfibre filter paper. Wash the precipitate remaining in the flask 3 times with 15-20 mL portions of pentane, each time decanting liquid into filter paper. Thoroughly wash flask with pentane and add washings to filter paper. Thoroughly wash precipitate on filter paper with pentane until discoloration is removed from the filter paper. Discard paper containing the precipitate.

NOTE: It is very important that the filter paper be kept wet throughout the washing and filtering process in step (e).

(f) Thoroughly wash and dry the flask used in this operation. Methylene Chloride washing is recommended.

(g) Using the heat lamp, evaporate the solution in the beaker until approximately 25 mL remains. Pour solution into the 125 mL flask previously used. Wash the beaker with pentane until clean, adding the washings to the flask.

(h) Evaporate the contents of the flask with the heat lamp until all the solvent has been removed. Then place on the hot plate and heat cautiously until a white vapor is noticed at the top of the flask.

(i) Remove the flask from the hot plate. Cool and weigh to the nearest 0.1 mg. Record this weight as \( B_p \) (weight of flask and residue).

(j) Add 75 mL of Petroleum Ether to the flask containing the residue and dissolve ALL the residue.

(k) Pour the solution into a 250 mL beaker and thoroughly wash the flask with Petroleum Ether. Add the washings to the solution in the beaker. If necessary, reduce the volume of the solution to 100 - 125 mL by evaporation with the heat lamp.

(l) Thoroughly wash and dry the flask. Methylene Chloride washing is recommended.

(m) Slowly add 10 ± 0.5 mL concentrated sulfuric acid to the solution in the beaker while stirring vigorously. Continue stirring for at least 3 minutes.

(n) Allow the solution to settle until clear.
Cover bottom of a 250 mL beaker with Sodium Hydroxide pellets, then immediately filter the liquid through a filter paper into the 250 mL beaker containing the Sodium Hydroxide pellets. Do not allow the sludge to leave the beaker. Wash the sludge in the beaker 3 times, using 15 - 20 mL portions of Petroleum Ether. Filter the washings into the pellets in the beaker, then discard filter paper.

CAUTION: Do not touch the pellets with bare hands.

CAUTION: Sludge contains large amounts of unreacted acid. Contact with Sodium Hydroxide pellets may cause explosion.

Using the heat lamp, evaporate this solution to approximately 50 mL, then filter into the 125 mL Erlenmeyer flask previously used, thoroughly washing beaker and pellets into filter paper, using a 30-40 mL portion of Petroleum Ether.

Discard the filter paper containing the Sodium Hydroxide pellets.

Evaporate the contents of the flask with heat lamp until all the solvent has been removed. Then place on the hot plate and heat cautiously until a white vapor is noticed at the top of the flask.

Remove the flask from the hot plate. Cool and weigh to the nearest 0.1 mg. Record this weight as "C" (weight of flask and residue).

CALCULATIONS AND REPORT

5. (a) Percent Asphaltenes:

\[ A = \frac{S_f - B_p}{S} \times 100 \]

Where:  
- \( S \) = Weight of sample, \( S_f - T_f \)  
- \( S_f \) = Weight of sample and flask, grams  
- \( T_f \) = Tare weight of flask, grams  
- \( A \) = Percent Asphaltenes.  
- \( B_p \) = Weight of flask and residue from Pentane washing, grams
(b) Percent Unreacteds:

\[ U = \frac{C - T_f}{S} \times 100 \]

Where:  
- \( U \) = Percent Unreacteds.  
- \( C \) = Weight of flask and residue from Petroleum Ether washing, grams  
- \( S \) = Weight of Sample, grams  
- \( T_f \) = Tare weight of flask, grams  

(c) CRR (Chemical Reactivity Ratio) Value:

\[ CRR = \frac{100 - (A + U)}{U} \]

Where:  
- \( CRR \) = Chemical Reactivity Ratio  
- \( A \) = Percent Asphaltenes  
- \( U \) = Percent Unreacteds.  

(d) Report Asphaltenes as a percentage of the asphalt sample, to the nearest 0.01%.

(e) Report CRR Value as a ratio of the Reacted and Unreacted asphalt components, to the nearest 0.01 unit, if this value is required.
FLOW SHEET - MODIFIED ROSTLER-STERNBERG ANALYSIS

NOTE:

A = Asphaltenes  Insoluble in nC5H12
*N1 = 1st Nitraffins Precipitated by concentrated H2SO4
*N2 = 2nd Nitraffins Precipitated by concentrated H2SO4
A1 = 1st Acidaffins Precipitated by concentrated H2SO4
A2 = 2nd Acidaffins Unreacted by concentrated H2SO4
P = Paraffins    Unreacted by concentrated H2SO4

* 1st and 2nd Nitraffins are Nitrogen Bases
RESIDUE BY EVAPORATION

(An Arizona Method)

1. **SCOPE**

1.1 This method describes a rapid procedure for determining the percent of asphaltic residue in all types of emulsified bituminous materials.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

2. **APPARATUS**

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Small glass beaker or similar container.

2.3 Ointment tins [180 mL (6 oz.)].

2.4 Hot plate capable of maintaining temperatures of 163 °C (325 °F) maximum.

2.5 Glass or metal stirring rod.
2.6 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.01 gram.

3. PRECAUTION

3.1 Care must be exercised in the use of apparatus and the handling of the tin so that material is neither lost nor additional material picked up. The use of the ointment tin lid serves as well as a clean place for resting the tin.

4. PROCEDURE

4.1 Thoroughly mix the sample using stirring rod.

Note: It is very important that the sample be completely mixed, making certain to mix material clinging to bottom and sides of container into sample. A sample that has "separated" may be rendered homogeneous through continued stirring. Never mix the sample by shaking. If a sample is received which has "broken", it shall be discarded and another sample obtained for testing.

4.2 Pour approximately 25 grams of the mixed material into the glass beaker or similar container.

4.3 Record weight of a 180 mL (6 oz.) ointment tin to the nearest 0.01 gram and place 5 ± 0.3 grams of material into the tin. Record weight of material in tin to the nearest 0.01 gram, as "A".

4.4 Repeat procedure in Subsection 4.3 for 2 additional samples.

4.5 Place the three samples on hot plate and slowly heat to a temperature that will prevent spattering or overcooking. (See Note in Subsection 4.7)
4.6 Heat samples at this temperature until the residue stops bubbling and appears smooth. This is an indication that the sample is nearing the end point (complete removal of water).

**Note:** Air bubbles should be removed occasionally by tapping tin on the hot plate or other hard surface and rotating the sample around the bottom of the tin. (Use of tongs or needle-nose pliers to handle tin is recommended.)

4.7 Raise the temperature of samples to 163 °C (325 °F) maximum. This may be accomplished by raising the temperature of the hot plate or by placing samples on another hot plate. If an additional hot plate is used and bubbling occurs the samples shall be placed back on the lower temperature hot plate for a short period of time and then returned to the higher temperature hot plate.

**Note:** The temperatures required vary with the type of material being tested. When testing the "ERA" grade materials and emulsions with low viscosities the temperatures used will normally be near the boiling point of water. When testing higher viscosity materials the temperatures may approach the 163 °C (325 °F) maximum allowed. Care must be exercised for all materials to prevent spattering and overheating.

4.8 Heat the samples until the first indication of smoking is detected, which shall be the determination of the removal of all water. The use of a black background is useful in observing the point of smoking.

**Note:** When testing "ERA - 1" grade materials, the determination of end point described in Subsection 4.8 shall not be used, rather when the material is free of all bubbles and has a completely translucent appearance.

4.9 Remove samples from hot plate and allow to cool.

4.10 Record the weight of each sample to the nearest 0.01 gram, as "B".
5. **CALCULATION**

5.1 Determine percent residue of each sample by the equation below, and record results to the nearest 0.1%:

\[
\text{Percent Residue} = \frac{B}{A} \times 100
\]

Where:  
A = Weight of Sample Before Heating  
B = Weight of Sample After Heating

Example:

\[
\text{%Residue} = \frac{3.18}{5.02} \times 100 = 63.35\%
\]

5.2 Determine the average percent residue of the material being tested by the equation below, and record the results to the nearest 0.1%.

\[
\text{Average %Residue} = \frac{(\text{Sum of the 3 individual samples Percent Residue})}{3}
\]

Example:

(For 3 samples of 63.4 %, 63.5%, and 63.7% residue)

\[
\text{Average %Residue} = \frac{(63.4 + 63.5 + 63.7)}{3} = 63.5\%
\]

Reported Example = 64%

6. **REPORT**

6.1 The average percent residue shall be reported to the nearest whole percent.
SERIES 700
CHEMICAL AND SPECIALTY
TESTING OF PAINT, VARNISH, LACQUER, 
AND RELATED MATERIAL 

(A Modification of Federal Test Method Standard 141)

SCOPE

1. (a) This group of tests describes the procedures for routine chemical 
and/or physical testing of paints and the raw materials used in the manufacture of these 
organic protective coatings for surface preservation or marking of wood, metal, and/or 
masonry installations.

(b) This test method may involve hazardous material, operations, or 
equipment. This test method does not purport to address all of the safety concerns 
associated with its use. It is the responsibility of the user to consult and establish 
appropriate safety and health practices and determine the applicability of any regulatory 
limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information 
regarding the procedure to be used for rounding numbers to the required degree of 
accuracy.

(d) Metric (SI) units and values are shown in this test method with 
English units and values following in parentheses. Values given for metric and English 
units may be numerically equivalent (soft converted) for the associated units, or they 
may be given as rounded or rationalized values (hard converted). Either the metric or 
English units along with their corresponding values shall be used in accordance with 
applicable specifications. See Appendix A2 of the Materials Testing Manual for 
additional information on the metric system.

APPARATUS & REAGENTS

2. (a) Requirements for the frequency of equipment calibration and 
verification are found in Appendix A3 of the Materials Testing Manual. All apparatus 
shall conform to those specifications given in the respective tests.
(b) All reagents shall conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society, where such specifications are available.

GENERAL

3. (a) Consistency - The determination of consistency of paint shall be as specified in ASTM Designation D 562, "Consistency of Paints Using the Stormer Viscosimeter."

(b) Drying Time - Determine the dry-to-touch and dry-hard times of paints by following the procedures specified in ASTM Designation D 1640, "Drying, Curing, or Film Formation of Organic Coatings at Room Temperature."

(c) Pigment Content - Follow ASTM Designation D 2371, "Pigment Content of Solvent-Reducible Paints," to determine percentage by weight. The remaining percentage may be considered as vehicle.

(d) Vehicle Composition - Follow ASTM Designation D 2621, "Infrared Identification of Vehicle Solids from Solvent-Reducible Paints."

(e) Density, kilograms per liter (pounds per gallon)

1) The apparatus shall consist of a smoothly finished aluminum, brass, stainless steel or plated cup provided with a snug-fitting, plug type cover having a small hole in its center. The capacity of the cup at 25.0 ± 0.5 °C shall be 83.2 ± 0.1 mL. The inside of the bottom of the cup shall be rounded. A convenient size for the apparatus is about 76 mm high by 38 mm in diameter. The cup shall be tared.

2) Pour the well-mixed paint, brought to a temperature of 25.0 ± 0.5 °C, into the cup until it is nearly full, put on the cover, rotate it firmly into place. Wipe off excess paint that exudes through the hole. Determine the weight of the sample to the nearest 0.1 gram. Multiply this weight by 0.012 to obtain the density in kilograms per liter (divide the weight of the sample by 10 to obtain the density in pounds per gallon).

3) This same procedure is applied for paste and semipaste, with care that all air pockets are removed from the material.
RAW MATERIALS, PIGMENTS

4. (a) White Pigments - Follow ASTM Designation D 34, "Chemical Analysis of White Pigments."

(b) Magnesium Silicate Pigments - Follow ASTM Designation D 717, "Analysis of Magnesium Silicate Pigment."

(c) Diatomaceous Silica - Follow ASTM Designation D 719, "Analysis of Diatomaceous Silica Pigment."

(d) Dry Red Lead - Follow ASTM Designation D 49, "Chemical Analysis of Dry Red Lead."

(e) Iron Oxide - Follow ASTM Designation D 50, "Chemical Analysis of Yellow, Orange, Red, and Brown Pigments Containing Iron and Manganese." Refer to Section 11 and 12 of same.

ALUMINUM PASTE


(a) Coarse Particles

(b) Easily Extracted Fatty and Oil Matter

(c) Nonvolatile Matter
SAMPLING AND SIEVING OF CRUMB RUBBER

(An Arizona Method)

1. SCOPE

1.1 This method describes the procedure for sampling and determining the sieve analysis of crumb rubber.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2 Sieves for fine screening conforming to ASTM E11, with a diameter of 8 inches, nested in sizes: #8, #10, #16, #30, #40, #50, #100, #200, and bottom pan.

2.3 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 grams.

2.4 Oven capable of maintaining a temperature of 140 ± 5 °F.

2.5 A small riffle splitter with one-half inch openings.

2.6 Mechanical sieve shaker which imparts a vertical, or lateral and vertical, motion to the sieves, causing the particles thereon to bounce and turn so as to present different orientations to the sieving surface.
2.7 Splitting blanket of adequate size for approximately 1500 grams of crumb rubber.

2.8 Talc (Talcum Powders) – Usually some mixture of magnesium silicate, silica, magnesium oxide, or magnesium-aluminum silicate. 100 percent of the particles shall pass a No. 200 sieve.

2.9 A one-quart glass jar with large opening, and lid.

2.10 Scoop, shovel, “sample thief” or other suitable device for obtaining samples.

3. **SAMPLING**

3.1 Samples shall be obtained at the hot plant or CRA (Crumb Rubber Asphalt) blending unit from bags selected by the Engineer.

3.2 The sample shall be obtained from the top of a randomly selected bag of crumb rubber, taken at a minimum of three random locations at least 3” below the surface, using a scoop, shovel, “sample thief”, or other suitable device. The combined sample size shall be approximately 1500 grams.

3.3 If the contractor suspects that segregation of the crumb rubber has occurred in the bag, the contractor may propose that the sample be obtained in a method which differs from that specified in Subsection 3.2 above. The proposed method of sampling must be approved by the Engineer and must render a representative sample. The sample shall be obtained by the contractor and witnessed by the Engineer.

4. **SAMPLE PREPARATION**

4.1 Dry the approximate 1500 gram sample to constant mass at a temperature of 140 ± 5 °F.

4.2 Spread the sample on the splitting blanket and break up clumps so that the material is free flowing.

4.3 Mix the material thoroughly by turning the entire sample over several times in the splitting blanket.
4.4 Obtain a representative 100 ± 10 gram test sample of the crumb rubber by splitting the entire sample through the splitter.

4.5 Place the test sample in the jar, breaking up any clumps of material. Weigh the sample to the nearest 0.1 gram and record as “Preweight of Crumb Rubber”.

4.6 Add 5.0 grams of talc to the jar. Place the lid on the jar and mix sample thoroughly by manually shaking until the rubber and talc are uniformly mixed (a minimum of one minute). Weigh the combined rubber and talc to the nearest 0.1 gram and record as “Preweight of Crumb Rubber and Talc”. Do not record this weight as “Corrected Dry Weight”.

4.7 Empty the sample of combined rubber and talc into the nest of sieves, thoroughly removing any material clinging to the jar and lid.

5. PROCEDURE

5.1 Subject the test sample to mechanical sieving for 10 ± 0.5 minutes. In no case shall fragments be turned or manipulated through the sieves by hand. If there are clumps of material retained on any of the screens, break these apart, and recombine the rubber and talc from all the sieves and bottom pan; place the material in the jar, reshapk, reweigh (recording new weight of combined rubber and talc), and resieve.

5.2 Determine the weight retained on each sieve and the bottom pan and record to the nearest 0.1 gram. Any material adhering to the bottom of a screen shall be brushed into the next finer screen.

5.3 Do not discard any of the sieved material until the sum of the individual weights retained is compared to the weight of the combined rubber and talc prior to sieving (“Preweight of Crumb Rubber and Talc”). If the difference between the two weights is less than or equal to 1.0% of the “Preweight of Crumb Rubber and Talc”, an adjustment in weight shall be made on the sieve which has the largest weight retained, except no adjustment shall be made on the minus #200 material. If the difference is greater than 1.0%, discard this test sample and obtain another 100 ± 10 gram sample as specified in Subsection 4.4 and repeat the test.
6. **CALCULATION**

6.1 If the weight of minus #200 material from sieving is 5.0 grams or less, record a zero as the minus #200 weight corrected for talc.

6.2 If the weight of minus #200 material from sieving is greater than 5.0 grams, subtract 5.0 grams from it and record this difference as the minus #200 weight corrected for talc.

**NOTE:** The adjustments made in Subsections 6.1 and 6.2 above are to account for the 5.0 grams of talc added to the sample.

6.3 After determining the minus #200 weight corrected for talc as described above, determine the “Corrected Dry Weight” by adding the weights retained on sieve size #8, #10, #16, #30, #40, #50, #100, #200 and the minus #200 corrected for talc. Record this sum as the "Corrected Dry Weight".

6.4 Determine the sieve factor for calculating the sieve analysis by dividing 100 by the "Corrected Dry Weight". Record the factor to at least six decimal places.

6.5 The percent passing for each sieve is determined by multiplying the weight retained on that sieve by the factor, and subtracting the result from the unrounded % passing the next larger sieve. Values for "weight retained times the factor" and "percent passing each sieve" shall be determined and used in the calculations to at least six decimal places. The percent passing value for each sieve is recorded in the sieve analysis to the nearest whole percent, except the percent passing the #200 sieve (corrected for talc) is recorded to the nearest 0.1 percent.

6.6 As a check on the sieve analysis, the weight of material passing the #200 (corrected for talc) is multiplied by the factor. The result of this calculation, rounded to the nearest 0.1 percent, should be the same as the value for the percent passing the #200 determined in the paragraph above.

6.7 If desired, obtain the percent retained on each sieve by subtracting the rounded % passing value for that sieve from the rounded % passing value for the next higher sieve.
7. **EXAMPLE**

7.1 Figure 1 is an example of the calculations when the weight of minus #200 material is 5.0 grams or less.

7.2 Figure 2 is an example of the calculations when the weight of minus #200 material exceeds 5.0 grams.

7.3 A blank Crumb Rubber Tabulation laboratory card is provided in Figure 3.
**EXAMPLE**

PREWEIGHT OF CRUMB RUBBER = 10317

PREWEIGHT OF CRUMB RUBBER AND TALC = 10817

**SIEVE FACTOR**

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<th>% RET</th>
<th>% PASS</th>
<th>SPEC</th>
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CORRECTED FOR TALC

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<th>SPEC</th>
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<tr>
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</table>

**FIGURE 1**
EXAMPLE

PREWEIGHT OF CRUMB RUBBER = 94.6

PREWEIGHT OF CRUMB RUBBER AND TALC = 99.6

SIEVE FACTOR = 1.057082

CORRECTED DRY WEIGHT = 94.6

CORRECTED FOR TALC = 94.3
FIGURE 3
HEATING AND DRYING MATERIALS IN MICROWAVE OVEN
(An Arizona Method)

SCOPE

1. (a) This method describes the procedure for the heating, drying, and/or determining moisture content of soils and aggregates, and bituminous mixtures utilizing a microwave oven.

(b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding number to the required degree of accuracy.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

(a) Microwave oven capable of variable heat intensity settings. When heating and drying aggregates that have been extracted from bituminous mixtures, proper ventilation of fumes from oven exhaust fan shall be provided. The microwave oven shall be operated in accordance with the manufacturer’s instructions.

(b) Sample containers suitable for microwave use.

(c) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.
HEATING AND DRYING SOILS AND AGGREGATE SAMPLES

3. The heating and drying of soils and aggregates is controlled by the intensity of heat generated that prevents aggregate breakage and sample splattering. When the percent moisture is determined, the method used with a microwave oven shall give results similar to those achieved with a conventional oven. In case of dispute, a conventional oven shall be utilized in accordance with the applicable AASHTO test method, T 255 "Total Moisture Content of Aggregate by Drying", or T 265 "Laboratory Determination of Moisture Content of Soils".

(a) The following describes the method for drying samples of soils and aggregates to constant weight.

1) Record the tare weight of a suitable size container to the nearest 0.1 gram. Place sample in container and determine and record the wet weight of sample to the nearest 0.1 gram.

2) Dry sample until a constant weight is obtained. The sample is considered to be at constant weight when further drying causes, or would cause, a difference in weight of not more than 0.1 gram.

(b) After the sample has been dried to constant weight, the percent moisture, if desired, is determined as follows:

1) Cool container and sample to room temperature. Determine and record the dry weight of sample to the nearest 0.1 gram.

2) Calculate and record the percent moisture to the nearest 0.1% by the following:

\[
\text{% moisture} = \frac{(\text{wet weight}) - (\text{dry weight})}{(\text{dry weight})} \times 100
\]

HEATING AND DRYING BITUMINOUS MIXTURES

4. Bituminous mixtures shall be heated in such a manner that controls the intensity of heat generated to prevent splattering, aggregate breakage, and asphalt being "burned off".

(a) The determination of constant weight and percent moisture, if desired, shall be determined in accordance with Arizona Test Method 406.
TENSILE PROOF DOWEL TEST

(An Arizona Method)

SCOPE

1. (a) This test method covers the determination of the tensile force necessary to break the bond between a steel reinforcing bar and portland cement concrete when utilizing an epoxy adhesive, under defined procedures for preparation, conditioning, and testing.

(b) Tensile strength is the maximum tensile load carried by the test specimen during a tension or pull-out test. The results are expressed in Newtons (pounds force).

(c) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(d) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(e) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:
(a) A rotary hammer drill with a 25.4 mm (1 inch) drill bit.

(b) Testing Machine, capable of maintaining a specified rate of loading and comprised of essentially the following:

1) Fixed Member - The upper member of the machine has grips for holding the test specimen in the machine. These grips shall secure the reinforcing bar in such a way that the concrete cylinder shall move into a seating alignment as soon as a load is applied.

2) Movable Member - This seating alignment shall bring the concrete cylinder into a uniform bearing against a steel plate that bears against the bottom of the movable or lower member.

MATERIALS

3. (a) The concrete cylinder shall be a 152.4 x 304.8 mm (6 x 12 inch) concrete cylinder fabricated according to AASHTO T 23 (ASTM C 31). The concrete utilized to fabricate the cylinder shall have a minimum compressive strength of 40 MPa (6000 psi) when tested in accordance with Arizona Test Method 314.

(b) The reinforcing bar shall be No. 20M (No. 6), conforming to the requirements of AASHTO M 31 (ASTM A 615) Grade 60, with a length of 815 mm (32 inches).

(c) The epoxy adhesive product, prepared according to the manufacturer's recommendations.

PREPARATION

4. (a) A 25.4 mm (1 inch) diameter hole is drilled in the center and parallel to the principal axis of the cylinder to a depth that will accommodate a full 150 mm (6 inches) of reinforcing bar. Compressed oil-free air shall be used to purge drill dust from the hole. Then, water under pressure along with a bottle brush will be used to flush out any remaining particles. Water under pressure will be continuously applied to the hole until the rinse water turns clear. Then the cylinder is moved to a storage area and set up side down to allow any free moisture to drain away. The cylinder will be dry before any epoxy adhesive is applied in fabricating the test specimen.
(b) If a low or medium viscosity epoxy is used, pour a sufficient quantity of epoxy into the hole, such that when a square end of reinforcing bar is pushed to the full depth of the drilled hole, the epoxy shall reach the top of the hole. A small amount of excess material extruded will be acceptable. The reinforcing bar should be rotated approximately one to three revolutions in an effort to more uniformly distribute the epoxy adhesive over the reinforcing bar and the wall of the drill hole.

(c) If a high viscosity or non-sag epoxy is used, use a spatula or putty knife in the application of the epoxy to place a sufficient quantity of the material into the drill hole. In addition, the length of reinforcing bar that is to be embedded, should be coated with epoxy before the square end of the reinforcing bar is inserted into the drill hole. A small amount of excess material extruded will be acceptable. The reinforcing bar should be rotated approximately one to three revolutions in an effort to more uniformly distribute the epoxy adhesive over the reinforcing bar and the wall of the drill hole.

(d) If a glass self-contained epoxy adhesive capsule is used, one end of the reinforcing bar shall be chisel pointed. The other end of the bar will have a square end, which shall be fitted into an attachent on a rotary hammer drill. The glass capsule shall be inserted into the drill hole and the reinforcing bar with the chisel point is then driven into the drill hole. The reinforcing bar shall be driven to the full depth of the drilled hole or refusal.

TEST SPECIMEN

5. Each test specimen shall consist of a fully assembled 152.4 mm (6 inch) diameter concrete cylinder with the rebar bonded within utilizing the epoxy adhesive product submitted for evaluation. The test shall consist of performing testing on two test specimens; one being tested at one day, and the other at three days, unless other time periods are agreed upon.

CONDITIONING

6. The conditioning of the test specimens shall be maintained, until such time as the tests are conducted, at a temperature of 24 ± 4 °C (75 ± 7 °F) and a relative humidity of 50 ± 2 percent.
PROCEDURE

7. (a) Place the test specimen in the test machine so the grips fully engage the reinforcing bar, while taking care to align the concrete cylinder and the bearing plate so that they will bear uniformly against the lower or movable member. The application of the tensile loading should begin as quickly as possible. An initial loading of 22 kN (5000 pounds force) shall be applied to the test sample. Loading shall then be applied at a rate of movement which corresponds to a loading rate within the range of 22 to 27 kN/minute (5000 to 6000 pounds force/minute) to failure.

(b) Record the maximum load carried by the test specimen at failure, to the nearest 450 N (100 pounds force).

(c) Record the type of failure that occurred. This will be based on a visual inspection. (See Note below).

NOTE: Adhesive failure refers to the lack of bond obtained by observing how much of a failure has occurred in the epoxy adhesive being in contact with the surfaces to be bonded, poor distribution of epoxy material. Concrete tensile failure could be described as being a complete failure that is preceded by developing a shallow surface cone.
REFLECTANCE, DRY OPACITY, AND YELLOWNESS INDEX OF TRAFFIC PAINT

(An Arizona Method)

SCOPE

1. (a) This method is used for integrated testing of traffic paints for their reflectance properties. It is applied primarily in testing water-borne white and yellow traffic paints. Yellowness Index is determined on white traffic paints.

(b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

(a) Black and white opacity charts - Leneta, form 2C.

(b) 0.25 mm (10 mil) doctor blade, 101.6 mm (4 inch) width.
PREPARATION OF TEST SPECIMENS (PAINT FILMS)

3. Two test films, on separate charts, are prepared for each paint sample as follows:

(a) Apply a bead of paint approximately 50.8 mm (2 inches) long and 5 milliliters in volume across the top of the upper (black) portion of a chart.

(b) Draw the paint down across both portions of the chart (black and white) using the 0.25mm (10 mil) doctor blade.

(c) Repeat, steps 3(a) and 3(b) using a bead of approximately 10 milliliters volume and the 0.38 (15 mil) doctor blade.

(d) Prior to testing, allow the test specimens to dry for a minimum of 24 hours at 25 ± 5 °C (77 ± 9 °F).

SETUP AND CALIBRATION OF REFLECTANCE METER

4. After an initial warmup time of a minimum of 30 minutes, calibrate the meter for use on the applicable filter channels (green "G", amber "A", and/or blue "B") as follows:

(a) Load the green filter in the filter rack of the "T" search unit and rotate the rack until the filter is in front of the lens.

(b) Select the green memory location by depressing the "G" button on the front face of the meter.

(c) Place the black cavity standard on the head of the search unit.

(d) Set the dark current (zero offset) by depressing first the "CHANGE" button and then the "ZERO" button on the face of the meter.
(e) Place the Photovolt ceramic reflectance standard plaque on the head of the search unit.

(f) Enter the standard reflectance value for the green filter on the meter display, by depressing first the "CHANGE" button, then the "STD" button, and finally, the "A", "B", and "G" buttons as necessary to increment the tens, units, and tenths digits respectively until the value is displayed. Then depress the "STD" button again. The standard reflectance value for the green filter is now stored in the memory, and the meter is now calibrated on the green filter channel.

(g) If reflectance data is to be obtained for a white paint sample, repeat steps 4(a) through 4(f) for the amber filter channel and again for the blue filter channel.

(h) For each channel used, immediately prior to obtaining reflectance data for paint samples, the calibration will be checked and, if necessary, corrected. First replace the black cavity standard on the head of the search unit and if drift has occurred, depress the "CHANGE" button and then the "ZERO" button. Next replace the standard plaque on the head of the search unit and if drift has occurred, depress the "CHANGE" button, and then depress the "STD" button twice.

MEASURING AND RECORDING REFLECTANCE DATA

5. (a) Obtain reflectance data for a white paint sample as follows:

1) After set up of the meter on the green filter channel, check and correct the calibration as described in paragraph 4(h). Measure the reflectance of the 0.25 mm (10 mil) specimen over the black and white portions of the chart. Record the results in the appropriate boxes of the reflectance data worksheet. (See Figure 1 for an illustration of a blank worksheet.) Then measure the reflectances of the 0.38 mm (15 mil) specimen over both portions of the chart and record similarly.

2) After set up of the meter on the amber filter channel, check and correct the calibration as described in paragraph 4(h). Measure and record reflectances of the 0.38 (15 mil) specimen over both the black and the white portion of the chart.

3) Repeat step 5(a)(2), on the blue channel.

(b) Obtain reflectance data for a yellow paint sample as in step 5(a)(1).
CALCULATING AND REPORTING REFLECTANCE PROPERTIES

6. (a) Report "Reflectance" of a white or yellow paint as the reflectance of the 0.38 mm (15 mil) specimen on the green channel over the white portion of the chart.

(b) Calculate and report "Dry Opacity" of a white or yellow paint as the ratio of reflectance of the 0.25 mm (10 mil) specimen over the black portion of the chart to the reflectance of the same over the white portion, both reflectances being measured on the green channel.

(c) Calculate and report "Yellowness Index" of a white paint according to the following formula: (Yellowness Index does not apply to a yellow paint.)

\[
\text{Yellowness Index} = \frac{\text{Amber} - \text{Blue}}{\text{Green}} \times 100
\]

Where: Amber, Blue, and Green are the measured reflectances of the 0.38 mm (15 mil) specimen over the white portion of the chart on the respective filter channels.

NOTE: Any reflectance and yellowness index properties derived from measurements made over the black portions of the charts are for informational purposes only.
REFLECTANCE DATA WORKSHEET

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<td>DRY OPACITY (BLACK/WHITE)</td>
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</table>

| 0.38 mm (15 mil) GREEN (G₂) REFLECTANCE | | | | | | |

| 0.38 mm (15 mil) AMBER (A) | | | | | | |
| 0.38 mm (15 mil) BLUE (B) | | | | | | |
| YELLOWNESS INDEX (ON WHITE PAINT) | | | | | | |
| Δ - B (100) G₂ | | | | | | |

FIGURE 1
CHLORIDE IN HARDENED CONCRETE

(An Arizona Method)

SCOPE

1. (a) This method covers the determination of chloride in concrete by a standard addition technique using a chloride electrode. Use of the standard addition technique minimizes the effects of interferences in the samples. "Total Chloride" is obtained by an acid digestion, and "Available Chloride" by water leaching of the concrete sample.

(b) This test method involves hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

(d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

REAGENTS

2. (a) Chloride standard solution, 1.000 mg/mL, (weigh 1.648 grams dried primary standard Sodium Chloride, and dilute to 1 liter with deionized water).

(b) Nitric acid, 12%. Caution must be exercised in preparing this solution. Measure 120 mL concentrated Nitric acid, slowly pour into a 1 liter (1000mL) beaker or flask with approximately 500 mL deionized water. Then, fill with deionized water to make 1 liter solution.
(c) Deionized water.

APPARATUS

3. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

(a) 300 mL Tall-Form beakers, calibrated to indicate 100 mL volume, with watch glasses.

(b) An analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.001 gram.

(c) Whatman # 43 filter paper, or equivalent.

(d) Low temperature hot plate.

(e) Magnetic stirrer and stirring bar.

(f) Orion 94-17 chloride electrode with Orion 90-01 reference electrode, or equivalent combination.

(g) Orion EA 940 specific ion meter, or equivalent.

(h) 1 mL pipette, accurate to 0.01 mL.

(i) Thermometer, accurate to at least 0.5 °C.

PROCEDURE

4. (a) Weigh a 1.000 gram representative homogeneous sample of finely pulverized concrete into a beaker. The sample should preferably pass a 75 μm (No. 200) sieve. If "ΔE", as calculated in paragraph 4(j), is less than 18 millivolts (or if "C_0", as calculated in paragraph 5(a), is equal to or greater than 0.010 mg/mL), the procedure shall be repeated using a smaller sample, recording the sample weight to the nearest 0.001 gram.

(b) To the beaker, add 25 mL nitric acid for "Total Chloride" or 25 mL deionized water for "Available Chloride", and cover with watch glass.
(c) Heat at a temperature just below the boiling point for 20 minutes, occasionally swirling solution. Do not allow solution to boil.

(d) Filter into a clean beaker and wash thoroughly using approximately 50 mL of deionized water.

(e) Dilute filtrate to 100 mL with deionized water. This is the "reading solution". Adjust solution temperature to 25 °C ± 0.5 °C and maintain at that temperature throughout remainder of test.

(f) Place stirring bar in beaker, place beaker on magnetic stirrer, insert electrodes, and initiate stirring. Stirring shall be at a constant moderate rate, such that the vortex created by stirring does not expose the tips of the immersed electrodes. The rate of stirring and the temperature (25 °C ± 0.5 °C) shall be constant throughout the remainder of the procedure, until the final reading has been obtained in paragraph (i) below.

(g) After reading has stabilized, record initial reading to the nearest millivolt as "E1".

(h) Add 1.00 mL chloride standard solution.

(i) After reading has stabilized, record final reading to the nearest millivolt as "E2".

(j) Calculate \( \Delta E = E_1 - E_2 \).

(k) Repeat steps (b) through (j) on a reagent blank in a clean beaker.

CALCULATIONS AND REPORT

5. (a) Calculate chloride concentration, "C₀" in sample reading solution and "Cₜ" in reagent blank reading solution, and record each to the nearest 0.001 mg/mL, by the following. (The formula is derived from the Nernst equation for a specific ion electrode. The relation holds in dilute solutions. See Appendix.)

\[
C₀ \text{ or } Cₜ = \frac{1}{(101) 10^{(\Delta E/S)}} - 100
\]
Where: \( \Delta E = E_1 - E_2 \)

\( S = \) Electrode slope at 25 ± 0.5 °C as determined in accordance with manufacturer's recommendations. (The slope "S" should equal approximately 59 millivolts for a properly functioning electrode. See Section 6, Appendix.)

(b) Calculate percent chloride concentration in concrete, "C" (Total Chloride or Available Chloride), and report to the nearest 0.001%, by the following:

\[
C = \frac{10 \ (C_o - C_b)}{\text{Sample Wt.}}
\]

**APPENDIX**

6. (a) The derivation of the formula for "C_o" and "C_b", the concentration of chloride in the reading solutions, is as follows:

(1) The chloride sensing electrode's behavior is given by the Nernst equation:

\[
E = E_0 - S \left[ \log \left( \frac{C_o}{C_b} \right) \right] = E_0 - \frac{2.303RT}{F} \left[ \log \left( \frac{C_o}{C_b} \right) \right]
\]

Where: \( E = \) the measured electrode potential in volts (difference of potential between chloride electrode and reference electrode).

\( E_o = \) the standard electrode potential.

\( S = \frac{2.303RT}{F} = \) the electrode slope. The theoretical value of "S" is given by:

\[
\frac{8.314}{2.303} \frac{\text{volt} \cdot \text{coul}}{\text{oK}} = \frac{298 \text{oK}}{96500 \text{coul}} = 0.059 \text{ volt}
\]

\( C = \) the chloride concentration in the reading solution in mg/mL.

\( k = \) a constant which converts concentration to activity.
For initial reading "E₁" = E₀ - S [log (kC₁)]
and final reading "E₂" = E₀ - S [log (kC₂)]

\[ \Delta E = E₁ - E₂ = S \log \left( \frac{C₂}{C₁} \right) \]

or

\[ \frac{C₂}{C₁} = 10^{\frac{\Delta E}{S}} \]

Since

\[ C₂ = \frac{C₁V₁ + CₛVₛ}{V₁ + Vₛ} \]

Where:

- \( C₂ \) = final chloride concentration in mg/mL
- \( C₁ \) = initial chloride concentration in mg/mL
- \( V₁ \) = initial volume = 100 mL
- \( Cₛ \) = standard chloride concentration = 1 mg/mL
- \( Vₛ \) = volume of chloride standard solution added = 1 mL

then

\[ \frac{C₂}{C₁} = \frac{C₁V₁ + CₛVₛ}{C₁(V₁ + Vₛ)} = 10^{\frac{\Delta E}{S}} \]

or

\[ C₁V₁ + CₛVₛ = C₁(V₁ + Vₛ)10^{\frac{\Delta E}{S}} \]

or

\[ CₛVₛ = C₁ \left[ (V₁ + Vₛ)10^{\frac{\Delta E}{S}} - V₁ \right] \]

Therefore

\[ C₁ = \frac{CₛVₛ}{(V₁ + Vₛ)10^{\frac{\Delta E}{S}} - V₁} \]

and

\[ C₀ \text{ or } C_b = \frac{1}{(101)10^{\frac{\Delta E}{S}} - 100} \]

(b) The formula for "C" (percent "Total Chloride" or "Available Chloride" concentration in concrete) is as follows:
\begin{align*}
C &= \frac{[C_o \text{ (mg/mL)} - C_b \text{ (mg/mL)}] \times 100 \text{ mL}}{\text{Sample Weight (g)}} \times \frac{1 \text{ g}}{1000 \text{ mg}} \times 100 \\
C &= \frac{10 \times (C_o - C_b)}{\text{Sample Wt.}}
\end{align*}
EXCHANGEABLE SODIUM IN TOPSOIL

(An Arizona Method)

1. SCOPE

(a) This test method is used to compare or qualify topsoil for cultivation of plants insofar as the proportion of exchangeable sodium among the four major cations (Sodium, Potassium, Calcium, and Magnesium) is related to plant welfare. "Exchangeable" is defined by the method itself and is based on the exchange of ammonium ion from the reagent Ammonium Acetate Solution with the four major cations under the conditions given by the method. The four cations are brought into solution by the reagent and their concentrations in solution are determined by Atomic Absorption or Flame Emission Spectrophotometry.

(b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

(a) Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

(1) No. 12 sieve.

(2) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.01 gram.

(3) Plastic vials, 1-1/4 inches in diameter and 4 inches in length, with caps.

(4) Reciprocating shaker capable of 180 cycles per minute, with carrier for holding vials in place.
(5) Sodium-free quantitative filter paper, 11 cm diameter.

(6) Atomic Absorption or Flame Emission Spectrophotometer capable of determining Sodium, Potassium, Calcium, and Magnesium.

3. REAGENTS

(a) Ammonium Acetate Solution, 1 Normal. (Measure 57 mL reagent glacial acetic acid into a 1 liter volumetric flask and dilute to 500 mL with deionized water. Add 69 mL reagent concentrated ammonium hydroxide. Dilute to approximately 950 mL with deionized water and mix. Adjust pH to 7.0 using glacial acetic acid or ammonium hydroxide dropwise as necessary. Dilute to the mark with deionized water.)

(b) Sodium Stock Solution, 1000 mg/liter. (Available as calibrated standard solution from chemical suppliers).

(c) Potassium Stock Solution, 1000 mg/liter. (Available as calibrated standard solution from chemical suppliers).

(d) Calcium Stock Solution, 1000 mg/liter. (Available as calibrated standard solution from chemical suppliers).

(e) Magnesium Stock Solution, 1000 mg/liter. (Available as calibrated standard solution from chemical suppliers).

(f) Hydrochloric Acid Solution, 50%. (Dilute concentrated reagent grade hydrochloric acid 1:1 with deionized water.)

(g) Lanthanum Chloride Reagent Solution, 0.26%. (Weigh 1.30 grams anhydrous reagent grade lanthanum chloride into a 500 mL volumetric flask. Dilute to the mark with deionized water.)

4. PROCEDURE

(a) Weigh 2.00 grams of soil, passing a No. 12 sieve into a plastic vial.

(b) Pipette 40.0 mL Ammonium Acetate Solution into the vial, and cap it.

(c) Place vial into carrier of reciprocating shaker. Shake at 180 cycles per minute for exactly 5 minutes.
(d) Immediately filter into a clean vial. (The folded filter paper can be placed onto the rim of the vial instead of using a funnel.) Cap the vial. This is the Original Extract Solution.

(e) Prepare a diluted solution by transferring a suitable aliquot from the Original Extract Solution to a volumetric flask, adding hydrochloric acid to 1% and lanthanum chloride to 26 mg/liter. (If a 50 mL flask is used, 1.0 mL of Hydrochloric Acid Solution and 0.5 mL of Lanthanum Chloride Reagent Solution is required.) The dilution is made with deionized water. This is the Diluted Extract Solution. The dilution should be sufficient to bring the absorbances of the four elements to values below those of the Standard Solution.

(f) Prepare a solution with concentrations of 1.00 mg/liter of sodium, 2.00 mg/liter of potassium, 5.00 mg/liter of calcium, and 0.50 mg/liter of magnesium. This solution should have 1% strength hydrochloric acid and 26 mg/liter lanthanum chloride. The dilution is made with deionized water. This is the Standard Solution.

(g) Prepare a solution with 1% Hydrochloric Acid and 26 mg/liter Lanthanum Chloride in deionized water. This is the Reagent Blank Solution.

(h) Perform an Atomic Absorption or Flame Emission Spectrophotometric analysis using the Diluted Extract Solution, the Standard Solution, and the Reagent Blank Solution. (The instrumental parameters, given in the table below are recommended for the analysis.) Determine the concentration of each of the four cations in the Original Extract Solution to the nearest 0.1 mg/liter. If the concentration of any element in the Diluted Extract Solution is higher than that in the Standard Solution, a more dilute Diluted Extract Solution should be prepared as in paragraph 4(e).

<table>
<thead>
<tr>
<th>ELEMENT</th>
<th>WAVELENGTH</th>
<th>FLAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium</td>
<td>589.2</td>
<td>Air-Acetylene</td>
</tr>
<tr>
<td>Potassium</td>
<td>766.5</td>
<td>Air-Acetylene</td>
</tr>
<tr>
<td>Calcium</td>
<td>422.7</td>
<td>Air-Acetylene</td>
</tr>
<tr>
<td>Magnesium</td>
<td>285.2</td>
<td>Air-Acetylene</td>
</tr>
</tbody>
</table>

5. **CALCULATIONS**

(a) Calculate the concentration of exchangeable cations in milliequivalents per 100 grams (meq /100 g) of soil using the following formulas:
\[ C'_{Na} \text{ (in meq/100 g)} = (0.087) \times (C_{Na}) \]
\[ C'_{K} \text{ (in meq/100 g)} = (0.051) \times (C_{K}) \]
\[ C'_{Ca} \text{ (in meq/100 g)} = (0.100) \times (C_{Ca}) \]
\[ C'_{Mg} \text{ (in meq/100 g)} = (0.165) \times (C_{Mg}) \]

Where: \( C_{Na}, C_{K}, C_{Ca}, \) or \( C_{Mg} \) is the concentration of each cation, in mg/liter, respectively in the Original Extract Solution.

(b) Compute the total Cation Exchange Capacity (CEC) of the soil, which is the sum of the four exchangeable cation concentrations, as follows:

\[ \text{CEC (in meq/100 g)} = C'_{Na} + C'_{K} + C'_{Ca} + C'_{Mg} \]

(c) Compute the Exchangeable Sodium Percentage (ESP), which is the exchangeable sodium proportion of the total Cation Exchange Capacity, as follows:

\[ \text{ESP} = \frac{C'_{Na}}{\text{CEC}} \times 100 \]

(d) Compute the Exchangeable Sodium (ES) in the soil, in parts per million, using the following formula:

\[ \text{ES (in ppm)} = (20) \times (C_{Na}) \]

6. REPORT

(a) Report Exchangeable Sodium Percentage (ESP) to the nearest 0.1 percent.

(b) Report Exchangeable Sodium (ES) to the nearest 1.0 ppm.
CALCIUM CARBONATE IN TOPSOIL
(NEUTRALIZATION POTENTIAL OF TOPSOIL)

(An Arizona Method)

1. SCOPE

1.1 This test method is used to determine the acid-consumption capacity or alkalinity of a topsoil as defined by the conditions of the test itself. The principal mineral components of the topsoil which give rise to this alkalinity are generally taken to be alkali and alkaline earth metal carbonates. They are represented by calcium carbonate, the most abundant member of the group.

1.2 A large excess of Hydrochloric Acid Standard Solution is added to ensure complete decomposition of the Calcium Carbonate. Then, the residual acid is backtitrated out with Sodium Hydroxide Standard Solution.

1.3 This test method involves hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.4 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 25 mL, volumetric pipette, accurate to ± 0.05 mL.

2.1.2 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.01 gram.
2.1.3 Filter paper equivalent to Whatman No. 40.

2.1.4 Filtration funnel, 7.5 cm, 60\(^\circ\), long stem.

2.1.5 300 mL beaker, tall form, graduated.

2.1.6 25 mL burette, accurate to \(\pm 0.01\) mL.

2.1.7 Stirring rod (glass or plastic).

3. REAGENTS

3.1 1.00 \(\pm\) 0.10 Normal Hydrochloric Acid Standard Solution. Label this solution “Reagent 1” or “R\(_1\)” on the bottle. This reagent is available from laboratory chemical suppliers as a standardized solution. Its normality should be checked regularly against “Reagent 2” below, as required by frequency of use. Record the actual normality of this solution as “N\(_1\)”.

3.2 1.00 \(\pm\) 0.10 Normal Sodium Hydroxide Standard Solution. Label this solution “Reagent 2” or “R\(_2\)” on the bottle. This reagent is available from laboratory chemical suppliers as a standardized solution. Its normality should be checked regularly against primary standard Potassium Biphthalate, as required by frequency of use. Record the actual normality of this solution as “N\(_2\)”.

3.3 Phenolphthalein Indicator (1% Phenolphthalein in 50% Isopropyl Alcohol).

3.4 Demineralized Water.

4. PROCEDURE

4.1 Weigh 1.00 \(\pm\) 0.01 gram of soil passing a No. 12 sieve into a 300 mL beaker.

4.2 Pipette 25.00 \(\pm\) 0.05 mL Hydrochloric Acid Solution (Reagent 1) into the 300 mL beaker.

4.3 Stir thoroughly with a stirring rod.

4.4 Let the mixture stand undisturbed for 15 minutes, or, if significant bubbling is observed, 30 minutes.
4.5 Filter the entire contents of the 300 mL beaker into another 300 mL beaker and rinse three times with demineralized water, being careful to keep the total volume at or under 100 mL.

4.6 Dilute to 100 mL, if necessary, with demineralized water, and add one drop of Phenolphthalein Indicator.

4.7 Backtitrate the residual Hydrochloric Acid with Sodium Hydroxide Solution (Reagent 2) to a Phenolphthalein endpoint.

4.8 Record the volume of Reagent 2 delivered, to the nearest 0.05 mL, as “$V_2$”.

5. **CALCULATIONS AND REPORT**

5.1 Calculate Calcium Carbonate, in percent, using the following formula:

$$\text{Calcium Carbonate, } \% = 5 \times \left[ 25 - \frac{N_2 \times V_2}{N_1} \right] \times N_1$$

5.2 Report Calcium Carbonate to the nearest 0.1%.
SULFATE IN SOILS
(An Arizona Method)

1. SCOPE

1.1 This test method describes a procedure for determining sulfate content in soil using a turbidimeter. The sulfate content determined is defined in terms of the method and may be called water soluble sulfate.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 The extraction procedure, Subsections 4.1 through 4.4, is the same as is used in Arizona Test Method 736, “Chloride In Soils”.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 Turbidimeter - with at least ± 2% accuracy to 1000 NTU (nephelometric turbidity units) and at least ± 5% accuracy to 4000 NTU, with sample cuvettes.

2.1.2 Balances or scales:

2.1.2.1 One balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity shall be at least 0.1 gram.
2.1.2.2 An analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M231, except the readability and sensitivity shall be at least 0.1 milligram.

2.1.3 Beaker(s) - three, 200 mL capacity each.

2.1.4 Erlenmeyer flask - 500 mL capacity with stopper.

2.1.5 Volumetric flask - 1000 mL capacity, accurate to 0.3 mL.

2.1.6 Reagent dispensing bottle - 500 mL capacity, capable of repeatedly dispensing 5 mL of reagent.

2.1.7 Reagent storage bottles - three, one-liter capacity, each with cap.

2.1.8 Dropping bottle - 60 mL, with dispensing tip or dropper.

2.1.9 Centrifuge tube - 50 mL, with cap.

3. REAGENTS

3.1 Conditioning Reagent.

3.1.1 Into a reagent storage bottle, add 300 mL demineralized water, 30 mL Reagent Grade Concentrated Hydrochloric Acid, 100 mL Reagent Grade Isopropyl Alcohol, 75 grams Reagent Grade Sodium Chloride, and 50 mL Reagent Grade Glycerol. Mix well. Transfer to a reagent dispensing bottle as needed.

3.2 Sulfate Standard Solution, 0.100 mg/mL (100 ppm).

3.2.1 Into a 1000 mL volumetric flask, add 147.9 mg Reagent Grade Anhydrous Sodium Sulfate. Fill the flask to the 1000 mL mark with demineralized water. Mix well. Transfer to a reagent storage bottle.

3.3 Barium Chloride Reagent Solution, 1%.

3.3.1 Into a 1000 mL volumetric flask, add 10 grams of Reagent Grade Barium Chloride Powder, Anhydrous. Fill the flask to the 1000 mL mark with demineralized water. Mix well. Transfer to a reagent dispensing bottle as needed, and the remainder to a reagent storage bottle.
3.4 Nitric Acid, 20%.

3.4.1 Measure 10 mL concentrated nitric acid into a suitable size beaker containing approximately 40 mL demineralized water and mix well. Transfer to a dropping bottle.

**CAUTION:** Exercise extreme caution in preparing and using the Nitric Acid solution. It must be properly labeled and treated as a hazardous material.

3.5 Demineralized water.

4. **PROCEDURE**

4.1 Weigh 100.0 ± 0.1 grams of soil passing a No. 10 sieve into a 500 mL Erlenmeyer flask.

4.2 Weigh 300.0 ± 0.1 grams demineralized water into the flask.

4.3 Stopper the flask, shake vigorously, and let the mixture stand undisturbed for one hour.

4.4 Carefully, with minimal disturbance of the sediment, decant 50 mL of the extract solution into a centrifuge tube.

4.4.1 If the decanted extract solution is clear, proceed to Subsection 4.5.

4.4.2 If the decanted extract solution is not clear, place the tube into the centrifuge and centrifuge at a minimum of 5000 RPM for 10 minutes. If the solution is still not clear, add 2 drops of 20% Nitric Acid solution to the tube and centrifuge again. Repeat until the centrifugate is clear.

4.5 Transfer 20.0 mL of the clear extract solution (or centrifugate) into a 200 mL beaker. This is the sample aliquot, “ALSAM.”

4.5.1 Dispense 5 mL of Conditioning Reagent and 5 mL of Barium Chloride Reagent Solution into the beaker, and dilute to 100 mL. This is the “Sample Solution.”

4.6 Transfer 20.0 mL of Sulfate Standard Solution into a second 200 mL beaker. This is the standard aliquot “ALSTD.”
4.6.1 Dispense 5 mL of Conditioning Reagent and 5 mL of Barium Chloride Reagent Solution into this beaker, and dilute to 100 mL. This is the “Standard Solution.”

4.7 Dispense 5 mL of Conditioning Reagent and 5 mL of Barium Chloride Reagent Solution into a third 200 mL beaker, and dilute to 100 mL. This is the “Reagent Blank Solution.”

4.8 Stir the three solutions and let them stand undisturbed for at least 15 minutes.

4.9 Rinse a sample cuvette with demineralized water and then with freshly stirred Reagent Blank Solution. Immediately transfer enough Reagent Blank Solution to fill the cuvette to the mark.

4.9.1 Place the cuvette into the turbidimeter and read the turbidity to the nearest 0.1 NTU. The turbidity of the Reagent Blank Solution is recorded as “TBBNK”.

4.10 Rinse a sample cuvette with demineralized water and then with freshly stirred Sample Solution. Immediately transfer enough Sample Solution to fill the cuvette to the mark.

4.10.1 Place the cuvette into the turbidimeter and read the turbidity to the nearest 0.1 NTU. Record the turbidity of the Sample Solution as “TBSAM.”

4.11 Rinse a sample cuvette with demineralized water and then with freshly stirred Standard Solution. Immediately transfer enough Standard Solution to fill the cuvette to the mark.

4.11.1 Place the cuvette into the turbidimeter and read the turbidity to the nearest 0.1 NTU. Record the turbidity of the Standard Solution as “TBSTD.”

4.12 Compare the values of “TBSAM” and “TBSTD.” If “TBSAM” is larger than “TBSTD”, repeat Subsections 4.5 and 4.5.1 using a suitably smaller sample aliquot. Record this volume, to the nearest 0.1 mL, as “ALSAM.” Repeat Subsections 4.10 and 4.10.1.

5. **CALCULATION AND REPORT**

5.1 Calculate sulfate content in the soil in parts per million, “S”, and report to the nearest 10 ppm as follows:
\[ S = 300 \times \frac{\text{ALSTD} \times \text{TBSAM}}{\text{ALSAM} \times \text{TBSTD}} \]

Where:

- ALSTD = 20.0 mL (Size of the standard aliquot).
- TBSTD = Turbidity corresponding to ALSTD.
- ALSAM = Size of the sample aliquot, mL.
- TBSAM = Turbidity corresponding to ALSAM.

**Note:** If the turbidity of the Reagent Blank Solution “TBBNK” is 0.40 or higher, for greater accuracy, the value of “TBBNK” shall be subtracted from “TBSAM” and from “TBSTD” before calculating S above.
DETERMINATION OF PORTLAND CEMENT CONTENT IN CEMENT TREATED BASE MATERIAL

(An Arizona Method)

SCOPE

1. (a) This test procedure covers a method for the determination of the percentage of portland cement in cement treated base material.

   (b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

   (a) Burette - 50 mL capacity with teflon stopcock and burette cover.

   (b) Burette clamps.
(c) Ring stand.

(d) Pipette - 10 mL capacity, accurate to 0.05 mL.

(e) Plastic dropper dispenser - 100 mL.

(f) Plastic wash bottles - 100 mL capacity, 500 mL capacity, 1000 mL capacity.

(g) Graduated cylinders - (transparent plastic), 100 mL capacity, 500 mL capacity, 1000 mL capacity.

(h) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

(i) An analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.0001 gram.

(j) Beakers - (plastic), 300 mL capacity, 2000 mL capacity, 4000 mL capacity.

(k) Stirring rods - stainless steel, 305 mm (12") long; glass, not over 150 mm (6 inches) long.

(l) Erlenmeyer flasks - (plastic), 500 mL capacity with stoppers.

(m) Variable control hot plate.

(n) Spatulas, cleaning brushes, watch glasses.

(o) Stop watch or equally efficient timing device.

(p) Plastic bottles - 3.8 liter (1 gal.), 1 liter (1 qt.).

(q) 19.0 liter (5 gallon) polyethylene jug.
REAGENTS

3. Reagents shall be prepared as described below. Distilled or de-ionized water shall be used in their preparation.

(a) 2-Propanol (Analyzed Reagent)

(b) Tartaric Acid 72% Solution (TAS).

1) For a 19.0-liter (5 gallon) batch, weigh 13680 grams of reagent grade D-Tartaric Acid into a 19.0 liter (5 gallon) polyethylene jug, or other suitable non-reactive container, precalibrated to indicate a 19.0 liter (5 gallon) volume.

2) Add boiling water to fill the jug to the 19.0 liter (5 gallon) mark. Mix thoroughly and let set overnight.

3) Filter into another 19.0 liter (5 gallon) jug and dilute to the 19.0 liter (5 gallon) mark.

(c) Potassium Hydroxide 1-Normal Titrating Solution (TS).

1) Weigh 56.1 grams reagent grade Potassium Hydroxide into a one liter polyethylene jug, or other suitable non-reactive container, precalibrated to indicate a one liter volume. The Potassium Hydroxide shall not be placed in a glass container.

2) Add sufficient water to dissolve the Potassium Hydroxide. Mix thoroughly and allow to come to room temperature. Dilute to the one liter mark. This solution shall not be stored in a glass container.

(e) 1% Methyl Red Indicator.

1) For a 50 mL batch, weigh 0.5 gram methyl red dye into a 100 mL beaker.

2) Add 25 mL 2-propanol and dissolve.

3) Fill to 50 mL mark with distilled water.

4) Transfer to the dropper dispenser.
PREPARATION OF STANDARD SAMPLES

4. (a) Standard samples shall be prepared using the same materials and mix design formula as the job from which field samples will be taken. A representative minimum 3000 grams of soil/aggregate material with no added cement shall be used to prepare the standard samples. From this, three representative portions to be used for preparing standard samples shall be obtained using a standard reduction method such as given in AASHTO R 76. The standard samples shall be prepared with differing cement contents bracketing the mix design cement content as follows:

   (1) 0.0% cement content.

   (2) The mix design cement content, expressed to the nearest 0.1%.

   (3) 5.0% more cement than the design content.

(b) The three components of the standard samples are cement, soil/aggregate material, and water. The total sample weight shall be 300 grams. The weight of water from the job is 30 grams. The dry components of the standard sample are cement and soil/aggregate material with a combined weight of 270 grams. The state of dryness of the soil/aggregate material used in preparing standard samples may be either saturated-surface-dry, air-dry, or oven-dry. It is important that the same state of dryness be obtained for all standard samples and for field test samples, [See paragraph 8(a)]. Use the following formulas to determine the dry components of each standard sample:

   \[ W_c = 300 \times \frac{C}{100} \]

   \[ W_{s/a} = 270 - W_c \]

Where:

- \( W_c \) = Weight of cement, grams
- \( W_{s/a} \) = Weight of soil/aggregate, grams
- \( C \) = Cement content, % of total sample

NOTE: In case a large quantity of large aggregate is present in a cement treated base job, a minimum of 15,000 grams of soil/aggregate material should be obtained for preparation of standard samples, and the following procedure followed:
(1) For each standard sample, 5000 grams of soil/aggregate material is obtained using the standard reduction method and separated into coarse and fine fractions.

(2) The coarse fraction is washed with distilled water and its adhering fines are added to the fine fraction.

(3) From the resulting combined fine fraction, a representative portion is obtained using the standard reduction method.

TEST PROCEDURE FOR STANDARD SAMPLES

5. (a) Transfer the dry components to the 2000 mL plastic beaker.

NOTE: It is preferable to weigh the cement as accurately as possible. Use an analytical balance to weigh the desired amount of cement. Then transfer the cement to the aggregate already in the plastic beaker.

(b) Mix thoroughly. AVOID PRODUCING DUST.

(c) Add 30 grams of water that is used on the job for mixing the batch.

NOTE: At this point it may be desirable to thoroughly mix the standard sample and transfer it to a suitable container and allow it to cure for a sufficient time to approximate the expected condition of the field sample at the time it is tested, after which the standard sample must be broken down as necessary to facilitate mixing and intimate contact with the reagent (TAS). An apparatus such as the one described in AASHTO T 87 may be utilized for this purpose. Care should be taken to avoid reducing the size of individual natural particles. Quantitatively transfer the standard sample to the 2000 mL beaker.

(d) Add 400 ± 0.1 mL of distilled water to the beaker.

(e) Stir thoroughly and vigorously for 3 ± 1 minutes.
Cover the 2000 mL beaker with a watch glass. Keep the beaker covered with a watch glass at all times, except when actually in use.

Let stand undisturbed for 40 ± 5 minutes.

Stir thoroughly and vigorously for 2 minutes. Let stand undisturbed for 1 to 2 minutes.

Accurately measure 300 mL Tartaric Acid 72% Solution (TAS) into a 500 mL graduated cylinder.

NOTE: Keep all containers of all reagents covered when not in use. Keep all stock containers sealed.

Add TAS slowly and with continued stirring to the beaker.

NOTE: If effervescence occurs, stop addition of TAS and stir vigorously until bubbling ceases, complete the addition of TAS, then stir at least 2 minutes continuously.

Let the sample stand undisturbed for 10 minutes.

Accurately measure 710 mL of 2-propanol into a 1000 mL graduated cylinder.

Add the 2-propanol, all at once, to the beaker, completely draining the cylinder. Immediately stir, rapidly and thoroughly, for exactly 2 minutes, so that all the contents of the beaker are washed with 2-propanol. Cover with a watch glass, and let the beaker stand undisturbed until a clear liquid layer forms.

Carefully pour only the clear liquid into a 500 mL plastic Erlenmeyer flask. Tightly stopper and seal the flask.

TITRATION PROCEDURE

6. (a) Fill the 50 mL burette completely with Potassium Hydroxide 1-Normal Titrating Solution (TS). Open the stopcock and let the burette drain completely.

(b) Refill the burette with TS, and adjust the bottom of the meniscus to read 0.0 mL.
NOTE: Make sure that there are no air bubbles anywhere in the system. Always check just below the stopcock for air bubbles when the burette is draining. Keep the top of the burette covered.

(c) Transfer an aliquot of 10.00 ± 0.05 mL of the clear liquid from the flask to a 300 mL plastic beaker using a 10 mL pipette.

(d) Add 100 mL distilled water to the beaker, using a 100 mL graduated cylinder.

(e) Stir the contents of the beaker with a glass stirring rod. Do not remove the stirring rod until the titration is completed.

(f) Add 2 drops of 1% Methyl Red Indicator and stir the solution. The test solution should now be a clear, dark-red color. The color change is as follows:

   Start: Clear, dark-red solution.
   To: Clear, pure orange solution.
   End: Clear, pure lemon-yellow solution.

(g) Rapidly, with stirring, add TS until the solution turns to an orange color.

(h) Stir the solution.

(i) Set the burette stopcock to deliver the TS one drop at a time, with stirring between each addition. As the end point is approached, the solution should turn to a clear red-orange or clear yellow-orange color.

(j) Decrease the rate of addition of TS, and stop addition when one drop produces a pure lemon-yellow clear solution. Allow the solution to stand undisturbed for one minute to check for color fading.

(k) If the color fades, add one more drop and again allow to stand undisturbed for one minute. Repeat until the color does not fade.

(l) Read the volume of TS used to the nearest 0.1 mL at the bottom of the meniscus.
(m) If time is available, run at least two titrations for each sample and obtain the average volume of TS used.

PREPARATION OF THE STANDARD CURVE

7. (a) The standard curve is prepared using linear graph paper.
(b) Plot the number of mL of TS, along the X-axis.
(c) Plot the percentages of portland cement, along the Y-axis.
(d) Draw the straight line of best fit for the three points.

TEST PROCEDURE FOR FIELD SAMPLE

8. (a) From a representative minimum 1000 gram field sample of the cement treated base material to be tested, at the same state of dryness as the standard samples, obtain a 270 ± 0.1 gram test sample using the standard reduction method used with the standard samples and transfer it to a 2000 mL plastic beaker.

NOTE: As necessary, the field sample must be broken down to facilitate weighing, mixing, and intimate contact with the reagent (TAS). An apparatus such as the one described in AASHTO T 87 may be utilized for this purpose. Care should be taken to avoid reducing the size of individual natural particles.

NOTE: If much large aggregate is present, a representative minimum of 5000 gram field sample should be used. It should be separated into coarse and fine fractions and the coarse fraction washed with distilled water to obtain a combined fine fraction from which the test sample is prepared as above. [See note following paragraph 4(b).]

(b) Add 400 ± 0.1 mL of distilled water to the beaker.
(c) Stir thoroughly and vigorously for 3 ± 1 minutes.
(d) Cover the beaker with a watch glass. Keep the beaker covered with a watch glass at all times, except when actually in use.
(e) Let the beaker stand undisturbed for 40 ± 5 minutes.

(f) Follow the procedure as described in paragraphs 5(h) through 5(n).

TITRATION PROCEDURE

9. Titrate in accordance with Section 6.

REPORT

10. Read results directly from the Standard Curve as percent portland cement in the field sample. Report the result to the nearest 0.1%.
TESTING OF THERMOPLASTIC PAVEMENT MARKING MATERIAL

(An Arizona Method)

SCOPE

1. (a) Thermoplastics to be tested according to the procedures given herein are mixtures of resins, mineral fillers, pigments, organic additives, and reflective glass spheres.

   (b) All testing is done on hot melt test specimens made from the commercial products. These include field test specimens obtained from highway striping operations and specimens produced according to Part I of this method.

   (c) This method is divided into two parts: Part I gives a procedure for producing hot melt test specimens. Part I also gives an alternate procedure for producing hot melt test specimens in accordance with AASHTO T 250. Part II gives the procedures for testing specimens.

   (d) This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

   (e) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (f) Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

PART I - PROCEDURE FOR PRODUCING HOT MELTS

APPARATUS

2. (a) Heating mantle - Glas-Col Model 620 or equivalent, capable of accepting a 4000 mL beaker.
(b) Stainless steel beaker - 4000 mL, Vollrath No. 84000 or equivalent, for use with heating mantle.

(c) Mixer - Variable-speed, air-driven, Cole-Parmer No. E-04685-00 or equivalent.

(d) Mixer stand - Cole-Parmer No. E-04546-00 or equivalent with 60 cm height and capability of holding mixer.

(e) Adjustable chuck - Cole-Parmer No. E-04423-00 or equivalent, capable of coupling mixer shaft to impeller assembly shaft.

(f) Impeller assembly - Any assembly with a 35-40 cm shaft of 0.5 cm minimum diameter, dual impellers of 8-10 cm diameter spaced 4-6 cm apart with one at the end of the shaft, capable of efficient mixing of 3000-3500 mL of fluid at 700-800 rpm. Alternatively, a heavy-duty “Jiffy” type impeller for one gallon paint containers is suitable.

(g) Temperature controller with thermocouple probe - Glas-Col Digitroll II, No. 104A PL612, with 6 in. type “J” iron-constantan thermocouple or equivalent, capable of controlling mixture temperature in conjunction with heating mantle at 218 °C.

(h) Ladles - Stainless steel, 30 mL and 60 mL capacity with long handles for sampling hot melts.

(i) Aluminum or stainless steel lid to cover beaker, with holes for impeller shaft and thermocouple, and turned-down edges for holding in place over beaker.

PROCEDURE

3. (a) Mount the mixer, with impeller assembly and lid, onto the mixer stand and note position of mixer which locates bottom of impeller assembly about 2-3 cm from bottom of beaker. Secure the mantle to the mixer stand in such a position as to permit the impeller assembly to be placed over the center of the beaker and to prevent relative movement between the impeller assembly and the beaker. Plug the mantle cord into the controller. Plug the controller into a wall outlet.

(b) Remove the mixer with impeller assembly and lid. Split out a 6000 gram sample of thermoplastic material and add enough to the beaker to come to within 5-6 cm of the top. Replace the mixer, impeller, and lid. (It may be necessary to first work the impeller assembly into the thermoplastic, and then replace the lid and mixer). Place the thermocouple in position through its hole in the lid, making sure it penetrates
into the thermoplastic. Set the controller parameters for automatic control of temperature at 218 °C, and start the heating cycle.

(c) When the thermoplastic has melted sufficiently, start the mixer at a low speed. As the temperature continues to rise, increase the mixer speed until it is evident that the entire mass of thermoplastic in the beaker is being turned over continuously. (Note this mixer speed for future use). Continue heating and mixing while periodically checking the level of thermoplastic in the beaker. If the level drops below 5-6 cm from the top of the beaker, add more thermoplastic as necessary. Continue this process for four hours after the material has reached the final temperature of 218 °C.

(d) At the end of the four hour period, turn the mixer off and remove it with the lid and impeller assembly. Immediately use the ladles to transfer enough of the melt to fill a set of suitable can lids, usually a one-gallon can lid and two one-quart can lids. Set the lids with the test specimens aside for cooling. These will be used for all of the tests of the thermoplastic material.

(e) Use heat resistant gloves to remove the beaker from the mantle and discard the remainder of the thermoplastic. Allow the beaker to drain completely while the material is hot.

Note: Alternatively, the procedure for producing hot melt test specimens for Reflectance, Color, and Yellowness Index as given in AASHTO T 250 may be used, if desired. If AASHTO T 250 is used, the amount of material specified may be increased to approximately 500 grams.

| PART II - TESTS |

**REFLECTANCE AND YELLOWNESS INDEX**

4. The apparatus and procedure for determining reflectance and yellowness is given in Sections 5 through 8.

**APPARATUS**

5. Reflectance and gloss meter - Photovolt 577 or equivalent, with "T" search head.
CALIBRATION OF METER

6. After an initial warmup time of a minimum of 30 minutes, calibrate the meter for use on the applicable filter channels (green "G", amber "A", and/or blue "B") as follows:

   (a) Load the green filter in the filter rack of the "T" search unit and rotate the rack until the filter is in front of the lens.

   (b) Select the green memory location by depressing the "G" button on the front face of the meter.

   (c) Place the black cavity standard on the head of the search unit.

   (d) Set the dark current (zero offset) by depressing first the "CHANGE" button and then the "ZERO" button on the face of the meter.

   (e) Place the Photovolt ceramic reflectance standard plaque on the head of the search unit.

   (f) Enter the standard reflectance value for the green filter on the meter display, by depressing first the "CHANGE" button, then the "STD" button, and finally, the "A", "B", and "G" buttons as necessary to increment the tens, units, and tenths digits respectively until the value is displayed. Then depress the "STD" button again. The standard reflectance value for the green filter is now stored in the memory, and the meter is now calibrated on the green filter channel.

   (g) If reflectance data is to be obtained for a white test specimen, repeat steps 6(a) through 6(f) for the amber filter channel and again for the blue filter channel.

   (h) For each channel used, immediately prior to obtaining reflectance data the calibration should be checked and, if necessary, corrected. First replace the black cavity standard on the head of the search unit and if drift has occurred, depress the "CHANGE" button and then the "ZERO" button. Next replace the standard plaque on the head of the search unit and if drift has occurred, depress the "CHANGE" button, and then depress the "STD" button twice.

MEASURING AND RECORDING REFLECTANCE DATA

7. (a) Obtain reflectance data for a white test specimen as follows:

       (1) After calibration of the meter on the green filter channel, measure the reflectance of the specimen. Record the result as "Green."
(2) Repeat step 7(a)(1) for the amber channel. Record the result as "Amber."

(3) Repeat step 7(a)(1) for the blue channel. Record the result as "Blue."

(b) Obtain reflectance data for a yellow test specimen as in step 7(a)(1) for the green channel only.

CALCULATING AND REPORTING REFLECTANCE PROPERTIES

8. (a) Report "Reflectance" of a white or yellow test specimen as the reflectance value "Green" obtained as in 7(a)(1) or 7(b) above, respectively.

(b) Calculate and report "Yellowness Index" of a white test specimen according to the following formula:

\[
\text{Yellowness Index} = \frac{\text{Amber} - \text{Blue}}{\text{Green}} \times 100
\]

BINDER AND GLASS BEAD CONTENT

9. The apparatus, reagents, and procedure for determining binder content and glass bead content is given in Sections 10 through 13.

APPARATUS

10. (a) A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.01 gram.

(b) Porcelain crucibles, 30 mL capacity.

(c) Muffle furnace, capable of holding a constant temperature of 538 °C and having a sufficiently large chamber for several crucibles.

(d) Drying oven, capable of holding a constant temperature of 100 °C.

(e) Mortar, 75-100 mL capacity, with pestle.

(f) Dessicator.

(g) Beakers, 400 mL capacity.
(h) Aluminum dishes, 57 mm diameter.

(i) Small spatula.

(j) Balance brush.

(k) Hot plate.

**REAGENT**

11. Hydrochloric acid solution, 50%, prepared by adding (cautiously) 500 mL concentrated hydrochloric acid to 500 mL demineralized water.

**PROCEDURE**

12. (a) Break up a test specimen into pieces sufficiently small to permit weighing of material into a crucible.

    (b) Weigh a crucible to the nearest 0.01 gram. Record the weight as “C”.

    (c) Zero the balance and weigh 10.00 ± 0.10 gram of test specimen into the crucible. Record the weight as “S”.

    (d) Preheat the furnace to 538 ºC.

    (e) Place the crucible (cautiously) into the furnace.

    (f) After one hour, remove the crucible from the furnace and place it into the dessicator for cooling.

    (g) After cooling, remove the crucible and weigh it, with its contents, to the nearest 0.01 gram. Record the weight as “R”.

    (h) Use the spatula to scrape the inside of the crucible and quantitatively transfer all of the residue into the mortar.

    (i) Using the pestle, gently grind the residue to a fine powder, taking care not to crush the glass beads.

    (j) Using the brush, quantitatively transfer all of the powdered residue to a beaker.
(k) Slowly add 150 mL of acid solution to the beaker, taking care to minimize possible splattering due to generation of carbon dioxide.

(l) Stir the mixture by swirling, place the beaker on the hot plate, and heat to boiling with frequently stirring.

(m) Remove the beaker from the hot plate and add 150 mL of demineralized water.

(n) Let stand until the beads settle and decant the suspension, being careful not to lose any beads.

(o) Add about 100 ml of water and repeat step 12(n) until the decanted water is clear.

(p) Place the beaker into the drying oven.

(q) When completely dried, remove the beaker, with beads, from the oven.

(r) Allow the beaker and beads to cool. Weigh to the nearest 0.01 gram and record the weight as "B1".

(s) Using the brush, quantitatively transfer the beads to a suitable receptacle for microscopic examination, if desired, or to a waste container.

(t) Reweigh the empty beaker to the nearest 0.01 gram. Record the weight as "B2".

CALCULATIONS AND REPORT

13. (a) Calculate and report the binder content, to the nearest 0.1%, using the following formula:

\[
\text{Binder Content, \%} = \frac{C + S - R}{S} \times 100
\]

(b) Calculate and report the bead content, to the nearest 0.1%, using the following formula:

\[
\text{Bead Content, \%} = \frac{B_1 - B_2}{S} \times 100
\]
CHLORIDE IN SOILS

(An Arizona Method)

1. SCOPE

1.1 This test method describes a procedure for determining chloride content in soil by a standard addition technique using a chloride electrode. The chloride content is defined in terms of the method and may be called water soluble chloride.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 The extraction procedure, Subsections 4.1 through 4.4, is the same as is used in Arizona Test Method 733, “Sulfate In Soils”.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus shall consist of the following:

2.1.1 Tall-form beakers - two, 200 mL each (Pyrex #1060 or equivalent), calibrated to indicate 100 mL volume.

2.1.2 An analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M231, except the readability and sensitivity of any balance utilized shall be at least 0.001 gram.

2.1.3 Magnetic stirrer and stirring bar.
2.1.4 Erlenmeyer flask(s) - 500 mL capacity with stopper(s).

2.1.5 Chloride electrode system - (Orion Ionplus #9617 combination chloride electrode or equivalent), suitable for use in a 200 mL tall-form beaker.

2.1.6 Specific ion meter - (Orion EA 940 or equivalent), suitable for use with the chloride electrode system.

2.1.7 Thermometer - accurate to at least 0.5 °C.

2.1.8 Pipettes - 1 mL, 5 mL, and 10 mL capacity, accurate to 1%.

2.1.9 Sample bottle - 200 mL capacity or larger, with cap.

2.1.10 Dropping bottle - 60 mL, with dispensing tip or dropper.

2.1.11 Centrifuge tube - 50 mL, with cap.

3. **REAGENTS**

3.1 Chloride Standard Solution, 1.000 mg/mL (1000 ppm).

3.1.1 Fisher Scientific # LC13000-1 or equivalent. Alternatively, the Chloride Standard Solution may be made by transferring 1.648 gram dried primary standard sodium chloride into a 1000 mL volumetric flask and diluting to 1000 mL with demineralized water.

3.2 Nitric Acid, 20%.

3.2.1 Measure 10 mL concentrated nitric acid into a suitable size beaker containing approximately 40 mL demineralized water and mix well. Transfer to a dropping bottle.

**CAUTION:** Exercise extreme caution in preparing and using the Nitric Acid solution. It must be properly labeled and treated as a hazardous material.

3.3 Demineralized Water.
4. **PROCEDURE**

4.1 Weigh 100.0 ± 0.1 grams of soil passing a No. 10 sieve into a 500 mL Erlenmeyer flask.

4.2 Weigh 300.0 ± 0.1 grams demineralized water into the flask.

4.3 Stopper the flask, shake vigorously, and let the mixture stand undisturbed for one hour.

4.4 Carefully, with minimal disturbance of the sediment, decant 50 mL of the extract solution into a 50 mL centrifuge tube.

4.4.1 If the decanted extract solution is not cloudy, proceed to Subsection 4.5.

**Note:** The decanted extract solution does not have to be clear, as it is required to be in Arizona Test Method 733.

4.4.2 If the decanted extract solution is cloudy, place the tube into the centrifuge and centrifuge at a minimum of 5000 RPM for 10 minutes.

4.5 Pipette a 10.0 mL aliquot of the extract solution (or centrifugate) into a 200 mL beaker. Record this volume as “A”. Dilute to the 100 mL mark with demineralized water and add three drops of Nitric Acid, 20%. This is the “Sample Reading Solution.”

4.6 Place a stirring bar into the beaker, place the beaker onto the magnetic stirrer, insert the electrodes, and initiate stirring. Stirring shall be at a constant moderate rate, such that the vortex created by stirring does not expose the tips of the immersed electrodes. The rate of stirring shall be constant throughout the procedure.

4.7 After the reading has stabilized, record the initial reading to the nearest millivolt, as “E₁”.

4.8 Add 1.00 mL of Chloride Standard Solution.

4.9 After the reading has stabilized, record the final reading to the nearest millivolt, as “E₂”.

4.10 Calculate $\Delta E$. ($\Delta E = E_1 - E_2$).
Note: "ΔE" is calculated and recorded initially as "ΔE₀" for the Sample Reading Solution. It is subsequently also calculated and recorded as "ΔEₐ" for the Reagent Blank Reading Solution.

4.11 If "ΔE₀" (for the Sample Reading Solution) is less than 18, repeat Subsections 4.5 through 4.10 with a suitably smaller size aliquot, recording this aliquot volume as “A.”

4.12 Prepare a Reagent Blank Reading Solution by placing 100 mL of demineralized water into a second 200 mL beaker and adding 3 drops of Nitric Acid 20%. Repeat Subsections 4.6 through 4.10, calculating and recording "ΔEₐ" (for the Reagent Blank Reading Solution).

5. CALCULATIONS AND REPORT

5.1 Calculate chloride concentration, "C₀," in the Sample Reading Solution and "Cₐ" in the Reagent Blank Reading Solution, and record each to the nearest 0.001 mg/mL, as follows:

\[
C₀ = \frac{1}{101 \times 10^{\frac{[\Delta E₀/S]}{100}}}
\]

\[
Cₐ = \frac{1}{101 \times 10^{\frac{[\Delta Eₐ/S]}{100}}}
\]

Where:

S = Electrode slope, as determined in accordance with the manufacturer’s recommendations. (The slope should equal approximately 59 millivolts for a properly functioning electrode.)

5.2 Calculate chloride concentration in the soil in parts per million, "C", and report to the nearest 10 ppm as follows:

\[
C = \frac{300,000 \times (C₀ - Cₐ)}{A}
\]
CHLORIDE IN CONCRETE ADMIXTURES

(An Arizona Method)

SCOPE

1. (a) This test method describes a procedure for determining the chloride content of products used as admixtures in Portland cement concrete. The method employs a standard addition technique using a chloride electrode.

   (b) This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

APPARATUS

2. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus shall consist of the following:

   (a) 300 mL tall-form beaker (Pyrex #1060 or equivalent), calibrated to indicate 100 mL volume.

   (b) An analytical balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M231, except the readability and sensitivity of any balance utilized shall be at least 0.001 gram.

   (c) Suitable magnetic stirrer and stirring bar.

   (d) Chloride electrode system (Orion Ionplus #9617 combination chloride electrode or equivalent) suitable for use in the 300 mL beaker.

   (e) Specific ion meter (Orion EA 940 or equivalent) suitable for use with the chloride electrode system.
(f) 3 mL disposable syringe with cap (Beckton Dickinson # 301112 or equivalent).

(g) 1.000 mL pipette or pipetter accurate to ± 0.001 mL.

REAGENTS

3. (a) Chloride standard solution, 1.000 mg/mL (1000 PPM), (Fisher Scientific #LC13000-1, or equivalent, or made by transferring 1.648 gram dried primary standard sodium chloride into a 1 liter volumetric flask and diluting to 1000 mL with demineralized water).

(b) Nitric acid, 8% (made by transferring 80 mL of concentrated nitric acid into a 1 liter volumetric flask containing approximately 500 mL demineralized water and diluting to 1000 mL with demineralized water. Caution must be exercised in preparing and using this solution! It should be properly labelled and treated as a hazardous material.

PROCEDURE

4. (a) Weigh 1.000 ± 0.010 gram sample of admixture into a beaker. (If liquid, weigh by difference using a syringe). Record the sample weight to the nearest 0.001 gram as "A."

(b) Transfer 25 mL nitric acid, 8% into the beaker and swirl to mix with the sample.

(c) Dilute to 100 mL with demineralized water.

(d) Place a stirring bar into the beaker, place the beaker onto the magnetic stirrer, insert the electrode(s) into the solution, and initiate stirring. Stir at a constant moderate rate such that the vortex created by stirring does not expose the tip(s) of the electrode(s) to the air. The rate of stirring and the temperature (25.0 ± 1.0 °C) shall be held constant throughout the procedure.

(e) After the reading has stabilized, record it to the nearest millivolt, as "E_1."

(f) Pipette 1.000 ± 0.001 mL chloride standard solution into the beaker.
(g) After the reading has stabilized, record it to the nearest millivolt, as "E₂."

(h) Calculate \( \Delta E = E_1 - E_2 \).

(i) If \( \Delta E \) is less than 18 mV, repeat steps (a) through (h) with a smaller size sample.

(j) Prepare a reagent blank solution by transferring 25 mL of Nitric Acid, 8% into a 300 mL beaker and diluting to 100 mL with demineralized water. Repeat steps (d) through (h).

CALCULATIONS AND REPORT

5. (a) Calculate chloride concentration, "C₀" in the sample reading solution and "Cₐ" in the reagent blank reading solution, and record each to the nearest 0.010 mg/mL, as follows:

\[
C₀ \text{ or } C_b = \frac{1}{(101)(10 \frac{\Delta E}{S}) - 100}
\]

Where: \( S \) = Electrode slope at 25 ± 1.0 °C as determined in accordance with manufacturer's recommendations. (The slope should equal approximately 59 millivolts for a properly functioning electrode.)

(b) Calculate chloride concentration in the admixture in parts per million, "C", and report to the nearest 10 PPM as follows:

\[
C = \frac{10^5(C₀ - C_b)}{A}
\]
MEAN MACROTEXTURE DEPTH OF MILLED PAVEMENT

(A Modification of Indiana Test Method 812-03T and ASTM E 965)

1. SCOPE

(a) This test method describes the procedure to determine the Mean Macrotecture Depth of a milled pavement surface.

(b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

(c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

(a) The apparatus shall consist of the following:

(1) Filler - Type 1 glass beads conforming to the requirements of AASHTO M 247.

(2) Spreader - A flat, stiff, hard disk made from methyl methacrylate (Plexiglas) with a thickness of 0.5 ± 0.1 inch, and a diameter of 8 inches ± 1/4 inch, with a handle attached to the top of the disk.

(3) Graduated Cylinder - Class B or better, Style III, 250 mL capacity graduated cylinder conforming to the requirements of ASTM E 1272, used to measure the amount of filler required for the test.

(4) Brushes - A stiff wire brush and a soft bristle brush, used to clean the pavement.

(5) Container - Small container, with a secure and easily removable cover, used to store 200 mL of filler.
(6) Wind Screen - A shield used to protect the test area from air turbulence created from wind or traffic.

(7) Measuring device - A tape measure, or ruler at least 12 inches long, graduated in at least 1/8 inch increments. A tape measure, or ruler at least 300 millimeters long, graduated in millimeter increments may be used if desired.

3. PREPARATION OF FILLER MATERIAL

(a) For each test location, one container with 200 mL of filler is prepared as follows:

(1) Fill the graduated cylinder to the 200 mL mark and gently tap the base of the cylinder several times on a rigid surface. Add more material to fill the graduated cylinder to the 200 mL mark. Gently tap the side of the cylinder to level the material. As necessary, add additional material, tap, and level until the filler is at the 200 mL mark.

(2) Place the measured amount of filler in the container.

4. PROCEDURE

(a) Randomly select a test location.

(b) Inspect the selected test location and ensure that it is dry, uniform, and free of unique or localized features such as cracks, joints, stripping, and patching. Clean the test location using the brushes to remove any residue, debris, or loosely bonded material. Position the wind screen around the test location.

(c) Holding the container above the test location at a height not greater than 4 inches from the pavement surface, pour the measured amount of filler into a conical pile on the test area.

(d) Lightly place the spreader on top of the conical pile, being careful not to compact the filler.

(e) Using the spreader, carefully spread the material into a circular patch filling the surface voids flush with the top of the milled pavement.

(f) Measure and record the diameter of the circular patch at four locations (intervals of 45°) to the nearest 1/8 inch. Convert each reading to the nearest millimeter (1 inch = 25.4 millimeters). If desired, measurements may be taken directly in millimeters and recorded to the nearest millimeter.
Example: (For a reading of 9-3/8"")

\[ 9-3/8" = 9.375" \]

\[ 9.375" \times 25.4 = 238.125 \text{ millimeters} = 238 \text{ millimeters (rounded to the nearest millimeter)} \]

5. **CALCULATIONS**

(a) Calculate the average diameter of the circular patch as follows:

\[ D_a = \frac{D_1 + D_2 + D_3 + D_4}{4} \]

Where:

- \( D_a \) = Average diameter of the circular patch, nearest millimeter
- \( D_1, D_2, D_3, D_4 \) = Individual diameters of the circular patch, nearest millimeter

(b) Using the average diameter of the circular patch, determine the Mean Macrotexture Depth by referencing Figure 1. The values for Mean Macrotexture Depth shown in Figure 1 have been determined using the equation for Mean Macrotexture Depth shown below. When necessary, use the equation to determine the Mean Macrotexture Depth for average diameters outside the range of values given in Figure 1.

Since: Volume = (Area) x (Height, or Depth)

Therefore: Height, or Depth \( = \frac{\text{Volume}}{\text{Area}} \)

so: Mean Macrotexture Depth, mm \( = \frac{V}{(\pi) \times \left(\frac{D_a}{2}\right)^2} \)

Where: \( V = \text{Volume of glass beads, 200,000 mm}^3 \ (200 \text{ mL}) \)
- \( D_a = \text{Average diameter of the circular patch, mm} \)

6. **REPORT AND EXAMPLE**

(a) An example of a completed Mean Macrotexture Depth test report is provided in Figure 2.

(b) A blank Mean Macrotexture Depth test report form is provided in Figure 3.
### MEAN MACROTERRACE DEPTH
(Based on 200 mL of Glass Beads and Average Diameter)

<table>
<thead>
<tr>
<th>Average Diameter (mm)</th>
<th>Mean Macrotexture Depth (mm)</th>
<th>Average Diameter (mm)</th>
<th>Mean Macrotexture Depth (mm)</th>
<th>Average Diameter (mm)</th>
<th>Mean Macrotexture Depth (mm)</th>
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**FIGURE 1**
# MEAN MACROTEXTURE DEPTH
## ARIZONA TEST METHOD 742

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<th>$D_2$ (mm)</th>
<th>$D_3$ (mm)</th>
<th>$D_4$ (mm)</th>
<th>$D_5$ (mm)</th>
<th>Mean Macrotexture Depth Spec.</th>
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<th>Fail (✓)</th>
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**EXAMPLE**
### MEAN MACROTEXTURE DEPTH

**ARIZONA TEST METHOD 742**

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<th>Offset</th>
<th>( D_1 ) (mm)</th>
<th>( D_2 ) (mm)</th>
<th>( D_3 ) (mm)</th>
<th>( D_4 ) (mm)</th>
<th>( D_5 ) (mm)</th>
<th>Mean Macrotexture Depth</th>
<th>Spec.</th>
<th>Pass (✓)</th>
<th>Fail (✗)</th>
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</table>

(03/18/08)
TITANIUM DIOXIDE IN PAINTS AND THERMOPLASTICS

(An Arizona Method)

1. SCOPE

1.1 This test method may be used to make a rapid estimation of the amount of Titanium Dioxide pigment in paints and thermoplastics, particularly those of white color used for traffic control markings.

1.2 The method is a non-instrumental, gravimetric method, which involves removing, successively, all accompanying constituents of the material, leaving only the Titanium Dioxide analyte. The isolated analyte is weighed directly.

1.3 The method is applicable for paints and for thermoplastics, with or without glass beads.

1.4 The procedure in this method consists of two parts. Part A is for paints; Part B is for thermoplastics.

1.5 This test method involves hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.6 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 600 mL beaker, graduated.
2.1.2 Whatman No. 40 ashless filter paper, 125 mm, or equivalent.

2.1.3 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except that the readability and sensitivity of any balance or scale utilized shall be at least 0.0001 gram.

2.1.4 50 mL porcelain crucible.

2.1.5 Filtration funnel, 7.5 cm, 60°, long stem.

2.1.6 Watch glass, 3 inch.

2.1.7 Glass stirring rod.

2.1.8 Muffle furnace, capable of maintaining a constant temperature of 450 °C.

2.1.9 Drying oven, capable of maintaining a constant temperature of 110 °C.

2.1.10 Dessicator.

2.1.11 Hot plate.

3. REAGENTS

3.1 Concentrated Hydrochloric Acid Solution.

3.2 Concentrated Nitric Acid Solution.

3.3 Demineralized water.

4. PROCEDURE

4.1 PART A (FOR PAINTS):

4.1.1 Weigh 5.0000 ± 0.2500 grams of paint into a crucible. Record the weight of the paint specimen as “B” to the nearest 0.0001 gram.

4.1.2 Place crucible and specimen into the drying oven at temperature of 110 °C.
4.1.3 After 2 hours, remove crucible, place into cold muffle furnace, and set the temperature to 450 °C.

4.1.4 Ignite specimen at 450 °C for 1 hour after the temperature of 450 °C is attained.

4.1.5 Remove crucible with ignition residue and allow to cool to room temperature.

4.1.6 Place crucible with residue into 300 mL beaker.

4.1.7 Cautiously, dropwise, add 30 mL Concentrated Hydrochloric Acid to crucible.

4.1.8 Cautiously add 10 mL Concentrated Nitric Acid to crucible.

4.1.9 After digestion of the residue is complete, add 250 mL of demineralized water to beaker, add 1 boiling chip, cover with watchglass, and place on the hot plate.

4.1.10 Heat beaker contents to boiling and boil for 15 minutes.

4.1.11 Remove beaker from hotplate and filter contents, carefully washing and decanting titanium dioxide, leaving boiling chip and glass beads, if any, in the beaker. Wash thoroughly with demineralized water.

4.1.12 Tightly fold filter and place into tared crucible. Record crucible weight as “A,” to the nearest 0.0001 gram.

4.1.13 Place crucible and contents into cold furnace and set temperature to 450 °C.

4.1.14 Ignite specimen at 450 °C for 1 hour after temperature of 450 °C is attained.

4.1.15 Remove crucible and allow to cool to room temperature in dessicator.

4.1.16 Weigh crucible and contents. Record weight as “C,” to the nearest 0.0001 gram.
4.2 **PART B (FOR THERMOPLASTICS):**

4.2.1 Weigh 5.0000 ± 0.2500 grams of thermoplastic into a crucible. Record the weight of the thermoplastic specimen as “B” to the nearest 0.0001 gram.

4.2.2 Place crucible and specimen into cold muffle furnace and set the temperature to 450 °C.

4.2.3 Proceed as specified in Subsections 4.1.4 through 4.1.16.

5. **CALCULATIONS AND REPORT**

5.1 Calculate Titanium Dioxide, in percent, using the following formula:

\[
\text{Titanium Dioxide, } \% = \frac{C - A}{B} \times 100
\]

5.2 Report Titanium Dioxide to the nearest 0.001%. 
ROCK SALT IN CRASH BARREL SAND

(An Arizona Method)

1. SCOPE

1.1 This test method is used to determine the amount of rock salt which has been mixed with sand for use in crash barrel cushions.

1.2 The method is a non-instrumental, gravimetric method, which involves removing the salt from the sand/salt mixture by dissolution and washing away the salt using demineralized water. The salt content is then determined by weight loss after drying.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. APPARATUS

2.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. Apparatus for this test procedure shall consist of the following:

2.1.1 1000 mL heavy-duty glass beaker, graduated; KIMAX No. 14005, or equivalent.

2.1.2 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except that the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.1.3 Stirring rod (glass, steel, or plastic).

2.1.4 Drying oven, capable of maintaining a constant temperature of 110 °C.

2.1.5 Spot plate (black).
3. **REAGENTS**

3.1 Silver Nitrate Test Solution, 1%. Weigh approximately 0.5 gram of Silver Nitrate crystals into a 100 mL beaker and dilute to 50 mL. Stir to dissolve, and transfer contents to an amber dropping bottle.

3.2 Demineralized water.

4. **PROCEDURE**

4.1 Weigh a clean beaker. Record the weight as “A” to the nearest 0.1 gram.

4.2 Weigh 500.0 ± 5.0 grams of sand/salt mixture into the beaker. Record the weight of the sand/salt mixture as “B” to the nearest 0.1 gram.

4.3 Add enough demineralized water to cover the specimen and fill the beaker to the 750 mL mark.

4.4 Stir the contents of the beaker vigorously and let the mix stand for one hour.

4.5 Decant and discard the supernatant extract solution.

4.6 Repeat Subsections 4.3 and 4.4, and then continue to Subsection 4.7.

4.7 Test two drops of the extract solution for the presence of chloride with two drops of the silver nitrate solution, in the spot plate.

4.8 If the test indicates the presence of chloride (by a white cloudy solution), repeat Subsections 4.3, 4.4, and 4.7. If the test indicates the absence of chloride, decant and discard the extract solution.

4.9 Place the beaker and wet sand residue into the drying oven. Dry thoroughly at 110 °C.

4.10 After drying, remove the beaker with dry sand residue from the oven and allow to cool to room temperature.

4.11 Weigh the beaker and contents. Record the weight as “C” to the nearest 0.1 gram.
5. **CALCULATIONS AND REPORT**

5.1 Calculate the salt content, in percent, using the following formula:

\[
\text{Salt, \%} = \frac{(A + B) - C}{B} \times 100
\]

5.2 Report as Percent Rock Salt, to the nearest 0.1\%.
SERIES 800
DESIGN
EVALUATION OF PROFILES

(A Modification of California Test 526)

1. SCOPE

1.1 This method describes the procedure for determining the Profile Index (PI) from profilograms of pavements made with a California type profilograph. Also described is the procedure used to locate individual high areas in excess of 0.3 inches when their reduction is required by the contract documents.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 See the archived ARIZ 801a for information regarding mechanical recording, determination of profile index by the manual method, or determination of high points by the manual method.

2. APPARATUS

2.1 The California Profilograph consists of a frame 25 feet in length supported upon wheels at both ends. The California Profilograph is fabricated with a truss structure for stability. Beams or other systems are not permitted.

The profilograph records the roadway surface profile from the vertical movement of a wheel attached to the frame at mid-point (profile wheel) and is in reference to the mean elevation of the points of contact with the roadway surface established by the support wheels.

The profilograph shall use an electronic recording device.

2.2 Electronic recorder – The record of the roadway surface is recorded on a scale of 1 inch = 25 feet longitudinally and 1 inch = 1 inch vertically (full scale). Data is
collected by means of a digital response resulting from the vertical movement of the profile wheel.

3. CALIBRATION

3.1 The profilograph must be calibrated both horizontally and vertically per the manufacturer’s recommendations. These calibrations and the profilograph repeatability should be evaluated prior to use in the project, weekly during use, and at any time verification may be necessary.

3.2 Vertical calibration is required after every profile wheel change and each reassembly of the profilograph. The air pressure of the profile wheel shall be checked daily to make sure it is within the manufacturer’s recommendation.

3.3 The following records shall be maintained:

3.3.1 Vertical and horizontal calibration following assembly and prior to use, weekly during use, and any other time vertical and horizontal calibration data is obtained.

3.3.2 Daily tire pressure checks and the associated calibration information if tire pressure changed beyond the manufacturer’s recommendation.

3.4 Horizontal calibration shall be performed on a straight, flat roadway test section at least 528 feet (0.1 mile) long measured accurately to within 1 foot (or 0.2%) of the length. The roadway test section shall be verified by a measuring tape or wheel.

3.5 Vertical calibration shall be performed on a flat, level area. Use vertical deflection standards that are flat plates of known thickness or a single device with graduated thickness. The thickness of the initial plate or initial step of the graduated device must not exceed 1.0 inch.

3.5.1 Raise the profile wheel and place it on the initial plate or initial step of the graduated device. Ensure that the initial plate or graduated device is firmly seated. This will place the profile wheel on a flat surface and establish a baseline value from which to measure subsequent elevations. Record the elevation displayed on the electronic profilograph.

3.5.2 Raise the profile wheel again and insert another plate on top of the initial baseline plate or slide the graduated device to the next elevation. Ensure that
the graduated device is firmly seated. Record the displayed elevation from the electronic profilograph. Perform this step for at least a 1 inch and 2 inch change in elevation from the initial elevation.

**Note:** Recorded elevations shall be accurate within 0.01 inches of the known thickness of the plates or graduations.

3.5.3 Reverse the process by removing the two individual plates one at a time or stepping down the graduated device and recording the change in elevation after the removal of each successive plate.

3.5.4 The calibration is considered complete if the recorded elevation returns to within 0.03 inches of the original starting position.

3.6 Adjust chart deviations in excess of 0.03 inches according to the manufacturer’s recommendations.

3.7 The profilograph must be able to demonstrate acceptable repeatability. Acceptable repeatability is defined as “after three tests, the difference in the measured PI must not exceed a PI of one between any two tests.” The pavement surface used for the repeatability test must have a PI value of 15 or less.

4. **PROCEDURE**

4.1 The profilograph shall be operated in accordance with the manufacturer’s instructions.

4.2 The instructions for assembling the profilograph are contained in a booklet supplied for each unit by the manufacturer.

4.3 Clear the intended profilograph path of all loose material and foreign objects.

4.4 If possible, move the profilograph about 30 feet forward to the starting point. Once there, initialize the recorder and make beginning notations.

4.5 The profilograph shall be moved at a speed no greater than a walk, approximately 2 to 3 mph, to eliminate as much bounce as possible. Higher speeds can result in a profilogram with excessive spikes (chatter) that is difficult to evaluate.
4.6. Steer the profilograph to stay within the prescribed testing path. Pertinent observation about surveyed location or unusual conditions may be made on the record only as they occur. Observe the recorder for any unusual operation.

4.7. Upon completion of a sampling path, make ending notations and review the recording. Repeat the procedure for successive sampling paths.

5. **CALCULATION**

5.1 PI is defined as – inches per 0.1 miles in excess of a zero (null) blanking band.

5.2 Zero (null) blanking band is defined as – a reference line that balances the profile above and below it. The blanking band shall be 0.2 inches unless otherwise specified.

5.3 The PI can be determined from the data collected by a profilograph using an electronic recording device capable of creating a profile trace (profilogram). The profilogram indicates the PI for the required distance as well as the location of all scallops.

5.4 The PI from an electronic profilograph shall be with a software program capable of generating a computerized profile trace from the collected data. The computer software shall be set with the following data filter settings:

5.4.1 The PI from an electronic profilograph is calculated according to the instructions in the software program.

5.4.2 Determination of high points in excess of 0.3 inches using an electronic profilograph will be automatic according to the instructions in the software program.

6. **REPORT**

6.1 Height of the blanking band to the nearest 0.05 inches

6.2 Cutoff height to the nearest 0.05 inches (for example 0.3 inches)

6.3 Profilograph make and model

6.4 Length of each segment for which the profile index is calculated
EFFECT OF WATER ON STRENGTH OF COMPACTED, TREATED AND UNTREATED BITUMINOUS MIXTURES (IMMERSION COMPRESSION TEST)

(A Modification of AASHTO Designation T 167 and ASTM D 1075)

1. SCOPE

1.1 This method covers measurement of the loss of strength resulting from the effect of water on compacted bituminous mixtures. A numerical index of retained strength is obtained by comparing the compressive strength of freshly molded and cured specimens with the compressive strength of duplicate specimens that have been immersed in water under prescribed conditions. Provisions are also given for testing specimens which have been treated with a mineral admixture. With some modifications this procedure may be used in testing recycled bituminous mixtures or emulsion mixes.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

2. REFERENCED DOCUMENTS

2.1 ARIZ 415Bulk Specific Gravity and Bulk Density of Compacted Bituminous Mixtures
ARIZ 815Marshall Mix Design Method for Asphalitic Concrete

2.2 AASHTO M 231Weighing Devices Used in the Testing of Materials
AASHTO T 167Compressive Strength of Hot Mix Asphalt
AASHTO T 316Viscosity Determination of Asphalt Binder Using Rotational Viscometer
3. **APPARATUS**

3.1 Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

3.2 Molds and Plungers – Molds and plungers conforming to the requirements of AASHTO T 167.

3.3 Support Bars – Steel bars to hold the mold cylinder one inch above the baseplate during molding operation.

3.4 Testing Machine – A testing machine conforming to the requirements of AASHTO T 167.

3.5 Ovens – A minimum of two ovens that are controllable to within ± 5 °F of any temperature specified in this method.

3.6 Hot Plate – A small controllable hot plate shall be provided under the mixing bowl to maintain the mix at the desired temperature during mixing.

3.7 Hot Water Bath – An automatically controlled water bath of sufficient size to permit total immersion of the test specimens. It shall have a perforated false bottom or be equipped with a shelf. Either one shall support the specimens at least 1 inch above the bottom of the bath. The bath and shelf or false bottom shall be either lined with or constructed of a non-reactive material. It shall provide accurate and uniform control of a temperature of 140 ± 2 °F.

3.8 Water Bath – A separate water bath conforming to Section 3.7 above and which provides accurate and uniform control for bringing immersed specimens to a temperature of 77 ± 2 °F for the compression test.

**Note:** The water used in both of the above baths shall be distilled water. Only one set of specimens shall be in the 140 °F bath at a time. The 140 °F bath shall be emptied, cleaned, and refilled with fresh distilled water for each set of specimens. The 77 °F water bath shall be drained and cleaned on a regular basis.
3.9 Air Bath – An automatically controlled air bath for storing the specimens at 77 ± 2 °F.

3.10 Balance – A balance capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance utilized shall be at least 0.1 gram.

3.11 Mixer – A commercial grade mixer conforming to the requirements of AASHTO T 167.

3.12 Miscellaneous Equipment – A flexible spatula for scraping the mixing bowl and a stiff spatula, approximately 1" wide and 8" long, for spading the mix in the mold. A supply of transfer plates approximately 5" x 5" made of glass or non-reactive material for transferring the molded specimens.

4. PREPARING AGGREGATE-MINERAL ADMIXTURE SAMPLES

4.1 Based on the stockpile composite aggregate gradation, the aggregate samples needed for the immersion compression (IMC) test are prepared as follows.

4.2 Dry the mineral aggregate from each individual stockpile at a temperature not exceeding any temperature restrictions in Section 5. Drying shall be performed until no further weight loss is obtained from continued drying.

4.3 Representative samples of aggregate material which are retained on the individual No. 8 and larger sieve sizes and the Minus No. 8 Material from each stockpile are used to prepare the samples for mix design testing.

4.4 Weigh up three 3400 gram samples of mineral aggregate plus the required percent of mineral admixture, by dry weight of the aggregate, to yield three sets of two IMC specimens.

**Note:** Generally the weight of mineral aggregate will provide specimens of acceptable heights, but adjustments may be necessary in some cases. Use the following equation below to adjust the weight of aggregate as necessary to conform to specimen height requirements of 4.000 ± 0.100 inches for IMC specimens.
The aggregate-mineral admixture samples shall be dried to constant weight within ± 5 °F of the laboratory mixing temperature and shall be at this temperature at the time of mixing with the asphalt binder. If necessary, a small amount of proportioned Minus No. 8 aggregate make-up material shall be added to bring samples to the desired weight.

5. LABORATORY MIXING TEMPERATURES AND BATCHING PROCEDURE

5.1 The rotational viscosity of the asphalt binder at 275 °F and 350 °F shall be determined in accordance with AASHTO T 316, and a viscosity-temperature curve developed in accordance with ASTM D 2493.

5.2 The laboratory mixing temperature range is defined as the range of temperatures where the un-aged asphalt binder has a rotational viscosity of 0.17 ± 0.02 Pascal-seconds. The actual laboratory mixing temperature used is normally selected at or near the mid-point of the range.

5.3 Alternatively, the viscosity-temperature curve may be found in the mix design report. For PG asphalt binders that have a maximum laboratory mixing temperature exceeding 325 °F or for modified asphalt binder, refer to the binder manufacturer to establish appropriate mixing temperature ranges. In no case shall the mixing temperature exceed 350 °F.

Note: When IMC testing is performed with Warm Mix Asphalt (WMA) technology, testing shall be performed with and without the WMA technology. The test results, both with and without the WMA technology, shall meet the minimum requirements of the specifications.

The WMA technology must be added to the mix before testing in accordance with the WMA technology manufacturer’s recommendations. The WMA technology shall be added at the rate anticipated to be used in the production of the asphaltic concrete.
The mixing temperature for the laboratory prepared samples shall be per the WMA technology manufacturer’s recommendations, but shall not exceed the maximum anticipated mixing temperature during field production. In making laboratory mixing temperature recommendations, the WMA technology manufacturer should consider the mixing temperature based on the viscosity-temperature curve for the asphalt which has been modified with the WMA technology as well as the minimum mixing temperature required for adequate coating.

5.4 Before each batch of asphaltic concrete is mixed, the asphalt binder shall be heated in a loosely covered container in a forced draft oven for approximately 2 hours or as necessary to bring the asphalt binder to within ± 5 °F of the laboratory mixing temperature. (Avoid prolonged heating of the binder.)

5.5 Calculate the weight of asphalt binder to be used as determined by the following equation:

\[
\text{Weight of Asphalt Binder} = \left( \frac{\text{Weight of Aggregate and Mineral Admixture}}{100 - \left( \frac{\text{Percent of Asphalt Binder}}{\text{Percent of Asphalt Binder}} \right)} \right) \times \text{Percent of Asphalt Binder}
\]

Percent of asphalt binder is based on the mix design asphalt binder content.

5.6 Preheat the mixing bowl and whip to within ± 5 °F of the laboratory mixing temperature. The aggregate-mineral admixture blend and the appropriate amount of asphalt binder shall be mechanically mixed together for 90 to 120 seconds within ± 5 °F of the required laboratory mixing temperature. After mechanical mixing, hand mixing shall be used as necessary to produce a well-coated homogeneous mixture.

5.7 Immediately after mixing, place the hot material on a tarp or a sheet of heavy paper large enough to manipulate the sample. Thoroughly scrape the bowl and whip and add this material to the sample. In a rolling motion, thoroughly mix the material. Leave the mound in a circular shape after rolling is completed. Spread the material into a circular mass. Spreading may be accomplished either by leveling the mound of material with a concrete trowel or hand float; or by placing a straightedge of sufficient length to span the final diameter of the circular mass over the center of the material and rotating it until the desired
height is obtained. Whichever method is utilized, the operator shall assure that the material is evenly distributed with as little segregation as possible. The thickness of the circular mass shall not exceed 3 inches. The circular mass shall be cut into 4 equal pie-shaped segments. Take opposite segments for each individual sample and use the entire batch.

6. MOLDING AND CURING TEST SPECIMENS

6.1 Place the mixtures in an oven maintained at 255 ± 5 °F for 2 hours ± 10 minutes. A mold and bottom plunger for each mixture shall also be heated to 255 ± 5 °F.

Note: For mixtures with WMA technology, the samples shall be at a compaction temperature of 255 ± 5 °F, unless an alternative compaction temperature is recommended by the WMA technology manufacturer and approved by the Engineer.

For WMA water foaming processes, if laboratory water foaming equipment is not available, the specimens for IMC testing may be fabricated from plant produced mix. The specimens shall be tested as described above except the specimens shall be compacted without allowing the mixture to cool after the sample is obtained. Reheating, aging, or curing will not be allowed.

6.2 Remove the bottom plunger and mold cylinder from the oven, and place the mold assembly on the baseplate (bottom plunger in place with the mold cylinder supported on the two steel bars). Place the paper disc on the bottom plunger to prevent material from adhering to the plunger. Place 1/2 of the mixture into the molding cylinder and spade the mixture vigorously with a heated, flat, metal spatula with a blade approximately 1" wide and 8" long, stiff enough to penetrate an entire layer of material, 15 times around the edge of the mold and 10 times at random into the mixture, penetrating the mixture to the bottom of the mold. Place the remaining half of the mixture into the mold and repeat the spading process, penetrating into the first lift of the mixture. The top of the mixture should be slightly rounded to aid in firm seating of the upper plunger.

6.3 Place a paper disc and then the upper plunger (which has been preheated) on the sample and compress the mixture under an initial load of 150 psi, to set it against the sides of the mold. Remove the support bars and permit full double-plunger action. Apply the load to the mixture at a rate of 0.2 inches per minute until a load of 2750 psi is reached. Hold the load at 2750 psi for 2 minutes.
Note: A load of 2750 psi will generally produce specimens that meet the criteria specified for air voids in Section 7. The loading may be varied, if necessary, to a minimum of 2000 psi. However, in all cases the requirements for air voids must be met. Record the load at which the specimens are prepared.

6.4 Remove the specimen from the mold. During and after extrusion from the mold, take care to maintain the specimen's shape and prevent tensile stresses in the specimen. Place the specimen on a transfer plate.

Note: The specimen may be allowed to cool for a maximum of 10 minutes before removal from the mold.

6.5 Place the specimen and plate in an oven at 140 ± 5 °F.

6.6 Repeat Sections 6.2 through 6.5 for the other mixtures.

6.7 Cure the specimens for 18 ± 0.5 hours at 140 ± 5 °F.

7. BULK DENSITY DETERMINATION

7.1 After removal from the 140 °F oven, allow the specimens to cool on the plate to room temperature.

Note: Cooling may be accomplished in a 77 °F air bath or if more rapid cooling is desired the specimen may be placed in front of a fan until cooled to room temperature.

7.2 Determine and record the bulk density in accordance with ARIZ 415 (Method A) and record the height of each specimen to the nearest 0.001 inches.

Note: The bulk densities between specimens shall not differ by more than 2.5 lbs/cu ft. If this criterion is not met the entire set of specimens shall be discarded and a new set prepared.

Note: The mixture shall be compacted to 7.0 +/- 1.0 percent air voids based on the mix design maximum specific gravity. The standard molding load of 2750 psi may be increased or decreased to achieve a target air voids.
7.3 Sort the six specimens into two groups of three specimens each, so that the average bulk density of the specimens in group 1 is essentially the same as for group 2. Test the specimens in group 1 as specified in Section 8. Test the specimens in group 2 as specified in Section 9.

8. DETERMINATION OF DRY STRENGTH

8.1 Bring the test specimens to testing temperature by storing in an air bath maintained at 77 ± 2 °F for 4 to 5 hours.

8.2 Remove each specimen from the air bath and test each specimen in axial compression without lateral support at a uniform rate of vertical deformation of 0.2 inches per minute. Record the load failure point for each specimen and the average load failure point. Determine the dry strength by converting the average load failure point to pounds per square inch.

Note: At least two of the individual load failure points shall be within ± 10% of the average load failure point for the three specimens. If this criterion is not met the entire set of 6 specimens shall be discarded and a new set prepared. If only two of the three specimens meet this criterion, a new average load failure point is determined using the two values.

9. DETERMINATION OF WET STRENGTH

9.1 Immerse the test specimens in a water bath maintained at 140 ± 2 °F, at least 1 inch below the top of the water, on a transfer plate, for 24 ± 0.5 hours.

9.2 Transfer the specimens to a 77 ± 2 °F water bath for 2 hours keeping them on the plates and making sure all specimens are totally immersed.

9.3 Remove each specimen from the water bath and test each specimen in axial compression without lateral support at a uniform rate of vertical deformation of 0.2 inches per minute. Record the load failure point for each specimen and the average load failure point. Determine the wet strength by converting the average load failure point to pounds per square inch.

Note: At least two of the individual load failure points shall be within ± 10% of the average load failure point for the three specimens. If
this criterion is not met the entire set of 6 specimens shall be discarded and a new set prepared. If only two of the three specimens meet this criterion, a new average load failure point is determined using the two values.

10. CALCULATIONS

10.1 The index of retained strength shall be expressed as the percentage of "dry strength" of the specimens. It shall be calculated as follows:

\[
\text{Index of Retained Strength} = \left( \frac{\text{Wet Strength of Specimens}}{\text{Dry Strength of Specimens}} \right) \times 100
\]

10.2 The index of retained strength shall be reported to the nearest whole percent.
## Density Determination of Dry IMC Specimens:

<table>
<thead>
<tr>
<th>SPEC. #</th>
<th>HEIGHT</th>
<th>SSD WT.</th>
<th>H₂O WT.</th>
<th>AIR WT.</th>
<th>SP. GR.</th>
<th>DENSITY</th>
<th>AVERAGE SP. GR.</th>
<th>AVG. IMC DENSITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.988</td>
<td>1830.8</td>
<td>1017.5</td>
<td>1823.4</td>
<td>2.242</td>
<td>139.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>3.988</td>
<td>1827.3</td>
<td>1015.2</td>
<td>1819.5</td>
<td>2.240</td>
<td>139.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3.986</td>
<td>1828.6</td>
<td>1015.2</td>
<td>1821.5</td>
<td>2.239</td>
<td>139.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>3.981</td>
<td>1829.5</td>
<td>1016.8</td>
<td>1820.6</td>
<td>2.240</td>
<td>139.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>3.988</td>
<td>1833.6</td>
<td>1020.5</td>
<td>1824.7</td>
<td>2.244</td>
<td>139.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>3.991</td>
<td>1837.1</td>
<td>1024.3</td>
<td>1826.9</td>
<td>2.248</td>
<td>140.0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[ \text{MIX DESIGN MAXIMUM DENSITY} = \frac{\text{AVG IMC DENSITY}}{150.5} \]

\[ \% \text{ AIR Voids} = \left( 1 - \frac{\text{AVG IMC DENSITY}}{\text{MIX DESIGN MAX DENSITY}} \right) \times 100 = 1 - \left( \frac{139.7}{150.5} \right) \times 100 = 7.2\% \]

LOAD: [ ] 250 psi [ ] Other: __________ psi

## Compressive Strength of Dry Specimens:

<table>
<thead>
<tr>
<th>SPEC. #</th>
<th>LOAD FAILURE POINT</th>
<th>AVG. LOAD FAILURE POINT</th>
<th>PSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>4170</td>
<td>4260</td>
<td>338.9</td>
</tr>
<tr>
<td>4</td>
<td>4280</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>4330</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

## Compressive Strength of Wet Specimens:

<table>
<thead>
<tr>
<th>SPEC. #</th>
<th>LOAD FAILURE POINT</th>
<th>AVG. LOAD FAILURE POINT</th>
<th>PSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3610</td>
<td>3780</td>
<td>300.7</td>
</tr>
<tr>
<td>2</td>
<td>3840</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>3890</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[ \text{INDEX OF RETAINED STRENGTH} = \left( \frac{\text{PSI (WET)}}{\text{PSI (DRY)}} \right) \times 100 = \left( \frac{300.7}{338.9} \right) = 89\% \]

**FIGURE 1**
CENTRIFUGE KEROSENE EQUIVALENT OF AGGREGATE
INCLUDING K-FACTOR
(A Modification of Test No. Calif. 303)

Scope

1. This method of test furnishes a measure of the surface capacity, including absorption, of both coarse and fine aggregates used in bituminous mixtures. The K-factor is an index that indicates the relative particle roughness or degree of porosity.

Apparatus

2. The apparatus shall consist of the following:

(a) Centrifuge. - A hand or power-driven centrifuge with cover, capable of exerting a force of 400 times gravity (400 G) on a 100-g. sample (See Figs. 1 and 2). The required rpm necessary to achieve this force is determined from the following formula:

\[ \text{Required rpm of centrifuge head} = \sqrt{\frac{16,000,000}{R}} \]

Where: \( R = \text{Radius in inches to center of gravity of sample} \)

(b) Centrifuge Cups. - Aluminum cups 2-13/16 in. in height and 2-1/16 in. in diameter (Fig. 3) with a perforated brass plate containing a minimum
FIGURE 2
DETAILED PLAN OF CENTRIFUGE PARTS

Material: "68 Hard Cast Bronze or equal.
Spider and Gimbals must balance.
All dimensions ±.005 except as noted.

Full Scale
DETAIL DRAWING OF
CENTRIFUGE CUP
Scale Full Size
Total weight 215 grams
± 3 grams.

HALF TOP VIEW
HALF BOTTOM VIEW

TOP COLLAR
BODY
Aluminum
Screen
BOTTOM RING

HALF ELEVATION
HALF SECTION

Steel

HALF SECTION

Do Not Undercut

Knut

HALF ELEVATION

TOP VIEW

BRONZE BOTTOM RING

FIGURE 3
of 100 holes, .062 inches in diameter, per square inch.

(c) Balance. - A torsion balance, 500 g. capacity, accurate to 0.1 g.

(d) Metal funnels. - Top diameter 3.5 in., height 4.15 in., orifice 0.5 in., with a piece of No. 10 sieve soldered to the top of the orifice.

(e) Class Beakers, 1500 ml. capacity.

(f) Timer with sweep second hand.

(g) Pans, 4.5 in. diameter, 1 in. deep.

(h) Ovens. - One oven capable of maintaining a temperature of 230°F ± 5°F; and an oven capable of maintaining a temperature of 140°F.

Materials

3. (a) Kerosene, 1 gallon.

(b) S.A.E. No. 10 lubricating oil.

NOTE: Kerosene and S.A.E. No. 10 lubricating oil shall be supplied as "standard" materials by Central Lab.

(c) Filter paper, 5.5 cm. diameter, Eaton Dikeman No. 613.

Preparation of Sample

4. (a) Separate the aggregate into two size groups; "C" material, used for K_c determination, passing the 3/8" sieve and retained on the No. 4 sieve; "F" material for K_f determination, all passing the No. 4 sieve.

(b) Determine Bulk O.D. specific gravity for fine aggregate ("F" material) by ARIZ 211 - Specific Gravity and Absorption of Fine Mineral Aggregate, and for coarse aggregate ("C" material) by AASHOT T 85 - Specific Gravity and Absorption of Coarse Aggregates.

NOTE: Bulk O.D. specific gravity which has been determined for plus No. 4 material using additional sizes may be used in lieu of running specific gravity for "C" material as described above.

(c) The gradation of the aggregate employed is used to calculate the surface area of the aggregates. This calculation consists of multiplying the total percent passing each sieve by a "surface area factor" as set forth in Table 1, and dividing by 100. The sum of all the results represents the equivalent surface area of the sample in terms of square feet per pound.

NOTE: All surface area factors must be used in the calculation. If a different series of sieves is used, different surface area factors are necessary. An example of calculations is given in Table II, section 13.

Procedure

5. Fine aggregate shall be tested as follows:

(a) Obtain a representative minimum 210 g. sample of Pass No. 4 material, place in an oven at 230°F ± 5°F, and dry to constant weight.

(b) Allow the sample to cool to room temperature and place exactly 100 g. in weighed centrifuge cups fitted with the perforated metal disk under-lying a disk of filter paper.

(c) Prepare the other centrifuge cup in the same manner, with another 100 g. of the same material.

(d) Place the centrifuge cups in a pan containing sufficient kerosene (11/2 in. deep) to saturate the samples by capillary action.

(e) When the samples are saturated, place the cups in the centrifuge, and spin for 2 minutes at a force of 400 G (for hand operated centrifuges this force can be developed by turning the handle about 45 revolutions per minute).

(f) Reweight the cups with the samples to the nearest 0.1 g. and subtract the original weight. The difference is the percent of kerosene retained (based on 100 g. of dry aggregate). Record the average of the two values for duplicate samples.

NOTE: If the difference between the two samples exceeds 0.30 grams the test should be run again, taking greater care in obtaining duplicate aggregate samples.

6. Coarse aggregate shall be tested as follows:

(a) Split out two 105 gram minimum representative samples of the passing 3/8 and retained No. 4 sieve material. (If a 1/4 in. sieve was employed in the coarse sieve analysis, the material retained

<table>
<thead>
<tr>
<th>TABLE I</th>
</tr>
</thead>
<tbody>
<tr>
<td>SURFACE AREA FACTORS</td>
</tr>
<tr>
<td>Total Percent Passing</td>
</tr>
<tr>
<td>Sieve No.</td>
</tr>
<tr>
<td>Surface Area Factor</td>
</tr>
</tbody>
</table>

*Assume surface area for all of plus No. 4 material = 2 sq. ft./lb.
on the 1/4 in. and No. 4 sieve shall be graded in proportion with the weight % retained on the respective sieves.) Place in an oven at 230° F ± 5° F. and dry to constant weight.

(b) Allow to cool at room temperature, then weigh out exactly 100 grams from one of the 105 gram samples and transfer to the funnel. Immerse material in S.A.E. No. 10 lubricating oil for 5 minutes by placing funnel in holder (a 500 ml. glass beaker works very well) and pouring adequate amount of oil over aggregate until sample is entirely covered.

(c) Remove funnel and place in a metal quart can. Drain for 2 minutes, then place can with funnel in an oven set at 160° F. for 15 minutes of additional draining.

(d) Obtain the tare weight of a pan, remove the sample from the oven and pour into the pan, allow to cool and then weigh to the nearest 0.1 gram. The difference between the original sample weight (100 grams) and the final sample weight (weight minus the tare weight of pan) is the percent oil retained.

(e) Subject the other 105 gram sample to the same procedure as shown in paragraphs (b), (c) and (d).

(f) Record the average of the two values for duplicate samples.

NOTE: If the difference between the two samples exceeds 0.40 gram the test should be run again, taking greater care in obtaining duplicate aggregate samples.

Calculations

7. \( K_f \)

(a) \( K_f \) is determined from the following factors:

1) Percent of kerosene retained, which represents the total effect of superficial area, the aggregate's absorptive properties, and surface roughness.

2) Computed surface area, based on particle size.

3) Percent passing No. 4 sieve.

(b) If the specific gravity for "g" is greater than 2.70 or less than 2.60 apply the correction to oil retained by the following formula:

\[
\text{Percent oil ret. corrected} = \left( \frac{\text{Percent oil ret.}}{\text{Spec. Gr. } "g"} \right) \times 2.65
\]

(c) Start in the lower left hand corner of the chart in Fig. 4 with the value for CKE corrected for specific gravity; follow the straightedge horizontally to the right to the intersection with calculated surface area; hold the point, move vertically upward to the intersection with the percent passing the No. 4 sieve, hold the point, and follow the straightedge horizontally to the right. The value obtained shall be the surface constant for the passing No. 4 fraction "F", or "Kf".

8. \( K_c \)

(a) \( K_c \) is determined from the percent of S.A.E. No. 10 oil retained, which represents the total effect of superficial area, the aggregate's absorptive properties, and surface roughness.

(b) If the specific gravity for "c" is greater than 2.70 or less than 2.60 apply the correction to oil retained by the following formula:

\[
\text{Percent oil ret. corrected} = \left( \frac{\text{Percent oil ret.}}{\text{Spec. Gr. } "c"} \right) \times 2.65
\]

(c) Start at the bottom of the chart in Fig. 5 with the corrected percent oil retained, follow the straightedge vertically upward to the intersection with the diagonal line, hold the point, and follow the straightedge horizontally to the left. This value is the surface constant for the retained coarse fraction "C", or "Kc".

9. \( K_m \)

(a) \( K_m \) represents the mean value of \( K \) for a given combination of coarse and fine materials on which \( K_c \) and \( K_f \) have already been determined.

(b) Use the chart in Fig. 6 to combine \( K_f \) and \( K_c \) for determination of "Km".

(c) Start in the lower left hand corner of the chart in Fig. 6 with the value of the total Surface Area, follow straightedge horizontally to the right to the percentage of coarse aggregate (+ No. 4), hold point, and follow straightedge vertically upward to the intersection with the difference between \( K_m \) and \( K_c \), hold point, and move horizontally to the right to the "correction to \( K_f \)". If \( K_c - K_f \) is positive \( K_m \) is calculated by \( K_m = K_f + \text{corr. to } K_f \); if \( K_c - K_f \) is negative \( K_m \) is calculated by \( K_m = K_f - \text{corr. to } K_f \). If \( K_c = K_f \), the correction is zero, and the chart does not have to be used, \( K_m \) simply equals \( K_f \).

10. Bitumen Ratio

(a) The chart in Fig. 7 is used, provided the following factors are known:

1) \( K_m \)

2) S.A. (Surface Area of Aggregate)

3) Combined Specific Gravity

\[
\text{Combined Sp. Gr.} = \frac{100 \times \% \text{ coarse}}{\text{sp. gr. coarse}} + \frac{\% \text{ fine}}{\text{sp. gr. fine}}
\]

Combined Sp. Gr.
(b) Start in the upper left-hand corner with S.A., follow a straight edge horizontally to the right to the intersection with the combined sp. gr., hold point, and move vertically downward to the intersection with known Kc, hold point, and move horizontally to the right. This value is for liquid asphalt SC-250, NC-250, and NC-250 only.

(c) If a heavier asphalt is used, refer to the chart in Fig. 8, by means of a straight edge, connect the point on Scale "A" which represents the grade of bitumen to be used with the point on Scale "B" representing the S.A. of the aggregate. Through the point of intersection on line "C" place the straight edge to connect with the previously determined bitumen ratio value on Scale "D". The intersection with Scale "E" represents the bitumen ratio corrected for viscosity of the bitumen.

11. Percentage of Recommended Bitumen

To convert bitumen ratio to percentage of bitumen on the basis of total mix, use the following formula:

\[ \text{% recommended bitumen} = \frac{\text{Bitumen Ratio}}{100 + \text{Bitumen Ratio}} \]

Report

12. Report the % bitumen recommended. Also record Kc, Kf, Km, the grade of bitumen used, specific gravity of coarse and fine aggregate, CKE values, and bitumen ratio.

Example

13. Using the information given in Table II, and that supplied in (a), the data in paragraph (b), (c), (d) and (e) was calculated by the procedure outlined above.

(a) Percent coarse = 38
Percent fine (pass #4) = 62
Bulk O.D. Sp. Gr. coarse = 2.589
Bulk O.D. Sp. Gr. fine = 2.613
Combined Bulk O.D. \( \frac{38}{2.589} + \frac{62}{2.613} = 2.604 \)

CKE Results

Fines - uncorrected ret. = 4.3
corrected ret. = 4.1 (No correction: 2.7 < sp. gr. > 2.6)

Coarse - uncorrected ret. = 3.7
corrected ret. = 3.6

(b) \( K_f = 1.25 \)
\( K_c = 1.54 \)
\( (K_c - K_f) = 1.54 - 1.24 = 0.30 \)
"Correction to \( K_f \) = 0.1 \( K_m = K_f + \text{correction to } K_f \) (\( K_c - K_f \) is positive)
\( K_m = 1.24 + 0.1 = 1.34 \)

(c) Bitumen Ratio = 5.4

(d) Corrected Bitumen Ratio = 6.3
(Corrected using AC-20 grade asphalt)

(e) Percent Bitumen recommended = 5.9%

---

**TABLE II**

**TOTAL SURFACE AREA CALCULATION**

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Percent Passing</th>
<th>x</th>
<th>S.A. Factor</th>
<th>( \frac{100}{=} )</th>
<th>Surface Area (S.A.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/8&quot;</td>
<td>100</td>
<td>-</td>
<td>-</td>
<td>2.0</td>
<td></td>
</tr>
<tr>
<td>1/8&quot;</td>
<td>96</td>
<td>-</td>
<td>-</td>
<td>1.24</td>
<td></td>
</tr>
<tr>
<td>No. 4</td>
<td>62</td>
<td>2</td>
<td>2</td>
<td>1.24</td>
<td></td>
</tr>
<tr>
<td>No. 8</td>
<td>31</td>
<td>4</td>
<td>4</td>
<td>1.76</td>
<td></td>
</tr>
<tr>
<td>No. 16</td>
<td>22</td>
<td>8</td>
<td>-</td>
<td>2.24</td>
<td></td>
</tr>
<tr>
<td>No. 30</td>
<td>16</td>
<td>14</td>
<td>14</td>
<td>4.20</td>
<td></td>
</tr>
<tr>
<td>No. 50</td>
<td>14</td>
<td>30</td>
<td>30</td>
<td>7.20</td>
<td></td>
</tr>
<tr>
<td>No. 100</td>
<td>12</td>
<td>60</td>
<td>60</td>
<td>15.36</td>
<td></td>
</tr>
<tr>
<td>No. 200</td>
<td>9.6</td>
<td>160</td>
<td>-</td>
<td>35.24 sq. ft./lb.</td>
<td></td>
</tr>
</tbody>
</table>

Total Surface Area = 35.24 sq. ft./lb.
CHART FOR DETERMINING $K_f$ FROM C.K.E.

C.K.E. Corrected = C.K.E. x $\frac{\text{sp. gr. fine}}{2.65}$

FIGURE 4
CHART FOR DETERMINING $K_C$
FROM
COARSE AGGREGATE ABSORPTION

Material Used
- Aggregate passing $\frac{3}{8}$" ret. * sieve
- Oil SAE 10

$\%$ oil ret. corrected = $\%$ oil ret. x \( \frac{\text{sp. gr. of aggregate}}{2.65} \)

$K_C$ may also be determined by the following:

$$K_C = 0.4(C) + (C - 2)(0.01) + 0.1$$

Where: $C =$ Percent Oil Retained (Corrected for Sp. Gr.)

---

**FIGURE 5**
CHART FOR COMBINING $K_f$ AND $K_c$
TO DETERMINE $K_m$

If $(K_c - K_f)$ is negative,

$$K_m = K_f - \text{correction to } K_f$$

If $(K_c - K_f)$ is positive,

$$K_m = K_f + \text{correction to } K_f$$

If $K_c = K_f$, $K_m = K_f$

FIGURE 6
CHART FOR DETERMINING BITUMEN RATIO

PROCEDURE

Given surface area, combined sp. gr., and $K_m$ of aggregate.

Find surface area on scale A. Proceed horizontally to line corresponding to combined sp. gr. of aggregate. Then down to line corresponding to $K_m$. Then horizontally to scale B for Bitumen Ratio.

Bitumen Ratio = lbs. of bitumen per 100 lbs. of aggregate and applies directly to asphalt of SC 250, MC 250, and RC 250 grades. The bitumen ratio is converted to percent bitumen (based on total mix) by the following:

% Bitumen = \( \frac{\text{Bitumen Ratio}}{100 + \text{Bitumen Ratio}} \times 100 \)

When heavier liquid or paving asphalts are used a correction must be made to the bitumen ratio. See Figure 8.
Paving Grade Asphalts

Liquid Asphalts
SC, MC, RC

PROCEDURE

By means of a straightedge, connect the point on scale A, which represents the grade of bitumen to be used, with the point on scale B representing the surface area of the aggregate. Through the point of intersection on line C, place a straightedge connecting with the previously determined bitumen ratio value on scale D. The intersection of the straightedge with scale E represents the bitumen ratio corrected for viscosity of the bitumen. Bitumen Ratio = lbs. of bitumen per 100 lbs. of aggregate.

The bitumen ratio is converted to percent bitumen (based on total mix) by the following:

\[
\% \text{ Bitumen} = \frac{\text{Bitumen Ratio}}{100 + \text{Bitumen Ratio}} \times 100
\]

CHART FOR CORRECTING BITUMEN RATIO DUE TO INCREASING VISCOSITY

FIGURE 8
MAXIMUM THEORETICAL SPECIFIC GRAVITY OF LABORATORY PREPARED BITUMINOUS MIXTURES (RICE TEST)

(A Modification of AASHTO T 209)

SCOPE

1. This method of test is intended for determining the maximum specific gravity of uncompacted bituminous mixtures which have been prepared in the laboratory.

APPARATUS

2. The apparatus shall consist of the following:

   (a) Balance - A balance accurate to 0.1 gram at the maximum weight to be determined.

   (b) Container - A heavy walled Erlenmeyer flask having a capacity of at least 1500 ml. strong enough to withstand a partial vacuum; the cover shall consist of a rubber stopper with a tight hose connection. A small piece of No. 200 wire mesh covering the hose opening shall be used to minimize the possibility of loss of fine material.

       NOTE: If a procedure which subjects multiple flasks to a vacuum simultaneously is used, the vacuum gauge shall be placed beyond the last bottle to insure that all the bottles are being subjected to the same amount of vacuum.

   (c) Glass cover plate large enough to cover mouth of the flask.

   (d) Vacuum pump for evacuating air from the container.

   (e) Ovens - One oven capable of maintaining a temperature of 255°F ± 5°F and one capable of maintaining approximately 300°F.

   (f) Distilled water - All water used in this procedure shall be distilled water.

   (g) Mixer - A mechanical commercial dough mixer with a minimum 10 quart capacity and equipped with a wire whip.
CALIBRATION OF FLASK

3. Calibrate the flask by accurately determining the weight of water at 77° ± 1°F required to fill it completely. Use of a glass cover plate is required to ensure accurate filling. Record the weight of the flask + water to the nearest 0.1 gram as "B".

PREPARATION OF SAMPLES

4. (a) One aggregate sample of 3000 grams, proportioned in that grading determined by design, is required for each test. The 3000 gram aggregate sample will provide 3 test samples when mixed and split as described below.

(b) To minimize stripping of the asphalt from the aggregate, pre-weigh a 300 to 500 gram sample of asphalt and add to this 1.0% of a suitable anti-stripping agent by weight of asphalt. This mixture will be designated as "binder" in this test.

NOTE: Pave Bond Special is the preferred anti-stripping agent.

(c) The percent binder used shall normally be 6.0%.

NOTE: A binder content of 6.0% will normally give satisfactory results, although aggregate characteristics and gradation may require adjustment in percentage of binder to be used.

(d) The temperature of the asphalt and aggregate at the time mixing begins shall be in accordance with the following:

<table>
<thead>
<tr>
<th>Asphalt Grade</th>
<th>Temperature Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC 20</td>
<td>300° ± 10°F</td>
</tr>
<tr>
<td>AC 30, AC 40</td>
<td>305° ± 10°F</td>
</tr>
</tbody>
</table>

(e) The aggregate shall be dried to a constant weight, at the temperature required as shown in paragraph 4 (d). Bring samples to desired weight by adding a small amount of proportioned Pass No. 8 makeup material.

(f) The aggregate and asphalt shall be mixed mechanically for 90 to 120 seconds and then hand mixed as necessary to ensure thorough coating.

NOTE: Before each batch is mixed, the mixing bowl and whip shall be at approximately the temperature specified in 4 (d).
(g) The mixed sample shall be placed on a tarp or sheet of heavy paper and in a rolling motion thoroughly mixed. The material shall be spread into a circular mass 1 1/2 to 2 inches thick. The circular mass shall be cut into 6 equal segments taking opposite segments for each individual sample.

(h) Place samples in an oven maintained at 255° ± 5°F and cure for 2 hours ± 5 minutes. After curing, spread each sample on a sheet of heavy paper or in a large flat bottom pan. Before the samples are completely cooled separate the particles of the mixture, taking care not to fracture the mineral particles, so that the particles of the fine aggregate portion are not larger than 1/4 inch. Allow the samples to cool to room temperature.

PROCEDURE

5. For each sample the procedure below shall be followed:

(a) Place the sample in the flask and determine weight of sample to the nearest 0.1 gram. This is designated as the "weight of sample in air", or "Wmm".

NOTE: Care should be taken in mixing and transferring of the samples to the flask so that there is no more than an 18 gram difference between the total weight of aggregate and binder before mixing and the total "weight of the samples in air".

(b) Add sufficient water, which has been treated with a wetting agent, to cover the sample.

NOTE: A suitable wetting agent such as Aerosol OT in a concentration of 0.01%, or 1 ml. of 10% solution per 1000 ml. of water, shall be used to facilitate the release of entrapped air.

(c) Remove entrapped air by subjecting the contents of the flask to a partial vacuum, with a minimum of 20 inches mercury for 15 ± 2 minutes. Agitate the contents three or four times within this period, to dislodge trapped air bubbles.

CAUTION: Do not agitate the sample too frequently or vigorously; this can cause stripping of the film from some particles, resulting in erroneous specific gravities.

(d) After the evacuation period, fill the flask completely with water, slide a preweighed glass cover plate over the mouth of the flask, and weigh immediately to the nearest 0.1 gram. The temperature of the flask, water, and sample shall be between 72°F and 80°F. Record as the weight of the "flask + water + sample", or "C".
(e) The entire contents of the flask shall be poured into a nest of sieves consisting of a No. 40 and a No. 200 screen.

NOTE: If stripping has occurred, as evidenced by discoloration of water in the flask, significant loss of minus No. 200 material may be expected. Provisions for the recovery and addition of this material to the Plus No. 200 material shall be made.

(f) Allow mix to drain through the sieves until excessive moisture is removed from mix. Spread the material retained on the No. 40 and No. 200 sieves in a pan and place before an electric fan to remove surface moisture. The air through the fan shall be at room temperature and no heat shall be used to dry the mix.

(g) After evaporation of excess moisture is observed, weigh mix at 15 minute intervals and when the weight loss is less than 0.5 gram for this interval the mix is considered to be surface dry. Record the surface dry weight as "Wsd". Intermittent stirring of the sample is required during the drying period. Conglomerations of mix shall be broken by hand. Care must be taken to prevent loss of particles of mixture.

NOTE: If the "Wsd" weight for any of the three samples is less than its corresponding "Wmm" weight the samples shall be discarded and a set of three new samples shall be prepared and tested.

**CALCULATIONS**

6. (a) The Volume of Voidless Mix, "Vvm", in ml. is determined for each sample by the following:

\[ Vvm = Wsd + B - C \]

Where:  
\( Wsd \) = Surface Dry Weight  
\( B \) = Weight of Flask + Water  
\( C \) = Weight of Flask + Water + Sample

(b) The Maximum Specific Gravity, "Gmm", is determined for each sample by the following:

\[ Gmm = \frac{Wmm}{Vvm} \]

(c) Compare the three individual values for maximum specific gravity. If the range of the three is within 0.014, all are used to determine the average maximum specific gravity as
shown in paragraph (d) below. If the range is greater than 0.014, the average of two may be used if they are within a range of 0.007. If values are not achieved within the above criteria, the samples shall be discarded and a set of three new samples shall be prepared and tested.

NOTE: For recycle mixes the above criteria shall be replaced with 0.024 for the range of three and 0.012 for two.

(d) The average maximum specific gravity of the bituminous mix is determined for the samples with acceptable maximum specific gravity values, and recorded to the nearest 0.001 unit.

(e) To determine the maximum density, the average maximum specific gravity is multiplied by 62.3 lbs/cu ft.

EXAMPLE

7. An example of the calculations is shown in Figure 1.

NOTE: The form shown in Figure 1 contains equations which shall be used to determine the combined bulk oven dry specific gravity of aggregate, the effective specific gravity of aggregate, and the asphalt absorption. These values are utilized in determining the voids relationships for the bituminous mixture. The areas provided on the form in Figure 1 may be conveniently used to perform these calculations.
### Table

<table>
<thead>
<tr>
<th>Flask Number</th>
<th>Weight of Flask</th>
<th>Weight of Sample in Air</th>
<th>Weight of Flask + Water</th>
<th>Surface Dry Weight</th>
<th>Volume of Voidless Mix (Wsd+B-C)</th>
<th>M.S. Density of Mix (Gmm)</th>
<th>Maximum Density of Mix (Asphalt SP. Gr.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1185.9</td>
<td>1059.6</td>
<td>3429.9</td>
<td>4046.7</td>
<td>1058.4</td>
<td>441.6</td>
<td>2.395</td>
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<tr>
<td>2</td>
<td>1280.0</td>
<td>1059.2</td>
<td>3499.2</td>
<td>4116.1</td>
<td>1060.3</td>
<td>443.4</td>
<td>2.389</td>
</tr>
<tr>
<td>3</td>
<td>1178.1</td>
<td>1059.2</td>
<td>3436.1</td>
<td>4053.6</td>
<td>1057.1</td>
<td>439.6</td>
<td>2.405</td>
</tr>
</tbody>
</table>

**Average:**

<table>
<thead>
<tr>
<th>Weight of Flask</th>
<th>Weight of Sample in Air</th>
<th>Weight of Flask + Water</th>
<th>Surface Dry Weight</th>
<th>Volume of Voidless Mix (Wsd+B-C)</th>
<th>M.S. Density of Mix (Gmm)</th>
<th>Maximum Density of Mix (Asphalt SP. Gr.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1180.9</td>
<td>1059.2</td>
<td>3436.0</td>
<td>4064.8</td>
<td>1060.4</td>
<td>441.6</td>
<td>2.395</td>
</tr>
</tbody>
</table>

**MIX WT. - TOTAL SAMPLE WT. = LOSS OF GRAMS = 84**

**Flask Number:**

<table>
<thead>
<tr>
<th>Flask Number</th>
<th>WT. of Sample and Flask</th>
<th>WT. of Sample, Flasks, Water and Glass Plate</th>
<th>WT. of Glass Plate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2243.5</td>
<td>4201.8</td>
<td>155.1</td>
</tr>
<tr>
<td>2</td>
<td>2339.2</td>
<td>4271.2</td>
<td>155.1</td>
</tr>
<tr>
<td>3</td>
<td>2235.3</td>
<td>4208.7</td>
<td>155.1</td>
</tr>
</tbody>
</table>

**Average:**

<table>
<thead>
<tr>
<th>WT. of Sample and Flask</th>
<th>WT. of Sample, Flasks, Water and Glass Plate</th>
<th>WT. of Glass Plate</th>
</tr>
</thead>
<tbody>
<tr>
<td>2243.5</td>
<td>4201.8</td>
<td>155.1</td>
</tr>
</tbody>
</table>

**Effective Specific Gravity of Aggregate (Gse) =

\[
\frac{(100) - (\% \text{ Asphalt})}{(Gmm) (\text{Asphalt SP. Gr.)}}
\]

\[
= \frac{(100) - (6.0)}{(2.392) (1.023)} = 2.615
\]

**Asphalt Absorption (Pba) =

\[
\frac{(Gse) - (Gsb)}{(Gse) \times (\text{Asphalt SP. Gr.)}} \times (100) =
\]

\[
= \frac{(2.615) - (2.576)}{(2.615) \times (2.576)} \times (100) = 0.59
\]

**Remarks:**

**Flask #3 eliminated due to being outside 0.04 allowance range.**

**Figure 1**
DESIGN OF SLURRY SEAL

(A Modification of Test No. Calif. 355)

Scope

1. This method is intended for use in determining the total liquid content for proper workability and the optimum asphalt content for a slurry seal. This test method also furnishes an index of the slurry seal mixture’s resistance to abrasion in the presence of water.

Apparatus

2. The apparatus shall consist of the following:

(a) Centrifuge, hand or power driven, capable of exerting a force of 400 times gravity (400 G) on a 100 g. sample.

Required rpm of centrifuge head = \( \sqrt{\frac{14,000,000}{r}} \)

Where:

\( r \) = radius in inches to center of gravity of sample.

(b) Centrifuge cups, 2 13/16 in. in height, 2 1/16 in. in diameter, complete with perforated brass plate 0.031 in. thick with minimum of 100 holes, 0.062 in. in diameter, per square inch.

(c) Balance, 500 g. capacity, accurate to 0.1 g.

(d) Drying oven or hot plate.

(e) Assortment of round metal pans.

(f) Scale, 2 kg. capacity, accurate to 1 g.

(g) Circular metal rings, 10 in. in inside diameter with vertical sides ¼ in. deep.

(h) Abrasion apparatus, as shown in Fig. 2.

(i) Mechanical mixer (bituminous mix) or other power device capable of driving abrasion apparatus at 33 rpm (Fig. 1).

(j) Special plastic cylinder, 500 ml., with valve at bottom (see Fig. 4).

(k) Kerosene.

(l) Filter paper, Eaton Dikemann Co., size 5½ cm., No. 611.

(m) Roofing Felt, 50-60 lb. weight.

Preparation of the Sample

3. (a) Weigh out approximately 105 g. of representative aggregates.

(b) Dry to constant weight and allow to cool to room temperature.

Determination of Surface Area and Absorption

4. (a) Place sample in a tared centrifuge cup containing filter paper and adjust the weight of the material to exactly 100 g.

(b) Place the cup and sample in the centrifuge, quickly pour about 20 ml. of kerosene over the sample and immediately centrifuge for 2 minutes at a force of 400 G.

NOTE: The centrifuge must be in operation within 5 seconds after adding kerosene, as an additional time allows the kerosene to be absorbed by the aggregate and will not represent true surface area.

(c) Weigh centrifuged sample to the nearest 0.1 g. and record the amount of kerosene retained. Correct for specific gravity as follows:
ABRADING ASSEMBLY

LIST OF PARTS

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Part Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 ea.</td>
<td>Bassik #900 Double Wheel Casters</td>
</tr>
<tr>
<td>2 ea.</td>
<td>1/2&quot; x 2&quot; Hex Bolt</td>
</tr>
<tr>
<td>2 ea.</td>
<td>1/2&quot; Hex Nut</td>
</tr>
<tr>
<td>6 ea.</td>
<td>1/2&quot; x 1&quot; Hex Bolt</td>
</tr>
<tr>
<td>1 ea.</td>
<td>3&quot; x 9&quot; x 1&quot; Aluminum Channel</td>
</tr>
</tbody>
</table>

Sufficient Lead to make total weight of apparatus 13.5 lbs.

*Wheels machined for true flat surface.*

FIGURE 2
Correction = \( \frac{\text{(g kerosene ret.) \times \text{avg. sp. gr. agg.}}}{2.65} \)

(d) Use the surface area chart in Fig. 3. Place the straightedge horizontally to the right to the point of intersection with the diagonal line and then move vertically downward to the intersection with the bottom scale to obtain the surface area.

(e) After step (c), place the sample and cup in a small pan containing a %-in. depth of kerosene and pour approximately 35 ml. of kerosene over the sample. Allow to soak for 10 minutes.

(f) Remove the sample from the bath and centrifuge for 2 minutes at a force of 400 G.

(g) Reweigh and record the amount of kerosene retained. Make a correction for specific gravity as in step (c).

(h) Subtract the correction result in (c) above from the result of (g) and record the difference as the absorption.

**Total Liquid Content of Mixture**

5. (a) Pour 500 g. of quartered aggregate into the tared plastic cylinder (Fig. 4). Add water slowly from the bottom by attaching a length of hose between a water faucet and the valve at the bottom of the graduate until free water appears on the surface of the sample.

(b) Record the total weight of the water.

NOTE: If the aggregate column breaks up during addition of water, rod with a small metal rod until the column goes back together.

(c) The above procedure will establish the maximum amount of liquid (emulsion plus water) that the aggregate can accommodate.

**Optimum Asphalt Content**

6. (a) Quarter out 3 separate 500-g. samples of aggregate for fabrication of test specimens.

(b) Add 5.0 percent water by weight of the dry material, to the first test sample and mix thoroughly. Plot the surface area of the material on Fig. 5 or Table 1 to determine the amount of emulsion required to give an asphalt film thickness of 9 microns. Add this amount of emulsion plus sufficient water to bring to the previously established total liquid content (section 5). This added amount of water will be in addition to the 5.0 percent added for initial mixing. Mix thoroughly.

(c) Place a metal ring on a disk cut from roofing felt to the outside diameter of the ring, and flow the mixed material into the ring. Strike off excess material felt to the outside diameter of the ring, and flow the mixed material into the ring. Strike off excess material and allow to stand at room temperature for 15 minutes.

(d) Remove the metal ring and place the test specimen with supporting disk in a 140°F oven to constant weight. Record the weight, after allowing specimen to cool to room temperature.

(e) In order to provide a range for visual inspection and evaluation of the abrasion test, repeat the above procedure on the other 2 samples using appropriate percentages of asphalt required (Fig. 5) to provide about 1 micron film thickness above and below the 9 micron level.

**Resistance to Abrasion**

7. (a) Place the 3 specimens that were prepared in section 6 in pans and cover each with 350 ml. of water at room temperature. Allow to soak for 1 hour.

(b) Place the pan with specimen and water in abrading unit and abraid for 15 minutes.

(c) Remove the specimen from the bath and dry in a 140°F oven to constant weight. Subtract this weight from the oven-dry weight before the abrasion test (section 6(d) above). This difference represents abrasion loss.

(d) Repeat steps (b) and (c) for the other 2 specimens. Multiply the grams lost in abrasion by 6.20. This equal grams abraided per square foot. The quantity 6.20 is a constant designating the area encompassed by the wheel track (1/6.20 sq. ft.). If the dimensions of the wheels are changed, this factor must be adjusted.
NOTE: Determining optimum asphalt content for a slurry seal mixture will depend on the judgment of the individual. This mixture after curing should present a dense, voidless mastic-like appearance, black in color, with no visible signs of free asphalt. Excessive loss in the abrasion test will normally be indication of insufficient asphalt and an excess of asphalt will usually be evidenced by the mixture picking up on the rubber wheels of the abrasion apparatus.

An abrasion loss in excess of 100 g. per square foot of surface on the specimen determined to have optimum asphalt content will be cause for rejection of the mixture. Causes of the excessive abrasion loss for this specimen can usually be attributed to hydrophylic aggregates or possibly to a faulty emulsion which tends to separate and inadequately coat the aggregate particles. Additional mixtures should be tested in which the emulsion in question is mixed with an aggregate of known non-hydrophylic characteristics. In addition, the aggregates in question should be tested with an emulsion from a different source to isolate the cause of the abrasion loss.
SURFACE AREA CHART


\[ \frac{2.65}{2} \]

FIGURE 3
NOTE: CYLINDER & BASE TO BE OF MOULDED TRANSPARENT ACRYLIC PLASTIC

500 ML CYLINDER
**NOTE:** Figures in parenthesis are percent of emulsion to give indicated percent of asphalt based on 60% asphalt in the emulsion. Use actual percent of asphalt if test report available.
DESIGN OF ASPHALTIC CONCRETE
FRICITION COURSE

(An Arizona Method)

1. SCOPE

1.1 This test method provides a standard methodology to be used for the development of an asphaltic concrete friction coarse mix design. It includes a means of determining the density and design binder content.

1.2 This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 A listing of subsequent Sections and Figures in this procedure is given below:

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<thead>
<tr>
<th>Section or Figure</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
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<td>13</td>
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</tbody>
</table>
2. TEST METHODS AND APPARATUS

2.1 This test method is used in conjunction with the test methods listed below. The required apparatus is shown in the individual test methods, as appropriate. This test method also requires specific apparatus for the draindown and unit weight procedure, as listed in Subsection 2.5. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual.

2.2

| ARIZ 201   | Sieving of Coarse and Fine Graded Soils and Aggregates |
| ARIZ 205   | Composite Grading                                      |
| ARIZ 210   | Specific Gravity and Absorption of Coarse Aggregate    |
| ARIZ 211   | Specific Gravity and Absorption of Fine Aggregate      |
| ARIZ 212   | Percentage of Fractured Coarse Aggregate Particles     |
| ARIZ 233   | Flakiness Index of Coarse Aggregate                     |
| ARIZ 238   | Percent Carbonates in Aggregate                         |
| ARIZ 242   | Sand Equivalent Test for Mineral Aggregate for Asphaltic Concrete Friction Course |
| ARIZ 248   | Alternate Procedures for Sieving of Coarse and Fine Graded Soils and Aggregates |
| ARIZ 251   | Combined Coarse and Fine Specific Gravity and Absorption |
| ARIZ 714   | Sieving of Granulated Rubber                           |
| ARIZ 806   | Maximum Theoretical Specific Gravity of Laboratory Prepared Bituminous Mixtures (Rice Test) |
2.3 AASHTO T 96  Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine  
AASHTO T 176  Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test  
AASHTO T 228  Specific Gravity of Semi-Solid Bituminous Materials  

Note: Perform testing by AASHTO T 228 at 77 °F.  

AASHTO M 231  Weighing Devices Used in the Testing Materials  
AASHTO T 316  Viscosity Determination of Asphalt Binder Using Rotational Viscometer  

2.4 ASTM D 7741  Measurement of Apparent Viscosity of Asphalt-Rubber or Other Asphalt Binders by Using a Rotational Hand Held Viscometer  

2.5 Apparatus for Draindown and Unit Weight Procedure  

2.5.1 Oven – An oven capable of maintaining constant temperatures between 220 and 335 °F.  

2.5.2 Compression Testing Machine – A testing machine conforming to the requirements of AASHTO T 167.  

2.5.3 Molds – Molding cylinders, and top and bottom molding plungers; the cylinders shall be 4.000 ± 0.005 inches inside diameter by approximately 6 inches in height. The top and bottom plungers shall be 3.985 ± 0.005 inches in diameter, the bottom plunger shall be approximately 2 inches in height and the top plunger shall be approximately 6 inches in height. A baseplate approximately 8” x 8” shall be provided to hold the mold and plunger assembly.  

Note: Identify the molding cylinders and record the tare weight to the nearest 0.1 gram, and the inside diameter recorded to the nearest 0.001 inch. Check the tare weight and inside diameter of the cylinders periodically to assure they have not changed due to use.  

2.5.4 Support Bars – Steel bars to hold the mold cylinder one inch above the baseplate during the molding operation.
2.5.5 A balance or scale capable of measuring the maximum weight to be determined and conforming to the requirements of AASHTO M 231, except the readability and sensitivity of any balance or scale utilized shall be at least 0.1 gram.

2.5.6 Hot Plate – Capable of maintaining a temperature of up to 325 ± 5 °F.

2.5.7 Miscellaneous mixing bowls, spoons, spatulas, scoops, pans, etc.

2.5.8 Thermometers – Dial type with glass face, or digital, capable of reading to the nearest 1 °F with a range of 50 to at least 350 °F.

2.5.9 1,000 ml Glass Beakers.

2.5.10 Glass Plate or Watch Glass large enough to cover Glass Beaker.

3. MATERIALS

3.1 Mineral Aggregate – The mineral aggregate used in the design shall be produced material from the source(s) for the project.

3.1.1 The composited gradation of the aggregate without admixture, and the composited gradation of the aggregate-mineral admixture blend shall comply with the grading limits of the specifications.

3.1.2 The composited mineral aggregate shall conform to the requirements of the specifications for Combined Bulk Oven Dry Specific Gravity (ARIZ 251), Combined Water Absorption (ARIZ 251), Sand Equivalent (ARIZ 242), Fractured Coarse Aggregate Particles (ARIZ 212), Flakiness Index (ARIZ 233), Percent Carbonates (ARIZ 238), and Abrasion (AASHTO T 96) when applicable.

3.2 Bituminous Material – The bituminous material used in the design shall be an asphalt binder, conforming to the requirements of Section 1005 of the specifications, which is to be used in the production of the asphaltic concrete. The specific gravity of the asphalt binder shall be determined in accordance with AASHTO T 228 (at 77 °F).

3.3 Mineral Admixture – Mineral admixture is required. The mineral admixture used in the design shall be the same type of material from the same supplier to be
used in production of friction course. The mineral admixture shall conform to the requirements of the specifications.

Note: The mineral admixture used for mix design testing should be stored in an airtight container and be no older than 3 months.

3.4 Crumb Rubber Asphalt (CRA) – The Crumb Rubber Asphalt shall be tested and conform to the specifications of ADOT Specifications Section 1009.

3.5 Crumb Rubber – The crumb rubber shall be tested for sieve analysis in accordance with ARIZ 714. It shall conform to ADOT Specifications Sections 414 & 1009.

4. DETERMINATION OF COMPOSITE GRADATION

4.1 Perform a coarse and a fine sieve analysis for the aggregate from each stockpile in accordance with ARIZ 201. The fine sieve sample size may be reduced if the minimum of 500 grams is not obtained from coarse sieving, or if there is insufficient material for performing other desired tests which utilizes the Minus No. 4 Material.

4.2 Compare the Percent Pass No. 8 from the Sieve Analysis to the actual Percent Pass No. 8 as follows:

4.2.1 The Minus No. 4 Material from each stockpile is separated into Retained No. 8 Material and Minus No. 8 Material, and the weights for each recorded.

4.2.2 Determine the actual Percent Pass No. 8 by dividing the weight of Minus No. 8 Material by the combined total weight of the Plus No. 8 and Minus No. 8 Material that was determined in Subsection 4.2.1.

4.2.3 Multiply this value by the Percent Pass No. 4 from the sieve analysis. Record the result to the nearest whole percent. This is the actual Percent Pass No. 8.

4.2.4 Compare the actual Percent Pass No. 8 to the Percent Pass No. 8 from the sieve analysis. If the difference between the two results is greater than 4.0%, a new fine sieve analysis must be performed. If the difference between the two results is still greater than 4.0% after performing a new fine sieve analysis, adjust the
fine sieve analysis by multiplying the Percent Pass for each sieve smaller than the No. 8 sieve by a factor obtained by dividing the actual Percent Pass No. 8 by the Percent Pass No. 8 from the sieve analysis.

4.3 Determine the mineral aggregate composite in accordance with ARIZ 205. This is used to establish the mix design gradation which is then compared to the specifications. If the gradation falls outside of the specification band, the stockpile percentages will be adjusted until the composite is within the band. (An example is given in Figure 1 & 2).

**Note:** Adjustments of stockpile percentages greater than 5% for any stockpile should not be made without the agreement of the supplier in writing.

4.4 The aggregate-mineral admixture blend composite is determined by adjusting the mineral aggregate composite (Percent Passing) for mineral admixture by performing the calculation for each sieve:

\[
\left( \frac{\% \text{ Pass Each Sieve}}{\text{Adjusted for Mineral Admixture}} \right) = \left( \frac{\% \text{ Pass Each Sieve in the Aggregate Composite}}{100} + \frac{\% \text{ Mineral Admixture}}{\text{Admixture}} \right) \times 100
\]

4.5 The composited gradation of the aggregate and the composited gradation of the aggregate-mineral admixture blend shall be shown on the design report, along with the percentage of each material.

4.6 The design gradation is adjusted for Minus No. 200 Material adhering to the coarse aggregate by washing a composited sample of Plus No. 4 Material in accordance with ARIZ 248, alternate #5 (Referee Method). Figure 3 gives an illustration of the calculations and the adjustment to the Minus No. 200 Material from washing.

If adjustments to the gradation are necessary due to the increase in Minus No. 200 Material, a maximum of a one percent change may be made for the Percent Retained on No. 200. If the adjusted Percent Pass No. 200 results in a greater amount of change, wash a composited sample as specified in ARIZ 248 Section 5 (Alternate #4) to determine the actual design gradation.
Note: It is recommended that the composited dust correction sample be tested before additional mix design testing is performed to assure that the design gradation meets governing specifications.

5. PREPARING SAMPLES FOR MIX DESIGNS USING STOCKPILE MATERIAL

5.1 Representative aggregate samples for each test are prepared utilizing the information from the composite for individual size fractions for No. 8 sieve size and larger from each stockpile; and the Minus No. 8 Material from each stockpile.

Note: Aggregate retained on the No. 8 and larger sized sieves may be weighed up by hand for each size fraction. The individual Minus No. 8 Materials may be obtained by first splitting each Minus No. 8 Material into pans of adequate size to hold approximately 1500 grams of material, and then using the split portions for weighing up samples by stirring and scooping out the required amount with a small scoop.

5.2 Table 1 shows the aggregate sample sizes, the number of samples required for each test listed, and which samples include mineral admixture. The aggregate weight shown for Maximum Theoretical Specific Gravity will provide three Rice test specimens. An example weigh-up sheet is shown in Figure 4.
<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Sample Size</th>
<th>No. of Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coarse Aggregate Specific Gravity/ Absorption (ARIZ 210)</td>
<td>Required grams of Mineral Aggregate as determined by the Nominal Maximum Aggregate Size [Excludes mineral admixture]</td>
<td>2</td>
</tr>
<tr>
<td>Fine Aggregate Specific Gravity/ Absorption (ARIZ 211)</td>
<td>1200 grams of Mineral Aggregate [Excludes mineral admixture]</td>
<td>2</td>
</tr>
<tr>
<td>Dust (ARIZ 248)</td>
<td>1000 grams Refer to Figure 4. [Excludes mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Sand Equivalent (ARIZ 242)</td>
<td>500 to 750 grams of Mineral Aggregate [Excludes mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Fractured Coarse Aggregate Particles (ARIZ 212)</td>
<td>Required grams of Mineral Aggregate as determined by test method [Excludes mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Flakiness Index (ARIZ 233)</td>
<td>#¼” Sieve 200g #4 Sieve 100g #8 Sieve 50g [Excludes mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Percent Carbonates (ARIZ 238)</td>
<td>300 grams of Mineral Aggregate [Excludes mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Abrasion (AASHTO T 96)</td>
<td>5000 grams of Mineral Aggregate as per test method for grading type [Excludes mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Maximum Theoretical Specific Gravity (Rice Test) (ARIZ 806, as modified in Section 11)</td>
<td>3030 grams of Mineral Aggregate [Includes mineral admixture]</td>
<td>1 [Yields 3 test specimens]</td>
</tr>
<tr>
<td>Draindown</td>
<td>960 grams Refer to Section 10. [Includes mineral admixture]</td>
<td>3</td>
</tr>
<tr>
<td>Bulk Unit Weight</td>
<td>960 grams Refer to Section 11. [Includes mineral admixture]</td>
<td>3</td>
</tr>
</tbody>
</table>
6. AGGREGATE SPECIFIC GRAVITIES AND ABSORPTION

6.1 Determine the bulk oven dry (OD) specific gravity, bulk saturated surface dry (SSD) specific gravity, apparent specific gravity, and absorption of the fine and coarse mineral aggregates in accordance with Arizona Test Method 211 and Arizona Test Method 210.

**Note:** Two samples of both the fine aggregate and coarse aggregate are to be tested and the results must not exceed the maximum range as listed below in Table 2. Additional samples are to be run as necessary until results fall within these ranges.

<table>
<thead>
<tr>
<th>Fine Specific Gravity</th>
<th>Acceptable Range of 2 Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Sp. Gr.(OD)</td>
<td>0.032</td>
</tr>
<tr>
<td>Bulk Sp. Gr.(SSD)</td>
<td>0.027</td>
</tr>
<tr>
<td>Apparent Sp. Gr.</td>
<td>0.027</td>
</tr>
<tr>
<td>Absorption, %</td>
<td>0.31</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Coarse Specific Gravity</th>
<th>Acceptable Range of 2 Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Sp. Gr.(OD)</td>
<td>0.250</td>
</tr>
<tr>
<td>Bulk Sp. Gr.(SSD)</td>
<td>0.020</td>
</tr>
<tr>
<td>Apparent Sp. Gr.</td>
<td>0.020</td>
</tr>
<tr>
<td>Absorption, %</td>
<td>0.25</td>
</tr>
</tbody>
</table>

6.2 The individual averages for the fine and coarse specific gravities are then utilized in calculating the final combined specific gravities.

6.3 The combined bulk (OD) specific gravity, bulk saturated surface dry (SSD) specific gravity, apparent specific gravity, and combined absorption for the coarse and fine mineral aggregates are calculated by the following:
Combined Bulk (OD) Specific Gravity = \frac{100}{\frac{P_c}{G_c} + \frac{P_f}{G_f}}

Where: \( P_c \) and \( P_f \) = Percent of Coarse Aggregate (Plus No. 8) from the composite and Percent of Fine Aggregate (Minus No. 8), respectively. (Use the nearest whole percent in calculations.).

\( G_c \) and \( G_f \) = Specific gravity of coarse and fine aggregate respectively.

Example:
\[
\begin{bmatrix}
P_c = 86 \\
P_f = 14 \\
G_c \ (OD) = 2.741 \\
G_f \ (OD) = 2.704
\end{bmatrix}
\]

Combined Bulk (OD) Specific Gravity = \frac{100}{\frac{86}{2.741} + \frac{14}{2.704}} = 2.736

Example:
\[
\begin{bmatrix}
\text{Combined Bulk (SSD) Specific Gravity} = 2.798 \\
\text{Combined Bulk (OD) Specific Gravity} = 2.736
\end{bmatrix}
\]

\[
(\text{Combined % Absorption}) = \frac{(2.798) - (2.736)}{(2.736)} \times 100 = 2.27\%
\]

6.4 The combined aggregate bulk specific gravity (OD) must be adjusted for admixture (\( G_{sb \ adj} \)). The calculations for this are as follows:

\[
G_{sb \ adj} = \frac{100 + P_{\text{admix}}}{\frac{100}{G_{sb}} + \frac{P_{\text{admix}}}{G_{\text{admix}}}}
\]

Where: \( P_{\text{admix}} \) = Percent of Mineral Admixture, by weight of the mineral aggregate (typically 1.0%)
7. ASPHALT MIXTURES TESTING TEMPERATURE REQUIREMENTS

7.1 Maximum Theoretical Specific Gravity (ARIZ 806), Draindown Determination, and Bulk Density Determination of friction course must be performed to complete the development of a mix design. Mix and cure samples for these tests at the temperatures specified below in Table 3. The Engineer will specify the laboratory mixing and curing temperatures for all the binders not listed in Table 3.

<table>
<thead>
<tr>
<th>Asphalt Binder Type</th>
<th>Mixing Temperature</th>
<th>Curing Temperature for Rice and Draindown Test</th>
<th>Curing and Compaction Temperature for Unit Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRA</td>
<td>325 ± 5 °F</td>
<td>325 ± 10 °F</td>
<td>230 ± 5 °F</td>
</tr>
<tr>
<td>ALL PG’s &amp; TR+</td>
<td>300 ± 5 °F</td>
<td>300 ± 10 °F</td>
<td>230 ± 5 °F</td>
</tr>
</tbody>
</table>

8. DETERMINATION OF MAXIMUM THEORETICAL SPECIFIC GRAVITY (with mineral admixture)
8.1 Determine the maximum theoretical specific gravity \( G_{mm} \) of the friction course mixture in accordance with ARIZ 806 and the temperature guidelines specified in Section 7. For design purposes the binder content will be 4.0% by weight of total mix.

**Note:** The final binder content selected for a friction course design is typically much higher than the 4.0% mandated above. However, using high asphalt percentages in conjunction with open-graded mixes often results in erroneous values when performing ARIZ 806. Occasionally, 4.0% binder does not provide an adequate asphalt film coating on the aggregate which results in a negative asphalt absorption calculation. If this happens, increase the percent of binder in increments of 0.5% until satisfactory test results are obtained.

8.2 Calculate the Effective Specific Gravity of the Aggregate/Admixture \( G_{se} \) using the following formula:

\[
G_{se} = \frac{100 - P_b}{100 - \frac{P_b}{G_{mm}}} - \frac{G_{mm}}{G_b}
\]

Where:
- \( G_{se} \) = Effective specific gravity of the combined aggregate and mineral admixture.
- \( P_b \) = Asphalt binder content at which the Rice test was performed (Rice \( P_b \)).
- \( G_{mm} \) = Measured maximum theoretical specific gravity of the mix at Rice \( P_b \).
- \( G_b \) = Specific gravity of the asphalt binder.

**Example:**

\[
\begin{bmatrix}
P_b = 4.0 \\
G_{mm} = 2.628 \\
G_b = 1.034
\end{bmatrix}
\]

\[
G_{se} = \frac{100 - 4.0}{100 - \frac{4.0}{2.628}} - \frac{2.628}{1.034} = 2.808
\]
8.3 Calculate the percent asphalt absorption ($P_{ba}$) by the following:

$$P_{ba} = \left( \frac{G_{se} - G_{sb}}{G_{sb} \times G_{se}} \right) \times G_b \times 100$$

Where:
- $P_{ba}$ = Absorbed asphalt, percent by total weight of mix.
- $G_{se}$ = Effective specific gravity of the combined aggregate and mineral admixture.
- $G_{sb}$ = Bulk oven dry specific gravity of the combined aggregate and mineral admixture.
- $G_b$ = Specific gravity of the asphalt binder.

Example:

$$\begin{align*}
G_{se} &= 2.808 \\
G_{sb} &= 2.739 \\
G_b &= 1.034
\end{align*}$$

$$P_{ba} = \left( \frac{2.808 - 2.739}{2.808 \times 2.739} \right) \times 1.034 \times 100 = 0.93$$

Report the asphalt absorption to the nearest 0.01%.

9. DETERMINATION OF DESIGN ASPHALT CONTENT

9.1 Design binder content is based on the binder draindown procedure as described in Section 10. The target percent of binder draindown is 0.25% and the intent is to determine a percent of binder as close as possible to the target draindown percent without exceeding it.

Note: Often draindown tests for AR-ACFC will indicate a design binder content that is over 10%. It is current ADOT practice not to issue a friction course design over 9.9%, regardless of what draindown test results indicate.

9.2 An initial binder percent for draindown testing is calculated using one of the two following formulas:
For ACFC and ACFC Special mixes:

\[
\text{ACFC Binder Content} = (0.20)(W) + \left( \frac{(61 + P)}{5} \right) \left( \frac{1}{C} \right) + 0.1
\]

Where:
\[
\begin{align*}
W &= \text{Percent Water Absorption} \\
P &= \text{Percent Pass #8} \\
C &= \text{Combined Bulk OD Specific Gravity}
\end{align*}
\]

Example:
\[
\begin{align*}
W &= 2.27 \\
P &= 14 \\
C &= 2.736
\end{align*}
\]

\[
\text{ACFC Binder Content} = (0.20)(2.27) + \left( \frac{(61 + 14)}{5} \right) \left( \frac{1}{2.736} \right) + 0.1 = 6.0\%
\]

For AR-ACFC mixes:

\[
\text{AR ACFC Binder Content} = \left[ (0.80)(W) + 8.6 \right] \left( \frac{2.620}{C} \right)
\]

Where:
\[
\begin{align*}
W &= \text{Percent Water Absorption} \\
C &= \text{Combined Bulk OD Specific Gravity}
\end{align*}
\]

Example:
\[
\begin{align*}
W &= 2.27 \\
C &= 2.736
\end{align*}
\]

\[
\text{AR ACFC Binder Content} = \left[ (0.80)(2.27) + 8.6 \right] \left( \frac{2.620}{2.736} \right) = 10.0\%
\]

Note: The equations above are used to give an initial binder content value only. This binder content will need to be adjusted based on the actual draindown test results. A minimum of three draindown tests will be run for the mix design; one at the design asphalt content, one at approximately 0.2% below the design content, and one at approximately 0.2% above the design content. Refer to Subsection 9.1.
10. DRAINDOWN DETERMINATION

10.1 Prepare a composited aggregate-admixture sample of approximately 960 grams and dry the sample to a constant weight.

10.2 Heat the composited sample, binder, metal mixing bowl, spoons, and scrapers to the temperature specified in Section 7. Determine the weight of a 1,000 milliliter glass beaker. Record the weight to the nearest 0.1 gram.

10.3 Once the binder and aggregate have reached the proper mixing temperature remove the mixing bowl from the oven and apply a light coat of aerosol based vegetable oil to the inner surface, and wipe any excess with a towel. Place the mixing bowl on a balance and zero the balance. Carefully add the heated aggregate into the mixing bowl. Determine the weight of the aggregate and, if needed, add Minus No. 8 Make-Up Material (See Table 1) to the sample to bring the weight back to 960 grams.

10.4 Create a small crater in the center of the aggregate and add the heated binder to the sample until the total weight of mix is reached.

Note: The total weight of mix is calculated by the following formula:

\[
\text{Total Weight of Mix} = \frac{\text{Wt. of Composited Aggregate (with Admixture)} \times 100}{100 - P_b}
\]

Where: \( P_b \) = Asphalt Binder Content

10.5 Place the mixing bowl on a hot plate and quickly and thoroughly hand mix the sample using a heated spoon. Transfer the sample into the glass beaker taking care to transfer as much of the mixture as possible. Use a scraper to remove mix material adhering to the bowl and spoon, and transfer it into the beaker.

10.6 Weigh and record the weight of the beaker and mixture to the nearest 0.1 gram. Place a glass plate or watch glass over the sample and return it to the oven at mix temperature for 1 hour ± 10 minutes.

10.7 Determine the weight of a sample pan and record the weight to the nearest 0.1 gram.

10.8 After the 1 hour curing, remove the glass beaker from the oven and remove the glass plate or watch glass. Flip the beaker over and quickly empty the asphalt
mixture into the sample pan. This must be done without shaking or tapping the glass beaker, nor allowing the beaker to drain for an extended time.

10.9 Place the beaker and the pan containing the sample in front of a fan and cool to room temperature. Weigh the beaker with the asphalt residue. Then determine the final weight of the pan and asphalt mixture. Record these values to the nearest 0.1 grams.

10.10 Determine the percent draindown by the following equation:

\[
\text{Percent Draindown} = \frac{(C - A) \times 100}{(C - A) + (D - B)}
\]

Where:

- **A** = Weight of Glass Beaker
- **B** = Weight of Sample Pan
- **C** = Final Weight of Beaker and Mixture Residue
- **D** = Final Weight of Pan and Mixture

Report the draindown value to the nearest 0.01%. An example of the calculations is shown in Figure 5.

11. **BULK DENSITY DETERMINATION**

11.1 Prepare a minimum of three composited aggregate-admixture samples of approximately 960 grams each and dry the samples to a constant weight.

11.2 Heat the composited samples, asphalt binder, metal mixing bowl, and spoons to the temperature specified in Section 7.

11.3 Once the binder and aggregate have reached the proper mixing temperature, remove the mixing bowl from the oven and apply a light coat of aerosol based vegetable oil to the inner surface, and wipe any excess with a towel. Place the mixing bowl on a balance and zero the balance. Carefully add the heated aggregate into the mixing bowl. Determine the weight of the aggregate and add Minus No. 8 Make-Up Material to the sample to bring the weight back to 960 grams.

11.4 Create a small crater in the center of the aggregate sample and add enough asphalt binder to the sample to bring the asphalt content up to the design binder content that was determined in Section 10.
11.5 Place the mixing bowl on a hot plate and quickly and thoroughly mix the sample using a heated spoon.

11.6 Place the mixed sample into a 230 ± 5 °F oven until a constant temperature is attained (approximately 1 hour). Also place a calibrated mold, bottom, and a top plunger in the oven.

11.7 Repeat steps 11.3 through 11.6 for the remaining two samples.

11.8 Remove the bottom plunger and mold cylinder from oven, place mold assembly on baseplate (bottom plunger in place with mold cylinder supported on the two steel bars). Apply a light coat of aerosol based vegetable oil to the mold and bottom plunger, and wipe any excess with a towel. Place all of the mixture into molding cylinder and spade mixture vigorously with heated flat metal spatula (approximately 1" wide and 8" long, stiff enough to penetrate the entire layer of material), 15 times around the edge of the mold and 10 times at random into the mixture, penetrating the mixture to the bottom of the mold. Smooth the surface of the mix to a slightly rounded shape.

11.9 Apply a light coat of aerosol based vegetable oil to the upper plunger, and wipe any excess with a towel. Place the upper plunger (which has been preheated) on the sample and compress the mixture under an initial load of 150 psi, to set it against the sides of the mold. Remove the support bars to permit full double-plunger action. Apply the load to the mixture at a rate of 0.2 inches per minute until a load of 2,000 psi is reached. Hold the load at 2,000 ± 100 psi for 2 minutes.

11.10 After compaction, the top plunger is removed, leaving the bottom plunger in place to support the specimen until the sample has cooled sufficiently to be handled safely with bare hands.

Note: Cooling may be achieved by the use of a 77 °F air bath or, if more rapid cooling is desired, fans may be used.

11.11 Repeat the procedure as described in Subsections 11.8 through 11.10 above, for the other two samples.

11.12 After the specimens have cooled, remove the bottom plunger and determine the weight of each mold with specimen to the nearest 0.1 gram. The weight of each
specimen is calculated by subtracting the weight of the corresponding mold cylinder.

11.13 While the specimens are still in the molds, the heights of each of the three specimens are determined to the nearest 0.001 inch.

11.14 The inside diameters of the molds used for each of the three specimens is recorded to the nearest 0.001 inch.

11.15 The bulk density is determined for each specimen by the following:

\[
\text{Bulk Density} = \frac{\text{Weight of Specimen (grams)}}{(d/2)^2 (h)(50.86)} \times 62.3
\]

Where:
- \( d \) = Inside Diameter of Mold (inches)
- \( h \) = Height of Specimen (inches)

Report the densities to the nearest 0.1 lbs./cu.ft.

Example:

\[
\begin{align*}
\text{Weight of Specimen} &= 1002.7 \text{ grams} \\
\text{Inside Diameter of Mold} &= 4.002 \text{ inches} \\
\text{Height of Specimen} &= 2.489 \text{ inches}
\end{align*}
\]

\[
\text{Bulk Density} = \frac{1002.7}{(4/2)^2 \times (2.489) \times (50.86)} \times 62.3 = 123.2 \text{ lbs./cu. ft.}
\]

Note: The range of densities between all 3 samples shall not differ by more than 2.0 lbs./cu.ft. If this density requirement is not met, discard the entire set of specimens and prepare a new set of specimens.

11.16 The average bulk density of the three specimens is determined and recorded to the nearest 0.1 lbs./cu.ft.

12. **EXAMPLE AND REPORT**

12.1 Report the test results and data obtained on the appropriate form. Liberal use of the remarks area to clarify and/or emphasize any element of the design is
strongly recommended. Information required in the mix design report is shown in the items listed below.

12.2 Name of the testing organization and the signature of the mix design engineer responsible for the mix design.

12.3 Aggregate:
- Source and identification
- Individual stockpile gradations
- Aggregate blend proportions and composite gradation, with and without mineral admixture
- Fine and coarse aggregate bulk (OD) specific gravity, bulk (SSD) specific gravity, and absorption
- Combined aggregate bulk (OD) specific gravity
- Combined aggregate absorption
- LA Abrasion at both 100 and 500 revolutions
- Sand Equivalent
- Fractured Coarse Aggregate Particles (percent with two fractured faces)
- Carbonates

12.4 Mineral Admixture:
- Type and source
- Percentage used
- Specific gravity

12.5 Combined Mineral Aggregate and Mineral Admixture:
- Combined bulk (OD) specific gravity (with admix)

12.6 CRA Binder Design for AR-ACFC only (from supplier), including:
- Source and grade of base asphalt cement
- Source and type of crumb rubber
- Crumb rubber gradation
- Proportions of asphalt cement and crumb rubber
- Percentage of crumb rubber, by weight of asphalt cement
- CRA binder properties, in compliance with Section 1009 of the ADOT Specifications
12.7 Asphalt Cement for ACFC, ACFC Special, and AR-ACFC mixes:
• Source and grade of asphalt cement
• Asphalt cement properties, in compliance with Section 1005 of the ADOT Specifications
• Asphalt cement specific gravity at 77 °F

12.8 Maximum Theoretical Specific Gravity of Asphalitic Mix:
• The binder content at which the Rice test was performed
• Calculated maximum theoretical specific gravity (G_{mm}) and density (pcf) at the binder and mineral admixture content at which the Rice was performed
• Effective specific gravity of the combined aggregate and mineral admixture (G_{se})

12.9 Asphalitic Mix Properties:
• Design binder content
• Bulk density (pcf)
• Asphalt absorption

12.10 Mix Design Summary:
• An example of a mix design summary report is shown in Figure 6
# Arizona Department of Transportation

ARIZONA TEST METHOD 814

Initial Composite Gradation

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<th>Design:</th>
<th>15-09015</th>
<th>Type:</th>
<th>407 SPECIAL</th>
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* Stockpile composite out of specification

**FIGURE 1**
### Adjusted Composite Gradation

**Project Number:** XM122 01X  
**Design:** 15-09015  
**Type:** 407 Special

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<th>Material</th>
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**FIGURE 2**
FIGURE 3

Arizona Department of Transportation
ARIZONA TEST METHOD 814

Determination Of Percent Pass No. 200 On Coarse Aggregate
And Total Percent Pass No. 200 In Composite
Using Arizona 248 Alternate Procedures for Sieving of
Coarse and Fine Graded Soils and Aggregates Alternate #5

Lab #: 15-09015 Project #: XM122 01X Date: 08/19/2015

\[
\begin{align*}
O \ D \ Weight \ of \ Unwashed \ Plus \ No. \ 4 &= 993.9 \\
O \ D \ Weight \ of \ Washed \ Plus \ No. \ 4 &= 990.9
\end{align*}
\]

\[
\frac{(O \ D \ Weight \ of \ Unwashed \ Plus \ No. \ 4) - (O \ D \ Weight \ of \ Washed \ Plus \ No. \ 4)}{O \ D \ Weight \ Unwashed \ Plus \ No. \ 4} \times 100 =
\]

\[
\frac{(993.9) - (990.9)}{993.9} \times 100 =
\]

\[0.30\% \ Pass \ No. \ 200 \ on \ Coarse \ Aggregate\]

Calculate Total Percent Pass No. 200 in Sample by the following, and record to the nearest 0.1 %:

\[
\left( \frac{Total \ Pass \ No. \ 200 \ in \ Sample}{% \ Pass \ No. \ 200 \ from \ fine \ Sieving} \right) = \left( 100 - \left( \frac{% Pass \ No. \ 200 \ Coarse \ Agg}{No. \ 4} \right) \right) \times \frac{% Pass \ No. \ 200}{100}
\]

Total % Pass No. 200 in Sample = \[\left( 1.3 \right) + \left( 100 - \left( \frac{35.34}{100} \right) \right) \times \left( 0.30 \right)\]

\[= 1.5\% \ Total \ Pass \ #200\]

FIGURE 3
### FIGURE 4

**Arizona Department of Transportation**

**ARIZONA TEST METHOD 814**

**Weigh UP Sheet**

<table>
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<tr>
<th>Material</th>
<th>Sieve Size</th>
<th>Accum. % Ret.</th>
<th>Rice (3030 g)</th>
<th>Bulk Unit Wt. (960 g)</th>
<th>Dust (1000 g)</th>
<th>Coarse Specific Gravity [Plus #8] (2000 g)</th>
<th>Fractured Faces [Plus #8] (300 g)</th>
<th>Abrasion (5000 g)</th>
<th>Percent Carbonates [Plus #4] (300 g)</th>
<th>Drain Down (960 g)</th>
<th>Flakiness Index (200 gr., 100 gr., &amp; 50 g)</th>
<th>Sand Equivalent [Minus #8] (600 g)</th>
<th>Fine Specific Gravity [Minus #8] (1200 g)</th>
<th>Make-up Material [Minus #8] (600 g)</th>
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<td>309</td>
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<td>788 (118)</td>
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<td>157</td>
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<td>474</td>
<td>1000</td>
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**FIGURE 4**
BULK DENSITY EQUATION:

\[
BD = \frac{\text{Weight of Specimen}}{((d/2)^2 \text{ Height of Specimen}) (50.86))} X 62.3
\]

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<tbody>
<tr>
<td>6.4</td>
<td>1</td>
<td>4</td>
<td>2.339</td>
<td>3563.8</td>
<td>2543.3</td>
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<td>2.145</td>
<td>133.6</td>
<td>2.136</td>
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<td></td>
<td>8</td>
<td>4</td>
<td>2.353</td>
<td>3615.1</td>
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<td></td>
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<td>4</td>
<td>2.352</td>
<td>3607.5</td>
<td>2586.8</td>
<td>1020.7</td>
<td>2.133</td>
<td>132.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

BINDING DRAIN DOWN TEST

% Binder: 6.4

\[
\text{Percent Draindown} = \frac{(C - A) \times 100}{(C - A) + (D - B)}
\]

A = Wt. of Beaker: 402.0

B = Wt. of Pan: 213.6

C = Wt. of Beaker (After): 404.5

D = Wt. of Pan & Binder: 1213.3

Percent Drain Down = \[
\frac{(404.5 - 402.0) \times 100}{(404.5 - 402.0) + (1213.3 - 213.6)} = 0.25\%
\]

FIGURE 5
This design meets ADOT specifications

General Design Information

<table>
<thead>
<tr>
<th>lab #</th>
<th>PG Grade</th>
<th>PG0-22 TR+</th>
<th>PG Source</th>
<th>Admix Source</th>
<th>Admix Specific Gravity</th>
<th>Total number of stockpiles</th>
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<tr>
<td>2015-10030</td>
<td></td>
<td></td>
<td>Acme Asphalt</td>
<td>Sugaville Products</td>
<td>3.140</td>
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Aggregate:

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<th>lab #</th>
<th>Description</th>
<th>Site #</th>
<th>Design %</th>
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<tbody>
<tr>
<td>2015-00119</td>
<td>3/8&quot; Coarse Agg</td>
<td>CM 9888</td>
<td>40</td>
</tr>
<tr>
<td>2015-00118</td>
<td>3/8&quot; Fine Agg</td>
<td>CM 9888</td>
<td>42</td>
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<tr>
<td>2015-00120</td>
<td>W.C.F.</td>
<td>CM 9888</td>
<td>18</td>
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Composite Gradation

<table>
<thead>
<tr>
<th>Sieve #</th>
<th>Specification Band</th>
<th>Gradation w/o admix</th>
<th>Gradation w/ admix</th>
<th>Field Target Band</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>% passing</td>
<td>% passing</td>
<td>% passing</td>
<td>min</td>
</tr>
<tr>
<td>1 1/2&quot;</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>1&quot;</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td></td>
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<tr>
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<td></td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>64</td>
<td>64</td>
<td>64</td>
<td></td>
</tr>
<tr>
<td>#4</td>
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<td>#8</td>
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<td>#10</td>
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<tr>
<td>#16</td>
<td>9</td>
<td>10</td>
<td>10</td>
<td></td>
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<tr>
<td>#30</td>
<td>5</td>
<td>6</td>
<td>6</td>
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<tr>
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<td>#100</td>
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<td>#200</td>
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Composite Aggregate Properties

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<tr>
<th>Property</th>
<th>Test Value</th>
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<th>max</th>
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<tr>
<td>L.A. Abrasion % at 100 revolutions (AASHTO T96)</td>
<td>4</td>
<td></td>
<td>9</td>
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<tr>
<td>L.A. Abrasion % at 500 revolutions (AASHTO T96)</td>
<td>17</td>
<td></td>
<td>40</td>
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<tr>
<td>Sand Equivalent (AZ 242)</td>
<td>74</td>
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<td>55</td>
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<tr>
<td>Two Fracture Flakes, % (AZ 212)</td>
<td>59</td>
<td></td>
<td>85</td>
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<tr>
<td>Fakirness Index, % (AZ 233)</td>
<td>21</td>
<td></td>
<td>25</td>
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<tr>
<td>Carbonates, % (AZ 238)</td>
<td>2.0</td>
<td></td>
<td>20</td>
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<tr>
<td>Combined O.D. Specific Gravity (AZ 210)</td>
<td>2.736</td>
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<td>2.35</td>
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<tr>
<td>Corrected Combined O.D. Specific gravity (with admix)</td>
<td>2.739</td>
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<td></td>
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<tr>
<td>Combined Water Absorption, % (AZ 814)</td>
<td>2.27</td>
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Calculated Mix Properties Results

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<thead>
<tr>
<th>Description</th>
<th>Design Values</th>
<th>Specification Limits</th>
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<tr>
<td>Design Binder Content</td>
<td>6.4</td>
<td></td>
</tr>
<tr>
<td>Bulk Density,pcf</td>
<td>133.1</td>
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</tr>
<tr>
<td>Asphalt Absorption %</td>
<td>0.93</td>
<td>&lt; 1.0</td>
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</tbody>
</table>

FIGURE 6
FIGURE 6 (continued)
MARSHALL MIX DESIGN METHOD
FOR ASPHALTIC CONCRETE
(An Arizona Method)

1. SCOPE

1.1 This method is used to design Asphaltic Concrete mixes using 4-inch diameter Marshall apparatus.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 A listing of subsequent Sections and Figures in this procedure is given below:

<table>
<thead>
<tr>
<th>Section or Figure #</th>
<th>Title</th>
<th>Page #</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section 2</td>
<td>Test Methods and Apparatus</td>
<td>2</td>
</tr>
<tr>
<td>Section 3</td>
<td>Materials</td>
<td>3</td>
</tr>
<tr>
<td>Section 4</td>
<td>Determination of Composite Gradation</td>
<td>4</td>
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<tr>
<td>Section 5</td>
<td>Preparing Aggregate/Mineral Admixture Samples for Mix Design Testing</td>
<td>5</td>
</tr>
<tr>
<td>Section 6</td>
<td>Aggregate Specific Gravities and Absorption</td>
<td>7</td>
</tr>
<tr>
<td>Section 7</td>
<td>Laboratory Mixing and Compaction Temperatures</td>
<td>9</td>
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<tr>
<td>Section 8</td>
<td>Preparation of Specimens for Bulk Specific Gravity/Bulk Density and Marshall Stability/Flow Determination</td>
<td>10</td>
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<td>Section 9</td>
<td>Bulk Specific Gravity/Bulk Density of Specimens</td>
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<td>Section 10</td>
<td>Marshall Stability and Flow Determination</td>
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<tr>
<td>Section 11</td>
<td>Maximum Theoretical Specific Gravity (Rice Test)</td>
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<tr>
<td>Section 12</td>
<td>Determination of Design Percent Asphalt Binder Content</td>
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<tr>
<td>Section 13</td>
<td>Immersion Compression (IMC) Test</td>
<td>20</td>
</tr>
<tr>
<td>Section 14</td>
<td>Mix Design Gradation Target Values</td>
<td>20</td>
</tr>
<tr>
<td>Section 15</td>
<td>Report</td>
<td>20</td>
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<tr>
<td>Figure 1</td>
<td>Example Mix Design Composite (With and Without Mineral Admixture)</td>
<td>23</td>
</tr>
<tr>
<td>Figure 2</td>
<td>Example Weigh Up Sheet #1</td>
<td>24</td>
</tr>
<tr>
<td>Figure 3</td>
<td>Example Weigh Up Sheet #2</td>
<td>25</td>
</tr>
<tr>
<td>Figure 4</td>
<td>Example Plots of Effective Voids, VMA, Dust/Bitumen Ratio, Bulk Specific Gravity, Marshall Stability, and Marshall Flow versus Percent Asphalt Binder Content</td>
<td>26</td>
</tr>
<tr>
<td>Figure 5</td>
<td>Example Mix Design Summary</td>
<td>27</td>
</tr>
</tbody>
</table>
2. TEST METHODS AND APPARATUS

2.1 This test method is used in conjunction with the test methods listed below. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. The required apparatus is shown in the individual test methods, as appropriate.

2.2 Arizona Test Methods:

- ARIZ 201 Sieving of Coarse and Fine Graded Soils and Aggregates
- ARIZ 205 Composite Grading
- ARIZ 210 Specific Gravity and Absorption of Coarse Aggregate
- ARIZ 211 Specific Gravity and Absorption of Fine Aggregate
- ARIZ 212 Percentage of Fractured Coarse Aggregate Particles
- ARIZ 238 Percent Carbonates in Aggregate
- ARIZ 247 Particle Shape and Texture of Fine Aggregate Using Uncompacted Void Content
- ARIZ 248 Alternate Procedures for Sieving of Coarse and Fine Graded Soils and Aggregates
- ARIZ 251 Combined Coarse and Fine Aggregate Specific Gravity and Absorption
- ARIZ 410 Compaction and Testing of Bituminous Mixtures Utilizing Four Inch Marshall Apparatus
- ARIZ 415 Bulk Specific Gravity and Bulk Density of Compacted Bituminous Mixtures
- ARIZ 416 Preparing and Splitting Field Samples of Bituminous Mixtures for Testing
- ARIZ 802 Effect of Water on Strength of Compacted Treated and Untreated Bituminous Mixtures (Immersion Compression Test)
- ARIZ 806 Maximum Theoretical Specific Gravity of Laboratory Prepared Bituminous Mixtures (Rice Test)

2.1.2 AASHTO Test Methods:

- AASHTO T 96 Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine
- AASHTO T 176 Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test
- AASHTO T 228 Specific Gravity of Semi-Solid Bituminous Materials

Note 1: Testing by AASHTO T 228 shall be performed at 77 °F.
3. MATERIALS

3.1 Mineral Aggregate - The mineral aggregate used in the design shall be produced material from the source(s) for the project.

3.1.1 The composited gradation of the aggregate without admixture, and the composited gradation of the aggregate-mineral admixture blend shall comply with the grading limits of the specifications.

3.1.2 The composited mineral aggregate shall conform to the requirements of the specifications for Sand Equivalent (AASHTO T 176), Combined Bulk Oven Dry Specific Gravity (ARIZ 251), Combined Water Absorption (ARIZ 251), Fractured Coarse Aggregate Particles (ARIZ 212), Uncompacted Void Content (ARIZ 247), and Percent Carbonates (ARIZ 238) when applicable.

3.1.3 Mineral aggregate from each source shall be tested separately for compliance to the project requirements for Abrasion (AASHTO T 96).

3.2 Bituminous Material - The bituminous material used in the design shall be an asphalt binder, conforming to the requirements of Section 1005 of the specifications, which is to be used in the production of the asphaltic concrete. The specific gravity of the asphalt binder shall be determined in accordance with AASHTO T 228 (at 77°F).

3.3 Mineral Admixture - Mineral admixture is required. The mineral admixture used in the design shall be the same type of material to be used in production of the asphaltic concrete. The mineral admixture shall conform to the requirements of the specifications.

3.4 Batch Plants - Mix designs developed for asphaltic concrete which is to be produced in a batch plant shall be prepared in accordance with this procedure, modified as follows:

3.4.1 Aggregate samples will be obtained from the hot bins for all testing except Sand Equivalent. Testing for Sand Equivalent will be performed on aggregate stockpile samples proportioned and composited to meet the
mix design composite without admixture. Any method may be used to combine the aggregate stockpile samples for Sand Equivalent testing as long as the resultant gradation is representative of the mix design composite gradation without admixture.

3.4.2 The mix design shall provide component percentages and composite gradations for both stockpile and hot bin materials.

4. DETERMINATION OF COMPOSITE GRADATION

4.1 The gradation of the aggregate from each individual component stockpile or bin shall be determined in accordance with ARIZ 248 using washed sieve analysis Alternate #3, Alternate #4, or Alternate #5. For Alternate #5, washing of the coarse aggregate may be performed on the composite Plus No. 4 material and applied to the composite percent pass the minus No. 200 determined from the unwashed coarse sieving and washed fine sieving of the individual stockpiles.

4.2 The composite gradation of the mineral aggregate is determined using desired percentages of each component based on washed sieve analysis. Mix designs may be developed based on bin or stockpile material, as appropriate for the respective mix production facility to be used.

4.3 The mineral aggregate composite shall be determined in accordance with ARIZ 205, paragraph 2(e).

4.4 The aggregate-mineral admixture blend composite is determined by adjusting the mineral aggregate composite (percent passing) for mineral admixture by performing the calculation in Equation 1 for each sieve:

\[
\text{Equation 1: } \left( \frac{\% \text{ passing each sieve in the aggregate composite}}{(100 + \% \text{ Mineral Admixture})} \right) = \left( \frac{\% \text{ passing each sieve in the aggregate composite}}{(100 + \% \text{ Mineral Admixture})} \right) \times 100
\]

4.5 The composited gradation of the aggregate and the composited gradation of the aggregate-mineral admixture blend shall be shown on the design report, along with the percentage of each material.
5. PREPARING AGGREGATE/MINERAL ADMIXTURE SAMPLES FOR MIX DESIGN TESTING

5.1 Based on the stockpile or bin composite aggregate gradation, the aggregate samples needed for mix design tests are prepared as follows.

5.1.1 Dry the mineral aggregate from each individual stockpile at a temperature not exceeding any temperature restrictions specified in subsequent test procedures. Drying shall be performed until no further weight loss is obtained from continued drying.

5.1.2 Representative samples of aggregate material which is retained on the individual No. 8 and larger sieve sizes and the minus No. 8 material from each stockpile or bin are used to prepare the samples for mix design testing.

5.1.3 Table 1 shows the aggregate sample sizes, the number of samples required for each test listed, and which samples include mineral admixture. The aggregate weight shown for Maximum Theoretical Specific Gravity will provide three Rice test specimens. The amount shown for Density-Stability/Flow will produce three Marshall specimens, and the amount shown for Immersion Compression (IMC) will produce two specimens. Example weigh-up sheets are shown in Figures 2 and 3.

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Sample Size</th>
<th>Number of Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fractured Coarse Aggregate Particles (ARIZ 212)</td>
<td>Required grams of Mineral Aggregate as determined by test method</td>
<td>1</td>
</tr>
<tr>
<td>Percent Carbonates (ARIZ 238)</td>
<td>300 grams of Mineral Aggregate</td>
<td>1</td>
</tr>
<tr>
<td>Abrasion (AASHTO T 96)</td>
<td>5000 grams of Mineral Aggregate as per test method for grading type</td>
<td>1</td>
</tr>
<tr>
<td>Fine Aggregate Specific Gravity/ Absorption (ARIZ 211)</td>
<td>1200 grams of Mineral Aggregate</td>
<td>1</td>
</tr>
<tr>
<td>Coarse Aggregate Specific Gravity/Absorption (ARIZ 210)</td>
<td>Required grams of Mineral Aggregate as determined by the Nominal Maximum Aggregate Size</td>
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### Table 1 (Continued)

<table>
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<tr>
<th>Test</th>
<th>Sample Description</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sand Equivalent</strong>&lt;br&gt;(AASHTO T 176)</td>
<td>500 to 750 grams of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td><strong>Uncompacted Void Content</strong>&lt;br&gt;(ARIZ 247)</td>
<td>Minimum 500 grams of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td><strong>Minus No. 8 Make-Up Material</strong></td>
<td>An adequate amount (normally 500 grams) of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td><strong>Marshall Density-Stability/Flow</strong>&lt;br&gt;(ARIZ 415 and ARIZ 410, as modified in Sections 9 and 10 respectively)</td>
<td>3300 grams of Mineral Aggregate (See Note 2) [Plus mineral admixture, by percent required by ARIZ 802 (as modified in Section 13)]</td>
<td>3 (See Note 3) [Each sample yields 1 set of 3 Marshall Specimens]</td>
</tr>
<tr>
<td><strong>Maximum Theoretical Specific Gravity (Rice Test)</strong>&lt;br&gt;(ARIZ 806, as modified in Section 11)</td>
<td>3000 grams of Mineral Aggregate [Plus mineral admixture, by percent required by ARIZ 802 (as modified in Section 13)]</td>
<td>1 [Yields 3 test specimens]</td>
</tr>
<tr>
<td><strong>Immersion Compression (IMC)</strong>&lt;br&gt;(ARIZ 802, as modified in Section 13)</td>
<td>3400 grams of Mineral Aggregate (See Note 2) [Plus required percent of mineral admixture]</td>
<td>3 [Each sample yields 1 set of 2 IMC Specimens]</td>
</tr>
</tbody>
</table>

**Note 2:** Generally the weight of mineral aggregate will provide specimens of acceptable heights, but adjustments may be necessary in some cases. Use Equation 2 to adjust the weight of aggregate as necessary to conform to specimen height requirements of 2.500 ± 0.200 inches for Marshall Density-Stability/Flow specimens and 4.000 ± 0.100 inches for IMC specimens.

\[
\text{Adjusted Weight of Aggregate} = \left( \frac{\text{Combined Bulk O.D.}}{\text{Agg. Specific Gravity}} \right) \times 2.650 
\]

**Note 3:** Requires one sample for each asphalt binder content to be tested (minimum of 3 asphalt binder contents, with 3 Marshall specimens at each asphalt binder content).

5.1.4 After the aggregate samples for the Marshall, Rice, and IMC specimens have been composited, add the required amount of mineral admixture, by dry weight of the aggregate, and mix thoroughly. Add 3% water, by dry weight of the aggregate, to each sample and mix thoroughly to wet the mineral admixture and aggregate surfaces.
5.1.5 Testing utilizing the prepared virgin aggregate-mineral admixture samples for Marshall, Rice, and IMC will be performed as specified in Sections 8-10, 11, and 13, respectively.

6. **AGGREGATE SPECIFIC GRAVITIES AND ABSORPTION**

6.1 Determine the Bulk Oven Dry, S.S.D., Apparent Specific Gravities and Absorption for the fine aggregate (minus No. 4) and the coarse aggregate (plus No. 4) in accordance with ARIZ 211 and ARIZ 210, respectively.

6.2 Determine the combined bulk oven dry specific gravity of the fine aggregate and the coarse aggregate without mineral admixture and also the combined water absorption of the fine aggregate and the coarse aggregate without mineral admixture, in accordance with ARIZ 251.

6.2.1 The combined aggregate bulk oven dry specific gravity without mineral admixture and the combined aggregate water absorption without mineral admixture are used only to determine compliance with specification requirements.

6.3 Using Equation 3, calculate the Combined Bulk Oven Dry (Gsb), S.S.D., and Apparent Specific Gravities of the aggregate-mineral admixture blend.

\[
\text{Combined Specific Gravity of Aggregate and Mineral Admixture Blend} = \frac{P_f + P_c + P_{\text{admix}}}{G_f} + \frac{P_f + P_c + P_{\text{admix}}}{G_c} + \frac{P_{\text{admix}}}{G_{\text{admix}}}
\]

Where:
- \(P_f, P_c\) = Weight percent of fine aggregate (minus No. 4) and coarse aggregate (plus No. 4) respectively. Determined from the aggregate composite without mineral admixture.
- \(P_{\text{admix}}\) = Percent mineral admixture by weight of the aggregate.
- \(P_f + P_c = 100\)
- \(P_f + P_c + P_{\text{admix}} = 100 + \% \text{ Mineral Admixture}\)
- \(G_f, G_c\) = Bulk Oven Dry, S.S.D., or Apparent specific gravity of the fine aggregate and the coarse aggregate respectively.
**Gadmix** = Specific gravity of the mineral admixture.

- Type I or II Cement = 3.14
- Type IP Cement = 3.00
- Hydrated Lime = 2.20

Example [for combined Bulk Oven Dry Specific Gravity (Gsb)]:

\[
P_f = 59 \\
P_c = 41 \\
G_f = 2.593 \\
G_c = 2.634 \\
G_{admix} = 3.14 \text{ (Type II Cement)} \\
P_{admix} = 1.0
\]

\[
\begin{align*}
\text{Example of Combined} \\
\text{Bulk Oven Dry Specific} \\
\text{Gravity (Gsb)} \text{ of Aggregate} \\
\text{and Mineral Admixture} \\
\end{align*}
\]

\[
\begin{align*}
G_{sb} &= \frac{P_f \times G_f + P_c \times G_c + P_{admix} \times G_{admix}}{P_f + P_c + P_{admix}} \\
&= \frac{59 \times 2.593 + 41 \times 2.634 + 1 \times 3.14}{2.614} = 2.614
\end{align*}
\]

6.4 Using Equation 4, calculate the Combined Water Absorption of the aggregate-mineral admixture blend.

**Equation 4:**

\[
\left( \text{Combined Water Absorption of Aggregate and Mineral Admixture Blend} \right) = \frac{(P_f \times A_f) + (P_c \times A_c) + (P_{admix} \times A_{admix})}{P_f + P_c + P_{admix}}
\]

Where:

- \( P_f, P_c \) = Weight percent of fine aggregate (minus No. 4) and coarse aggregate (plus No. 4) respectively. Determined from the aggregate composite without mineral admixture.
- \( P_{admix} \) = Percent mineral admixture by weight of the aggregate.
- \( P_f + P_c = 100 \)
- \( P_f + P_c + P_{admix} = 100 + \% \text{ Mineral Admixture} \)
- \( A_f, A_c \) = Percent water absorption of the coarse aggregate and the fine aggregate respectively.
- \( A_{admix} \) = Percent water absorption of mineral admixture (assumed to be 0.0%).
Example: \[
P_f = 59 \\
P_c = 41 \\
A_f = 1.28 \\
A_c = 0.79 \\
P_{\text{admix}} = 1.0 \\
A_{\text{admix}} = 0.00
\]

\[
\begin{align*}
\text{Combined Water Absorption of Aggregate and Mineral Admixture} & = \frac{(59 \times 1.28) + (41 \times 0.79) + (1.0 \times 0.00)}{59 + 41 + 1.0} = 1.07
\end{align*}
\]

7. LABORATORY MIXING AND COMPACTION TEMPERATURES

7.1 The rotational viscosity of the asphalt binder at 275 °F and 350 °F shall be determined in accordance with AASHTO T 316, and a viscosity - temperature curve developed in accordance with ASTM D 2493.

7.2 The laboratory mixing temperature range is defined as the range of temperatures where the un-aged asphalt binder has a rotational viscosity of 0.17 ± 0.02 Pascal·seconds. The actual laboratory mixing temperature used is normally selected at or near the mid-point of the range.

7.3 The laboratory compaction temperature range is defined as the range of temperatures where the un-aged asphalt binder has a rotational viscosity of 0.28 ± 0.03 Pascal·seconds. The actual laboratory compaction temperature used is normally selected at or near the mid-point of the range.

7.4 The viscosity - temperature curve shall be included in the mix design report. For PG asphalt binders that have a maximum laboratory mixing temperature exceeding 325 °F or a maximum laboratory compaction temperature exceeding 300 °F, the laboratory mixing and compaction temperature ranges shall be specified in writing by the asphalt binder supplier. A viscosity-temperature curve will meet this requirement for written documentation if the viscosity-temperature curve is developed and submitted by the binder supplier and includes language that the recommended laboratory mixing and compaction temperatures are within acceptable ranges, and shall include a statement indicating the maximum laboratory mixing temperature to which the binder can be heated without damage. The laboratory mixing and compaction temperature ranges, as
well as the actual laboratory mixing and compaction temperatures used, shall be reported on the mix design.

8. PREPARATION OF SPECIMENS FOR BULK SPECIFIC GRAVITY/BULK DENSITY AND MARSHALL STABILITY/FLOW DETERMINATION

8.1 Specimens shall be prepared as described herein, utilizing the apparatus specified in ARIZ 410.

*Note 4:* Normally a range of 3 different asphalt binder contents at 0.5% increments will provide sufficient information, although in some cases it may be necessary to prepare additional sets of samples at other asphalt binder contents.

8.2 The aggregate-mineral admixture samples, prepared as specified in Section 5, shall be dried to constant weight at the laboratory mixing temperature $\pm 5^\circ F$ and shall be at this temperature at the time of mixing with the asphalt binder. If necessary, a small amount of proportioned minus No. 8 aggregate make-up material shall be added to bring samples to the desired weight.

8.3 Before each batch of asphaltic concrete is mixed, the asphalt binder shall be heated in a loosely covered container in a forced draft oven for approximately 2 hours or as necessary to bring the asphalt binder to the laboratory mixing temperature $\pm 5^\circ F$.

8.4 Using Equation 5, calculate the weight of asphalt binder to be used for each asphalt binder content:

\[
\text{Weight of Asphalt Binder} = \frac{\text{Weight of Aggregate and Mineral Admixture}}{100 - \text{Percent of Asphalt Binder}} \times \text{Percent of Asphalt Binder}
\]

Example (for Marshall specimens with 4.5% Asphalt Binder) (See Note 4):

\[
\text{Weight of Asphalt Binder} = \frac{3333}{100 - 4.5} \times 4.5 = 157.1 \text{ grams}
\]
Note 5: Before each batch is mixed, the mixing bowl and whip shall be heated to the laboratory mixing temperature ± 5 °F.

8.5 The aggregate-mineral admixture blend and the appropriate amount of asphalt binder shall be mixed together for 90 to 120 seconds at the required laboratory mixing temperature ± 5 °F. Mechanical mixing is required.

Note 6: Although a wide range of mixers may provide the desired well-coated homogeneous mixture, commercial dough mixers with whips are often used. Minimum recommended capacity of the mixing bowl is 12 quarts.

Note 7: After mechanical mixing, hand mixing shall be used as necessary to produce a well-coated homogeneous mixture.

8.6 Immediately after mixing, each batch of asphaltic concrete shall be thoroughly blended and spread according to the procedures described in ARIZ 416. The circular mass shall be cut into 6 equal pie-shaped segments. Take opposite segments for each individual specimen and use up the entire batch.

8.7 Each individual specimen shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The specimens shall be oven-cured for 2 hours ± 10 minutes at the laboratory compaction temperature ± 5 °F.

8.8 A mold assembly (base plate, mold, and collar) shall be heated to the laboratory compaction temperature ± 5 °F. The face of the compaction hammer shall be thoroughly cleaned and heated on a hot plate set at the laboratory compaction temperature ± 5 °F. A suitable shield, baffle plate, or sand bath shall be used on the surface of the hot plate to minimize localized overheating of the face of the hammer.

8.9 Place a 4-inch diameter paper disc in the bottom of the mold before the mixture is introduced. Quickly place the mixture into the mold using a transfer bowl or other suitable device. Spade the mixture vigorously with a heated flat metal spatula, with a blade approximately 1-inch wide and 6-inches long and stiff enough to penetrate the entire layer of material, 15 times around the perimeter and 10 times at random into the mixture, penetrating the mixture to the bottom of the mold. Smooth the surface of mix to a slightly rounded shape.
8.10 Before compaction, put the mold containing the mixture in an oven as necessary to heat the mixture to the required laboratory compaction temperature ± 5 °F. The specimen shall not be heated for more than one hour after being placed in the mold. Prior to compaction, it shall be verified that the mixture is at the required temperature. An electronic temperature probe with a digital display or a long-stem metal thermometer with a dial face shall be used. The use of an infrared temperature gun is not allowed.

8.11 Immediately upon removing the mold assembly loaded with mixture from the oven, place a paper disc on top of mixture, place the mold assembly on the compaction pedestal in the mold holder, and apply 75 blows with the compaction hammer. Remove the base plate and collar, and reverse and reassemble the mold. Apply 75 compaction blows to the face of the reversed specimen.

**Note 8:** The compaction hammer shall apply only one blow after each fall, that is, there shall not be a rebound impact. The compaction hammer shall meet the requirements specified in Subsection 2.4 of ARIZ 410.

8.12 Remove the collar and top paper disc. Remove the base plate and remove the bottom paper disc while the specimen is still hot.

**Note 9:** Paper discs need to be removed while the specimen is hot. The discs are difficult to remove after the specimens have cooled.

8.13 Allow each compacted specimen to cool until they are cool enough to be extruded without damaging the specimen.

**Note 10:** Generally specimens can be extruded without damage when they are at a temperature of approximately 77 to 90 °F.

**Note 11:** Cooling may be accomplished at room temperature, or in a 77 °F air bath. If more rapid cooling is desired, the mold and specimen may be placed in front of a fan until cool.

8.14 Extrude the specimen by orienting the mold so that the ram pushes on the bottom face (base plate face) of the specimen. If any specimen is deformed or damaged during extrusion, the entire set of specimens at that asphalt binder content shall be discarded and a new set prepared.
8.15 Immediately upon extrusion, measure and record the height of the specimen to the nearest 0.001 inch and determine and record its weight in air to the nearest 0.1 gram. Compacted specimens shall be 2.500 ± 0.200 inches in height. If this criteria is not met for the specimens at each asphalt binder content, the entire set of specimens at that asphalt binder content shall be discarded and a new set prepared after necessary adjustments in the weight of aggregate have been made using Equation 2 (see Note 2 in Table 1).

8.16 Repeat the procedures in Subsections 8.4 through 8.15 for the required specimens.

9. **BULK SPECIFIC GRAVITY/BULK DENSITY OF SPECIMENS**

9.1 Determine the bulk specific gravity and bulk density of the three compacted specimens at each asphalt binder content in accordance with Method A of ARIZ 415. The determination of the "Weight in Water" and "S.S.D. Weight" of each specimen will be completed before the next specimen is submerged for its "Weight in Water" determination.

**Note 12:** Specimens fabricated in the laboratory that have not been exposed to moisture do not require drying after extrusion from the molds. The specimen weight in air obtained in Subsection 9.1 is its dry weight.

9.2 For each asphalt binder content, the densities of individual compacted specimens shall not differ by more than 2.0 pcf. If this density requirement is not met, the entire set of specimens at that asphalt binder content shall be discarded and a new set of specimens prepared.

9.3 For each set of three specimens, determine the average bulk specific gravity ($G_{mb}$), and using that value calculate the average bulk density ($G_{mb}$ multiplied by 62.3 lb./cu. ft.).

10. **MARSHALL STABILITY AND FLOW DETERMINATION**

10.1 Specimens shall be tested for Marshall Stability and Flow utilizing the apparatus specified in ARIZ 410.

10.2 For each specimen, determine the uncorrected stability, the stability (corrected for height), and the flow according to ARIZ 410, Subsections 4.6 through 4.11. In addition, determine the average stability (corrected for height) and average flow for each set of specimens prepared at each asphalt binder content.
11. **MAXIMUM THEORETICAL SPECIFIC GRAVITY (RICE TEST)**

11.1 The maximum theoretical specific gravity of the mixture shall be determined in accordance with ARIZ 806 with the following modifications:

11.1.1 Prepare the specimens including mineral admixture according to the procedures described in Section 5 and Subsections 8.2 through 8.5 using 5.5% or 6.0% asphalt binder by total mix weight. A liquid anti-stripping agent is not used.

11.1.2 Immediately after mixing, the material shall be thoroughly blended and spread according to the procedures described in ARIZ 416. The circular mass shall be cut into 6 equal pie-shaped segments. Take opposite segments for each individual test sample and use up the entire batch.

11.1.3 Each individual test sample shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The test samples shall be oven-cured for 2 hours ± 10 minutes at the laboratory compaction temperature ± 5 °F.

11.1.4 After curing, each test sample shall be spread on a sheet of heavy paper or in a large flat bottom pan. Before the samples are completely cooled, separate the particles of the mixture, taking care not to fracture the mineral aggregate particles, so that the particles of the fine aggregate portion are not larger than ¼ inch. Allow the test samples to cool to room temperature.

11.1.5 Using Equation 6, calculate the effective specific gravity of the combined aggregate and mineral admixture ($G_{se}$).

\[
G_{se} = \frac{100 - P_b}{\frac{100}{G_{mm}} - \frac{P_b}{G_b}}
\]

**Where:**

- $G_{se}$ = Effective specific gravity of the combined aggregate and mineral admixture.
- $P_b$ = Asphalt binder content at which the Rice test was performed (Rice $P_b$).
- $G_{mm}$ = Measured maximum theoretical specific gravity of the mix at Rice $P_b$.
- $G_b$ = Specific gravity of the asphalt binder.
Example: \[ P_b = 5.5 \]
\[ G_{mm} = 2.436 \]
\[ G_b = 1.032 \]

\[
G_{se} = \frac{100 - 5.5}{100 - 5.5} \times \frac{2.436}{1.032} = 2.645
\]

11.1.6 Using Equation 7, calculate the maximum theoretical specific gravity (\( G_{mm} \)) for each asphalt binder content.

Note 13: \( G_{se} \) is considered constant regardless of the asphalt binder content.

**Equation 7:**
\[
G_{mm} = \frac{100}{P_{sa} \cdot \frac{G_{se}}{G_b} + P_b}
\]

Where:
- \( G_{mm} \) = Calculated maximum theoretical specific gravity of the mix at each asphalt binder content \( P_b \).
- \( P_{sa} \) = Aggregate and mineral admixture content, percent by total weight of mix (100-\( P_b \)).
- \( P_b \) = Percent asphalt binder content, by total weight of mix, for each asphalt binder content.
- \( G_{se} \) = Effective specific gravity of the combined aggregate and mineral admixture.
- \( G_b \) = Specific gravity of the asphalt binder.

Example (for 4.5% asphalt binder content) (See Note 4):

\[
P_{sa} = 95.5
\]
\[
P_b = 4.5
\]
\[
G_{se} = 2.645
\]
\[
G_b = 1.032
\]

\[
G_{mm} = \frac{100}{95.5 \cdot \frac{2.645}{1.032} + 4.5} = 2.471
\]
12. DETERMINATION OF DESIGN PERCENT ASPHALT BINDER CONTENT

12.1 The design percent asphalt binder content is determined as follows in Subsections 12.2 through 12.3.

12.2 For each asphalt binder content used, calculate effective voids ($V_a$), percent absorbed asphalt ($P_{ba}$), voids in mineral aggregate ($V_{MA}$), percent effective asphalt binder content ($P_{be}$), and the ratio of the mix design composite gradation target for the No. 200 sieve, including mineral admixture, to the effective asphalt binder content (Dust/Bitumen Ratio) using the following equations.

12.2.1 Using Equation 8, calculate the effective voids ($V_a$). The calculated $G_{mm}$ values for the respective asphalt binder contents are used to determine the corresponding effective voids content of the compacted Marshall specimens at each asphalt binder content.

$$V_a = \left( \frac{G_{mm} - G_{mb}}{G_{mm}} \right) \times 100$$

Where: $V_a$ = Effective voids in the compacted mixture, percent of total volume.

$G_{mm}$ = Calculated maximum theoretical specific gravity of the mix at each asphalt binder content $P_b$.

$G_{mb}$ = Bulk specific gravity of compacted mixture specimens.

Example (for 4.5% asphalt binder content) (See Note 4):

$G_{mm} = 2.471$

$G_{mb} = 2.290$

$$V_a = \frac{2.471 - 2.290}{2.471} \times 100 = 7.3$$

12.2.2 Using Equation 9, calculate the percent absorbed asphalt ($P_{ba}$).

**Note 14**: The percent absorbed asphalt ($P_{ba}$) is the same for all asphalt binder contents.
Equation 9: 

\[ P_{ba} = \left( \frac{G_{se} - G_{sb}}{G_{sb} \times G_{se}} \right) \times G_b \times 100 \]

Where:
- \( P_{ba} \) = Absorbed asphalt, percent by total weight of mix.
- \( G_{se} \) = Effective specific gravity of the combined aggregate and mineral admixture.
- \( G_{sb} \) = Bulk oven dry specific gravity of the combined aggregate and mineral admixture.
- \( G_b \) = Specific gravity of the asphalt binder.

Example:
- \( G_{se} = 2.645 \)
- \( G_{sb} = 2.614 \)
- \( G_b = 1.032 \)

\[ P_{ba} = \frac{2.645 - 2.614}{2.614 \times 2.645} \times 1.032 \times 100 = 0.46 \]

12.2.3 Using Equation 10, calculate voids in mineral aggregate (VMA) of the compacted Marshall specimens at each asphalt binder content.

Equation 10: 

\[ VMA = 100 - \left( \frac{G_{mb} \times P_{sa}}{G_{sb}} \right) \]

Where:
- \( VMA \) = Voids in the mineral aggregate, percent of bulk volume.
- \( G_{mb} \) = Bulk specific gravity of compacted mixture specimens.
- \( P_{sa} \) = Aggregate and mineral admixture content, percent by total weight of mix (100-P\(_b\)).
- \( G_{sb} \) = Bulk oven dry specific gravity of the combined aggregate and mineral admixture.

Example (for 4.5% asphalt binder content) (See Note 4):

- \( G_{mb} = 2.290 \)
- \( P_{sa} = 95.5 \)
- \( G_{sb} = 2.614 \)
12.2.4 Using Equation 11, calculate the percent effective asphalt binder content ($P_{be}$) of the compacted Marshall specimens at each asphalt binder content.

**Equation 11:**

$$P_{be} = P_b - \left( \frac{P_{ba} \times P_{sa}}{100} \right)$$

Where:

- $P_{be}$ = Percent effective asphalt binder content of the mixture (free binder not absorbed).
- $P_b$ = Asphalt binder content, percent by total weight of mix.
- $P_{ba}$ = Absorbed asphalt, percent by total weight of mix.
- $P_{sa}$ = Aggregate and mineral admixture content, percent by total weight of mix (100-$P_b$).

Example (for 4.5% asphalt binder content) (See Note 4):

- $P_b = 4.5$
- $P_{ba} = 0.46$
- $P_{sa} = 95.5$

$$P_{be} = 4.5 - \frac{0.46 \times 95.5}{100} = 4.06$$

12.2.5 Using Equation 12, calculate the ratio of the mix design composite gradation target for the No. 200 sieve, including mineral admixture, to the effective asphalt binder content (Dust/Bitumen Ratio) of the compacted Marshall specimens at each asphalt binder content.

**Equation 12:**

$$\text{Dust} / \text{Bitumen Ratio} = \frac{\text{Mix Design}}{\text{Passing No. 200 (Including Mineral Admix.)}}$$

Where: $P_{be}$ = Percent effective asphalt binder content of the mixture (free binder not absorbed).
Example (for 4.5% asphalt binder content) (See Note 4):

\[
P_{be} = 4.06
\]

\[
\text{Dust/Bitumen Ratio} = \frac{5.0}{4.06} = 1.23
\]

12.3 If the specified value for effective voids (V_a) falls between the resultant V_a values determined for two of the asphalt binder contents used, straight-line interpolation shall be used to calculate the bulk specific gravity (G_{mb}) at each 0.1% asphalt binder content increment between the two corresponding measured values of G_{mb}. Using equations 7 and 8, determine the values for G_{mm} and V_a at each interpolated asphalt binder content. Select the asphalt binder content which results in V_a being as close as possible to the specified value.

12.3.1 Using equations 10, 11, and 12, calculate VMA, P_{be}, and the Dust/Bitumen Ratio at the selected asphalt binder content. Using straight-line interpolation, determine the values for stability and flow that correspond to the selected asphalt binder content. If the values for VMA, Dust/Bitumen Ratio, stability, and flow are within the limits of the specifications, the selected asphalt binder content shall be the design asphalt binder content.

12.3.2 On separate graphs (using the same scale for percent asphalt binder for each graph), plot the values for bulk specific gravity, effective voids, VMA, Dust/Bitumen Ratio, stability, and flow for each set of three specimens and at the design asphalt binder content, versus the percent asphalt binder. Draw a smooth “best fit” curve based on the plotted points, as shown in Figure 4. The graphs in Figure 4 are for illustrative purposes only. Straight-line interpolation, as specified in Subsections 12.3 and 12.3.1, is used for mix design calculations.

12.3.3 If it is not possible to obtain specification compliance within the range of asphalt binder contents used, a determination must be made to either redesign the mix (different aggregate gradation and/or source) or prepare additional specimens at other asphalt binder contents for testing and volumetric analyses.
13. IMMERSION COMPRESSION (IMC) TEST

13.1 The Immersion Compression Test shall be performed in accordance with ARIZ 802, modified as follows:

13.1.1 Prepare the asphaltic concrete sample including mineral admixture according to the procedures described in Section 5 and Subsections 8.2 through 8.5 at the design asphalt binder content.

13.1.2 Immediately after mixing, each batch of asphaltic concrete shall be thoroughly blended and spread according to the procedures described in ARIZ 416. The circular mass shall be cut into 4 equal pie-shaped segments. Take opposite segments for each individual sample and use up the entire batch.

13.1.3 Each individual sample shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The samples shall be oven-cured for 2 hours ± 10 minutes at 255 ± 5 ºF.

13.1.4 Compacted specimens shall be 4.000 ± 0.100 inches in height.

14. MIX DESIGN GRADATION TARGET VALUES

14.1 The target values for the aggregate composite without mineral admixture, and the aggregate-mineral admixture blend composite, for the asphaltic concrete mixture shall be expressed as percent passing particular sieve sizes as required by the specifications for the project. An example mix design composite, with and without mineral admixture, is shown in Figure 1.

15. REPORT

15.1 Report the test results and data obtained on the appropriate form. Liberal use of the remarks area to clarify and/or emphasize any element of the design is strongly recommended. Information required in the mix design report includes the items listed below:

15.1.1 Name and address of the testing organization and the signature and seal of the mix design engineer responsible for the mix design.
15.1.2 Aggregate:
- Source and identification
- Individual stockpile or bin gradations
- Aggregate blend proportions and composite gradation, with and without mineral admixture
- Fine and coarse aggregate specific gravities (Bulk Oven Dry, SSD, Apparent) and absorption
- Combined aggregate specific gravities (Bulk Oven Dry, SSD, Apparent)
- Combined aggregate absorption
- Aggregate quality
  - LA Abrasion
  - Sand Equivalent
  - Fractured Coarse Aggregate Particles (percentage with one fractured face and percentage with two fractured faces)
  - Uncompacted Void Content (when applicable)
  - Carbonates (when applicable)

15.1.3 Mineral Admixture:
- Type and source
- Percentage used
- Specific gravity

15.1.4 Combined Mineral Aggregate and Mineral Admixture
- Combined specific gravities [Bulk Oven Dry \( (G_{sb}) \), SSD, Apparent] and absorption of the combined aggregate and mineral admixture
- Effective specific gravity of the combined aggregate and mineral admixture \( (G_{se}) \)

15.1.5 Asphalt Binder:
- Source and grade
- Specific gravity (at 77 °F)
- Viscosity - Temperature Curve and the following:
  - Laboratory mixing temperature range and actual laboratory mixing temperature used.
  - Laboratory compaction temperature range and actual laboratory compaction temperature used.

15.1.6 Maximum theoretical specific gravity \( (G_{mm}) \) and density (pcf) at the asphalt binder content at which the Rice test was performed \( (Rice \ P_b) \)
15.1.7 Mixture Compaction Trials:
- Percent asphalt binder content \( (P_b) \), by weight of the bituminous mixture
- Aggregate and mineral admixture content \( (P_{sa}) \)
- Calculated maximum theoretical specific gravity \( (G_{mm}) \) and density \( (pcf) \)
- Bulk specific gravity \( (G_{mb}) \) and bulk density \( (pcf) \) of Marshall specimens
- Percent effective voids \( (V_a) \)
- Percent voids in mineral aggregate \( (VMA) \)
- Dust/Bitumen Ratio
- Percent absorbed asphalt \( (P_{ba}) \)
- Percent effective asphalt binder content \( (P_{be}) \)
- Marshall stability (nearest 10 pounds)
- Marshall flow (0.01 inch)

15.1.8 Plots of the following properties versus percent asphalt binder content (See examples in Figure 4):
- Percent effective voids \( (V_a) \)
- Percent voids in mineral aggregate \( (VMA) \)
- Dust/Bitumen Ratio
- Bulk specific gravity \( (G_{mb}) \)
- Marshall stability
- Marshall flow

15.1.9 Mixture Properties at Design Percent Asphalt Binder Content:
- Percent asphalt binder content \( (P_b) \), by weight of the bituminous mixture
- Aggregate and mineral admixture content \( (P_{sa}) \)
- Calculated maximum theoretical specific gravity \( (G_{mm}) \) and density \( (pcf) \)
- Bulk specific gravity \( (G_{mb}) \) and bulk density \( (pcf) \) of Marshall specimens
- Percent effective voids \( (V_a) \)
- Percent voids in mineral aggregate \( (VMA) \)
- Dust/Bitumen Ratio
- Percent absorbed asphalt \( (P_{ba}) \)
- Percent effective asphalt binder content \( (P_{be}) \)
- Marshall stability (nearest 10 pounds)
- Marshall flow (0.01 inch)
- Immersion Compression wet strength (psi)
- Immersion Compression dry strength (psi)
- Index of retained strength
15.1.10 Mix Design Summary. The mix design summary shall contain the information shown in Figure 5. [An example Mix Design Summary for an Asphaltic Concrete Marshall Mix Design.]

### Example Mix Design Composite
**(With and Without Mineral Admixture)**

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* 1.0 percent admixture, by weight of dry mineral aggregate, is used.

**FIGURE 1**
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<td>---</td>
<td>68</td>
<td>34</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>5.800</td>
<td>10.000</td>
<td>---</td>
<td>---</td>
<td>186</td>
<td>93</td>
<td>66</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Washed Crushed Fines</td>
<td>1/4&quot;</td>
<td>0.027</td>
<td>10.027</td>
<td>11</td>
<td>85</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>0.297</td>
<td>10.324</td>
<td>14</td>
<td>114</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>5.913</td>
<td>16.237</td>
<td>---</td>
<td>---</td>
<td>306</td>
<td>153</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>20.783</td>
<td>37.000</td>
<td>---</td>
<td>---</td>
<td>727</td>
<td>364</td>
<td>304</td>
<td>---</td>
<td>---</td>
</tr>
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<td></td>
<td>1/4&quot;</td>
<td>4.530</td>
<td>42.700</td>
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<td>972</td>
<td>---</td>
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<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>2.370</td>
<td>45.070</td>
<td>113</td>
<td>904</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>5.670</td>
<td>50.740</td>
<td>---</td>
<td>---</td>
<td>842</td>
<td>421</td>
<td>---</td>
<td>---</td>
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</tr>
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<td></td>
<td>#8</td>
<td>16.280</td>
<td>57.000</td>
<td>---</td>
<td>---</td>
<td>1173</td>
<td>585</td>
<td>489</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>1/2&quot; Aggregate</td>
<td>3/4&quot;</td>
<td>5.775</td>
<td>72.775</td>
<td>184</td>
<td>1469</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>1/2&quot;</td>
<td>16.236</td>
<td>89.011</td>
<td>382</td>
<td>3057</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>2919</td>
</tr>
<tr>
<td></td>
<td>3/8&quot;</td>
<td>5.808</td>
<td>94.819</td>
<td>453</td>
<td>3625</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>5000</td>
</tr>
<tr>
<td></td>
<td>1/4&quot;</td>
<td>3.432</td>
<td>98.251</td>
<td>495</td>
<td>3961</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>0.396</td>
<td>98.647</td>
<td>500</td>
<td>4000</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.429</td>
<td>99.076</td>
<td>---</td>
<td>---</td>
<td>1181</td>
<td>580</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.924</td>
<td>100.000</td>
<td>---</td>
<td>---</td>
<td>1200</td>
<td>800</td>
<td>500</td>
<td>---</td>
<td>---</td>
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</tbody>
</table>
## Example Weigh Up Sheet #2

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SIEVE</th>
<th>INDIV. PERCENT RETAINED</th>
<th>ACCUM. PERCENT RETAINED</th>
<th>RICE (3000 grams)</th>
<th>MARSHALL (3300 grams)</th>
<th>IMMERSION COMPRESSION (3400 grams)</th>
<th>MINUS NO. 8 MAKE-UP MATERIAL (500 grams)</th>
<th>ACCUMULATIVE WEIGHT (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crusher Fines</td>
<td>#4</td>
<td>0.840</td>
<td>0.840</td>
<td>25</td>
<td>28</td>
<td>29</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>3.360</td>
<td>4.200</td>
<td>126</td>
<td>139</td>
<td>143</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>5.800</td>
<td>10.000</td>
<td>300</td>
<td>330</td>
<td>340</td>
<td>66</td>
<td></td>
</tr>
<tr>
<td>Washed Crusher Fines</td>
<td>1/4&quot;</td>
<td>0.027</td>
<td>10.027</td>
<td>301</td>
<td>331</td>
<td>341</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>0.297</td>
<td>10.324</td>
<td>310</td>
<td>341</td>
<td>351</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>5.913</td>
<td>16.237</td>
<td>487</td>
<td>536</td>
<td>552</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>20.763</td>
<td>37.000</td>
<td>1110</td>
<td>1221</td>
<td>1258</td>
<td>304</td>
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</tr>
<tr>
<td>3/8&quot; Aggregate</td>
<td>3/8&quot;</td>
<td>1.170</td>
<td>38.170</td>
<td>1145</td>
<td>1260</td>
<td>1298</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1/4&quot;</td>
<td>4.530</td>
<td>42.700</td>
<td>1281</td>
<td>1409</td>
<td>1452</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>2.370</td>
<td>45.070</td>
<td>1352</td>
<td>1487</td>
<td>1532</td>
<td>---</td>
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<tr>
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<td>#8</td>
<td>5.670</td>
<td>50.740</td>
<td>1522</td>
<td>1674</td>
<td>1725</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>16.260</td>
<td>67.000</td>
<td>2010</td>
<td>2211</td>
<td>2278</td>
<td>489</td>
<td></td>
</tr>
<tr>
<td>3/4&quot; Aggregate</td>
<td>3/4&quot;</td>
<td>5.775</td>
<td>72.775</td>
<td>2183</td>
<td>2402</td>
<td>2474</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1/2&quot;</td>
<td>16.236</td>
<td>89.011</td>
<td>2670</td>
<td>2937</td>
<td>3026</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3/8&quot;</td>
<td>5.808</td>
<td>94.819</td>
<td>2845</td>
<td>3129</td>
<td>3224</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1/4&quot;</td>
<td>3.432</td>
<td>98.251</td>
<td>2948</td>
<td>3242</td>
<td>3341</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>0.396</td>
<td>98.647</td>
<td>2959</td>
<td>3255</td>
<td>3354</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.429</td>
<td>99.076</td>
<td>2972</td>
<td>3270</td>
<td>3369</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>0.924</td>
<td>100.000</td>
<td>3000</td>
<td>3300</td>
<td>3400</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>Weight of Mineral Admixture (Type II Cement)</td>
<td>[1.0%, by weight of Dry Mineral Aggregate]</td>
<td>30</td>
<td>33</td>
<td>34</td>
<td>---</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total weight of Mineral Aggregate and Mineral Admixture</td>
<td>3030</td>
<td>3333</td>
<td>3434</td>
<td>---</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

FIGURE 3
Example Plots of Effective Voids, VMA, Dust/Bitumen Ratio, Bulk Specific Gravity, Marshall Stability, and Marshall Flow versus Percent Asphalt Binder Content

The above graphs are for illustrative purposes only. Straight-line interpolation, as specified in Subsections 12.3 and 12.3.1, is used for mix design calculations.

FIGURE 4
## Asphalitic Concrete Marshall Mix Design
### Example Mix Design Summary

**Date:** February 22, 2013  
**Contractor:** Best Paving Contractors, Inc.  
**Mix Designation:** ADOT 416 (3/4 Inch Special Mix)  
**ADOT Project Number:** STP-888-Z(888)A  
**ADOT TRACS Number:** H888801C  
**Mix Design Laboratory:** XYZ Testing Laboratories  
**Mix Design Engineer:** Joe Designer  
**Project Name:** Anywhere - Everywhere  
**Grade/Specific Gravity of Asphalt Binder:** PG 70-10/1.032

### COMPOSITE GRADATION:

<table>
<thead>
<tr>
<th>Mineral Aggregate</th>
<th>Percent Used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crusher Fines</td>
<td>10</td>
</tr>
<tr>
<td>Washed Crusher Fines</td>
<td>27</td>
</tr>
<tr>
<td>3/8&quot; Aggregate</td>
<td>30</td>
</tr>
<tr>
<td>3/4&quot; Aggregate</td>
<td>33</td>
</tr>
<tr>
<td>Mineral Admixture</td>
<td>1.0</td>
</tr>
</tbody>
</table>

### DESIGN DATA:

<table>
<thead>
<tr>
<th>% Asphalt Binder</th>
<th>4.5</th>
<th>5.0</th>
<th>5.2</th>
<th>5.5</th>
<th>Spec.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Specific Gravity</td>
<td>2.290</td>
<td>2.309</td>
<td>2.311</td>
<td>2.314</td>
<td></td>
</tr>
<tr>
<td>Bulk Density (lb./cu. ft.)</td>
<td>142.7</td>
<td>143.9</td>
<td>144.0</td>
<td>144.2</td>
<td></td>
</tr>
<tr>
<td>Stability (lb.)</td>
<td>4120</td>
<td>4550</td>
<td>4530</td>
<td>4500</td>
<td>2000 Min.</td>
</tr>
<tr>
<td>Flow (0.01 inch)</td>
<td>13</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>8 - 16</td>
</tr>
<tr>
<td>Effective Void (%)</td>
<td>7.3</td>
<td>5.9</td>
<td>5.5</td>
<td>5.0</td>
<td>5.5 ± 0.2</td>
</tr>
<tr>
<td>VMA (%)</td>
<td>16.3</td>
<td>16.1</td>
<td>16.2</td>
<td>16.4</td>
<td>15.0 - 18.0</td>
</tr>
<tr>
<td>Effective Asphalt, Total Mix (%)</td>
<td>4.06</td>
<td>4.56</td>
<td>4.76</td>
<td>5.06</td>
<td></td>
</tr>
<tr>
<td>Dust/Bitumen Ratio</td>
<td>1.23</td>
<td>1.10</td>
<td>1.05</td>
<td>0.99</td>
<td>0.6 - 1.2</td>
</tr>
<tr>
<td>Absorbed Asphalt (%)</td>
<td>0.46</td>
<td>0 - 1.0</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Max. Theoretical Specific Gravity at 5.2% Asphalt Binder = 2.446  
Max. Theoretical Density at 5.2% Asphalt Binder = 152.4 lb/cu. ft. |

### AGGREGATE PROPERTIES:

<table>
<thead>
<tr>
<th>Test Characteristic</th>
<th>Fine</th>
<th>Coarse</th>
<th>Combined</th>
<th>Spec.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk O.D. Specific Gravity</td>
<td>2.593</td>
<td>2.634</td>
<td>2.610</td>
<td>2.350 - 2.850</td>
</tr>
<tr>
<td>SSD Specific Gravity</td>
<td>2.626</td>
<td>2.655</td>
<td>2.638</td>
<td>-----</td>
</tr>
<tr>
<td>Apparent Specific Gravity</td>
<td>2.682</td>
<td>2.690</td>
<td>2.685</td>
<td>-----</td>
</tr>
<tr>
<td>Absorption</td>
<td>1.28</td>
<td>0.79</td>
<td>1.08</td>
<td>0 - 2.5%</td>
</tr>
<tr>
<td>Sand Equivalent</td>
<td>75</td>
<td>Min. 55</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Uncompacted Void Content</td>
<td>47.4</td>
<td>Min. 45.0%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carbonates</td>
<td>N/A</td>
<td>-----</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fractured Coarse Aggregate Particles</td>
<td>-----</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At least one fractured face</td>
<td>100</td>
<td>Min. 92%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>At least two fractured faces</td>
<td>95</td>
<td>Min. 85%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abrasion (Type B)</td>
<td>-----</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 Rev., % Loss</td>
<td>4</td>
<td>9% Max.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>500 Rev., % Loss</td>
<td>22</td>
<td>40% Max.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### RECOMMENDED ASPHALT BINDER CONTENT:

- **(By weight of total mix)**  
- **5.2%**

### ADDITIONAL DATA / INFORMATION:

- Source of Mineral Aggregate: Excellent Aggregate Materials, Inc.  
- Source of Asphalt Binder: Superb Asphalt Suppliers  
- Mineral Admixture Type/Specific Gravity: Type II Cement/3.14  
- Laboratory Mixing Temperature Range: 320 °F to 331 °F  
- Laboratory Compaction Temperature Range: 298 °F to 307 °F  
- Actual Laboratory Mixing Temperature Used: 325 °F  
- Actual Laboratory Compaction Temperature Used: 300 °F

### REMARKS:

**FIGURE 5**
DESIGN OF EXPOSED AGGREGATE SEAL COATS

(An Arizona Method)

SCOPE

1. (a) This method describes the procedure for calculating design quantities of materials for exposed aggregate seal coats utilizing cover aggregates and emulsified asphalt. The resultant quantities are to be adjusted for field conditions, as necessary.

   (b) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (c) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

DATA

2. The following values are to be determined by the method indicated:

   H = Average least dimension of cover aggregate (Median Aggregate size modified for Flakiness Index - Arizona Test Method 233).

   G = Bulk Specific Gravity (Arizona Test Method 210 - Saturated surface dry basis).

   R = Residual bitumen of emulsified asphalt, expressed as a decimal (AASHTO T 59).
T = Traffic factor from the following:

When the ADT is less than or equal to 2000,
\[ T = 0.85 - [0.0001 \times (\text{ADT})] \]

When the ADT is more than 2000, \( T = 0.65 \)

NOTE: Traffic must be considered by lanes - for instance on a divided highway, the passing lane and distress lanes should have a higher traffic factor than the travel lane.

E = Wastage factor due to cover aggregate lost to whip off by traffic and to variation in spread (1 + Estimated Waste, for example, 1.05 for 5% wastage).

V = Voids in Aggregate by the following:

Metric:
\[ V = 1 - \frac{W_L}{(998) \times (G)} \]

Where: \( W_L = \) Loose unit weight of aggregate in kg/m³ by AASHTO T 19 (Shoveling Procedure).

English:
\[ V = 1 - \frac{W_L}{(62.3) \times (G)} \]

Where: \( W_L = \) Loose unit weight of aggregate in lbs./cu. ft. by AASHTO T 19 (Shoveling Procedure).
S = Surface Texture Correction from below:

<table>
<thead>
<tr>
<th>Surface Texture</th>
<th>Surface Texture Correction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>liters/square meter</td>
</tr>
<tr>
<td></td>
<td>gals./sq. yd.</td>
</tr>
<tr>
<td>Very Porous</td>
<td>0.41</td>
</tr>
<tr>
<td>Rough and Porous</td>
<td>0.27</td>
</tr>
<tr>
<td>Slightly Rough, Black</td>
<td>0.14</td>
</tr>
<tr>
<td>Black and Smooth (Hard, no aggregate</td>
<td>0.00</td>
</tr>
<tr>
<td>embedment predicted)</td>
<td></td>
</tr>
<tr>
<td>Soft and Rich (Aggregate embedment</td>
<td>-0.05* to -0.23*</td>
</tr>
<tr>
<td>predicted)</td>
<td>-0.01* to -0.05*</td>
</tr>
<tr>
<td></td>
<td>* Depending upon aggregate</td>
</tr>
<tr>
<td></td>
<td>embedment estimation.</td>
</tr>
</tbody>
</table>

CALCULATIONS

3. The required amount of cover material and bituminous material shall be calculated by the following:

(a) Cover Material Requirement:

Metric:

For 100% pass 9.5 mm sieve,

\[ C = [0.6163 \times (G) \times (H) \times (E)] \times (2 - V) \]

For 100% pass 12.5 mm sieve,

\[ C = [0.5078 \times (G) \times (H) \times (E)] \times (2 - V) \]

Where: \( C = \) Cover Material in \( \text{kg/m}^2 \)

Convert cover material from \( \text{kg/m}^2 \) to \( \text{m}^3/\text{m}^2 \)

by the following equation:

\[ \frac{C(\text{in kg/m}^2)}{W_c(\text{in kg/m}^3)} = C(\text{in m}^3/\text{m}^2) \]
English:

For 100% pass 3/8" sieve,
\[ C = [28.4 \times (G) \times (H) \times (E)] \times (2 - V) \]

For 100% pass 1/2" sieve,
\[ C = [23.4 \times (G) \times (H) \times (E)] \times (2 - V) \]

Where: \( C \) = Cover Material in lbs./sq. yd

Convert cover material from lbs./sq. yd. to cubic yards per square yard by the following equation:

\[
\frac{C \text{ (in lbs./sq. yd.)}}{[W_L \text{ (in lbs./cu. ft.)}] \times 27}
\]

(b) Bituminous Material Requirement:

Metric:
\[ B = \frac{0.3622 \times (H) \times (T)}{R} + S \]

Where: \( B \) = Bituminous Material in L/m²

English:
\[ B = \frac{2 \times (H) \times (T)}{R} + S \]

Where: \( B \) = Bituminous Material in gals./sq. yd.

EXAMPLE

4. The following example will serve to illustrate the calculations:

(a) Data:

\[
\begin{align*}
H &= 5.50 \text{ mm} \\
G &= 2.630 \\
R &= 0.65 \\
T \text{ (for 1500 ADT)} &= 0.85 - [0.0001 \times (1500)] = 0.70 \\
E &= 1.06
\end{align*}
\]
V = 0.46 (W_L = 1418 kg/m^3) 
S = Rough and Porous = 0.27 L/m^2 

(b) Cover Material Requirement:

For 100% pass 12.5 mm sieve,

\[ C = [0.5078 (G) (H) (E)] (2 - V) \]
\[ = [0.5078 (2.630) (5.50) (1.06)] (2 - 0.46) \]
\[ = 11.99 \text{ kg/m}^2 \]

Convert 11.99 kg/m^2 to m^3/m^2 by:

\[ \frac{11.99 \text{ kg/m}^2}{1418 \text{ kg/m}^3} = 0.008 \text{ m}^3/\text{m}^2 \]

(c) Bituminous Material Requirement:

\[ B = \frac{0.3622 (H) (T)}{R} + S \]
\[ = \frac{0.3622 (5.50) (0.70)}{0.65} + 0.27 \]
\[ = 2.42 \text{ liters per square meter} \]

REPORT

5. Report the following:

(a) The amount of cover material required, in kg/m^2 (lbs./sq. yd.), to the nearest 0.01 kg/m^2 (0.1 lbs./sq. yd.).

(b) The amount of cover material required, in m^3/m^2 (cubic yards per square yard), to the nearest 0.001 m^3/m^2 (0.001 cubic yards per square yard).

(c) The amount of bituminous material required, to the nearest 0.01 L/m^2 (0.01 gals./sq. yd.).
METHOD OF TEST FOR DETERMINING THE QUANTITY OF ASPHALT REJUVENATING AGENT REQUIRED FOR AN ASPHALTIC PAVEMENT

(A Modification of California Test No. 345)

SCOPE

1. (a) This method of test describes the procedure for determining the quantity of asphalt rejuvenating agent an asphaltic pavement will absorb in a specified period of time.

   NOTE: This test is used primarily for maintenance purposes.

   (b) This test method may involve hazardous material, operations, or equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of any regulatory limitations prior to use.

   (c) See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

   (d) Metric (SI) units and values are shown in this test method with English units and values following in parentheses. Values given for metric and English units may be numerically equivalent (soft converted) for the associated units, or they may be given as rounded or rationalized values (hard converted). Either the metric or English units along with their corresponding values shall be used in accordance with applicable specifications. See Appendix A2 of the Materials Testing Manual for additional information on the metric system.

APPARATUS

2. Apparatus for this test procedure shall consist of the following:
(a) 1 - Aluminum template 152 mm (6") diameter with handle.

(b) 1 - Caulking gun with a 127 mm (5") piece of 6.35 mm (1/4") copper tubing with cap.

(c) 1 - 25 mL graduated cylinder (plastic or glass).

(d) 1 - 127 mm (5") inch trowel.

(e) 2 - small stiff bristle brushes.

(f) 2 - 3.8 liter (1 gallon) friction top cans. One for new medium weight chassis grease. One for used grease.

(g) 1 - 203 mm (8") spatula.

(h) 2 - pieces of yellow lumber crayon.

(i) 1 - stop watch with 60 second dial.

MATERIAL

3. The following materials are required:

(a) Medium weight chassis grease.

(b) A liter (quart) can containing a dilution of 2 parts rejuvenating agent and one part distilled water.

PROCEDURE

4. (a) With lumber crayon and template, draw a 152 mm (6") diameter circle on the pavement.

(b) Extrude grease from the caulking gun on the circle. The diameter of the grease should be about 6 mm (1/4").

(c) Run finger around the outside of the grease ring, pushing a small amount of grease into the pavement. This will form a sealed reservoir for the asphalt rejuvenating agent (2:1 dilution).
(d) Measure 8.3 mL of asphalt rejuvenating agent, 2:1 dilution, in the graduate and pour into grease ring. Start stop watch and quickly brush agent uniformly over the area within the grease ring.

(e) Record the time required for the rejuvenating agent to completely penetrate the surface. Complete penetration is generally indicated by a loss in color of the rejuvenating agent, giving a dull dark appearance to the surface.

NOTE: Loss of color can also occur when evaporation of the water takes place leaving a thin film of the residue on the surface of the pavement giving a shiny appearance. In this case penetration has not occurred.

(f) If 8.3 mL is absorbed within a 15 minute interval, make a new grease ring and repeat test with additional testing solution in increments of 8.3 mL until time of penetration just exceeds 15 minutes.

(g) If 8.3 mL is not absorbed within the 15 minute period, repeat the test using 4.1 mL.

NOTE: 4.1 mL is equivalent to 0.23 L/m² (0.05 gal./sq. yd.) spread rate.
8.3 mL is equivalent to 0.45 L/m² (0.10 gal./sq. yd.) spread rate.
16.6 mL is equivalent to 0.91 L/m² (0.20 gal./sq. yd.) spread rate.

(h) If an application rate of 0.91 L/m² (0.20 gal./sq. yd.) [16.6 mL] is exceeded, the effect of rejuvenating agent on the properties of the asphaltic pavement (for example, stability/flow, voids, etc.) shall be evaluated.

(i) Graduate can be easily cleaned with water.

(j) Pick up grease with trowel and place in 3.8 liter (1 gallon) can for used grease. Do not mix used grease with new grease.
PROCEDURE FOR DETERMINING PAVEMENT REQUIREMENTS

5. The following procedure is recommended for obtaining the average absorption rate or quantity of rejuvenating agent required for a given section of highway.
   (a) In any travel lane, determine the absorption at 8 meter (25 foot) intervals in the outer wheel path, inner wheel path, and between wheel paths, respectively. The three readings should be averaged to obtain the reading for the test area.
   (b) This procedure should be repeated at intervals of approximately 600 meters (2000 feet).

REFERENCE

6. "Method of Test for Determining Quantity and Rate of Absorption of Reclamite into an Asphalt Pavement" by Golden Bear Oil Company, Bakersfield, California.
EVALUATION OF PAVEMENT SMOOTHNESS

(An Arizona Method)

SCOPE

1. This test method describes the procedure for determining the smoothness of pavements. Profile measurements are taken on designated area(s) of the pavement. Those measurements are analyzed and reported as International Roughness Index (IRI) values, inches per mile (in/mi).

APPARATUS

2. The equipment used to perform profile testing shall be a General Motors Research (GMR)-type Inertial Profiler, meeting the requirements of ASTM E 950.

METHOD OF MEASURING

3. The test vehicle is driven in the wheel paths of each lane a sufficient number of times to collect two valid individual sets of profile measurements. The speed at which profile measurements are taken shall be a constant speed (± 2 mph), between 30 and 65 mph.

CALCULATIONS

4. (a) Profile measurements taken using the GMR-type Inertial Profiler will be reduced to IRI smoothness values, in accordance with the requirements of ASTM E 1926. IRI smoothness values will be recorded to the nearest 0.1 in/mi for each 0.1 mile increment.

(b) For each individual set of IRI smoothness values, the average value will be determined.

(c) The set of IRI smoothness values with the lowest average value will be used for acceptance purposes.
(d) If necessary, 0.1 mile increment IRI values used for acceptance purposes shall be adjusted to account for any exception areas required by the contract documents.

REPORT

5. A report will be prepared showing:

(a) The date of test.

(b) The name of the test operator(s).

(c) Identification of test vehicle.

(d) Speed of test vehicle during testing.

(e) Project number, if applicable.

(f) The individual IRI value, adjusted as necessary per paragraph 4(d), and recorded to the nearest 0.1 in/mi for each 0.1 lane-mile increment (Actual Smoothness Value “AS”).

(g) The location of each 0.1 lane-mile increment for the corresponding individual IRI value.
1. SCOPE

1.1 This method is used to design Asphaltic Concrete (Asphalt-Rubber) [AR-AC] mixes using 4-inch diameter Marshall apparatus.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 A listing of subsequent Sections and Figures in this procedure is given below:

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<th>Title</th>
<th>Page #</th>
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<td>Figure 5</td>
<td>Example Mix Design Summary</td>
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</table>
2. TEST METHODS AND APPARATUS

2.1 This test method is used in conjunction with the test methods listed below. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. The required apparatus is shown in the individual test methods, as appropriate.

2.2 Arizona Test Methods:

- ARIZ 201 Sieving of Coarse and Fine Graded Soils and Aggregates
- ARIZ 205 Composite Grading
- ARIZ 210 Specific Gravity and Absorption of Coarse Aggregate
- ARIZ 211 Specific Gravity and Absorption of Fine Aggregate
- ARIZ 212 Percentage of Fractured Coarse Aggregate Particles
- ARIZ 238 Percent Carbonates in Aggregate
- ARIZ 247 Particle Shape and Texture of Fine Aggregate Using Uncompacted Void Content
- ARIZ 248 Alternate Procedures for Sieving of Coarse and Fine Graded Soils and Aggregates
- ARIZ 251 Combined Coarse and Fine Aggregate Specific Gravity and Absorption
- ARIZ 410 Compaction and Testing of Bituminous Mixtures Utilizing Four Inch Marshall Apparatus
- ARIZ 415 Bulk Specific Gravity and Bulk Density of Compacted Bituminous Mixtures
- ARIZ 416 Preparing and Splitting Field Samples of Bituminous Mixtures for Testing
- ARIZ 806 Maximum Theoretical Specific Gravity of Laboratory Prepared Bituminous Mixtures (Rice Test)

2.1.2 AASHTO Test Methods:

- AASHTO T 96 Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine
- AASHTO T 176 Plastic Fines in Graded Aggregates and Soils by Use of the Sand Equivalent Test
- AASHTO T 228 Specific Gravity of Semi-Solid Bituminous Materials

**Note 1:** Testing by AASHTO T 228 shall be performed at 77 °F.
3. MATERIALS

3.1 Mineral Aggregate - The mineral aggregate used in the design shall be produced material from the source(s) for the project. Use of natural sand is not permitted in AR-AC mixtures.

3.1.1 The composited gradation of the aggregate without admixture, and the composited gradation of the aggregate-mineral admixture blend shall comply with the grading limits of the specifications.

3.1.2 The composited mineral aggregate shall conform to the requirements of the specifications for Sand Equivalent (AASHTO T 176), Combined Bulk Oven Dry Specific Gravity (ARIZ 251), Combined Water Absorption (ARIZ 251), Fractured Coarse Aggregate Particles (ARIZ 212), Uncompacted Void Content (ARIZ 247), and Percent Carbonates (ARIZ 238) when applicable.

3.1.3 Mineral aggregate from each source shall be tested separately for compliance to the project requirements for Abrasion (AASHTO T 96).

3.2 Bituminous Material - The bituminous material used in the design shall be asphalt-rubber material [hereinafter Crumb Rubber Asphalt (CRA)], conforming to the requirements of Section 1009 of the specifications, which is to be used in the production of the AR-AC. The specific gravity of the CRA and of the asphalt cement used in the CRA shall be determined in accordance with AASHTO T 228 (at 77 °F).

3.3 Mineral Admixture - Mineral admixture is required. The mineral admixture used in the design shall be the same type of material to be used in production of the AR-AC. The mineral admixture shall conform to the requirements of the specifications.

3.4 Batch Plants - Mix designs developed for asphaltic concrete which is to be produced in a batch plant shall be prepared in accordance with this procedure, modified as follows:

3.4.1 Aggregate samples will be obtained from the hot bins for all testing except Sand Equivalent. Testing for Sand Equivalent will be performed on aggregate stockpile samples proportioned and composited to meet the mix design composite without admixture. Any method may be used to combine the aggregate stockpile samples for Sand Equivalent testing as long as the resultant gradation is representative of the mix design composite gradation without admixture.
3.4.2 The mix design shall provide component percentages and composite gradations for both stockpile and hot bin materials.

4. DETERMINATION OF COMPOSITE GRADATION

4.1 The gradation of the aggregate from each individual component stockpile or bin shall be determined in accordance with ARIZ 248 using washed sieve analysis Alternate #3, Alternate #4, or Alternate #5. For Alternate #5, washing of the coarse aggregate may be performed on the composite Plus No. 4 material and applied to the composite percent pass the minus No. 200 determined from the unwashed coarse sieving and washed fine sieving of the individual stockpiles.

4.2 The composite gradation of the mineral aggregate is determined using desired percentages of each component based on washed sieve analysis. Mix designs may be developed based on bin or stockpile material, as appropriate for the respective mix production facility to be used.

4.3 The mineral aggregate composite shall be determined in accordance with ARIZ 205, paragraph 2(e).

4.4 The aggregate-mineral admixture blend composite is determined by adjusting the mineral aggregate composite (percent passing) for mineral admixture by performing the calculation in Equation 1 for each sieve:

\[
\text{Equation 1:} \quad \left( \frac{\% \text{ passing each sieve in the aggregate composite}}{(100) + (\% \text{ Mineral Admixture})} \right) = \text{[Adjusted for Mineral Admixture]}
\]

4.5 The composited gradation of the aggregate and the composited gradation of the aggregate-mineral admixture blend shall be shown on the design report, along with the percentage of each material.

5. PREPARING AGGREGATE/MINERAL ADMIXTURE SAMPLES FOR MIX DESIGN TESTING

5.1 Based on the stockpile or bin composite aggregate gradation, the aggregate samples needed for mix design tests are prepared as follows.
5.1.1 Dry the mineral aggregate from each individual stockpile at a temperature not exceeding any temperature restrictions specified in subsequent test procedures. Drying shall be performed until no further weight loss is obtained from continued drying.

5.1.2 Representative samples of aggregate material which is retained on the individual No. 8 and larger sieve sizes and the minus No. 8 material from each stockpile or bin are used to prepare the samples for mix design testing.

5.1.3 Table 1 shows the aggregate sample sizes, the number of samples required for each test listed, and which samples include mineral admixture. The aggregate weight shown for Maximum Theoretical Specific Gravity will provide three Rice test specimens and the amount shown for Density-Stability/Flow will produce three Marshall specimens. Example weigh-up sheets are shown in Figures 2 and 3.

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Sample Size</th>
<th>Number of Samples</th>
</tr>
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<tbody>
<tr>
<td>Fractured Coarse Aggregate Particles (ARIZ 212)</td>
<td>Required grams of Mineral Aggregate as determined by test method [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Percent Carbonates (ARIZ 238) (When required)</td>
<td>300 grams of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Abrasion (AASHTO T 96)</td>
<td>5000 grams of Mineral Aggregate as per test method for grading type [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Fine Aggregate Specific Gravity/ Absorption (ARIZ 211)</td>
<td>1200 grams of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Coarse Aggregate Specific Gravity/Absorption (ARIZ 210)</td>
<td>Required grams of Mineral Aggregate as determined by the Nominal Maximum Aggregate Size [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Sand Equivalent (AASHTO T 176)</td>
<td>500 to 750 grams of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td>Table 1 (Continued)</td>
<td></td>
<td></td>
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<tr>
<td>---------------------</td>
<td></td>
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</tr>
<tr>
<td><strong>Uncompacted Void Content (ARIZ 247)</strong></td>
<td>Minimum 500 grams of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td><strong>Minus No. 8 Make-Up Material</strong></td>
<td>An adequate amount (normally 500 grams) of Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
</tr>
<tr>
<td><strong>Marshall Density-Stability/Flow (ARIZ 415 and ARIZ 410, as modified in Sections 8 and 9 respectively)</strong></td>
<td>3000 grams of Mineral Aggregate (See Note 2) [Plus 30 grams of mineral admixture]</td>
<td>3 (See Note 3) [Each sample yields 1 set of 3 Marshall Specimens]</td>
</tr>
<tr>
<td><strong>Maximum Theoretical Specific Gravity (Rice Test) (ARIZ 806, as modified in Section 10)</strong></td>
<td>3000 grams of Mineral Aggregate [Plus 30 grams of mineral admixture]</td>
<td>1 [Yields 3 test specimens]</td>
</tr>
</tbody>
</table>

**Note 2:** Generally 3000 grams of mineral aggregate will provide Marshall Density-Stability/Flow specimens of acceptable heights, but adjustments may be necessary in some cases. Use Equation 2 to adjust the weight of aggregate as necessary to conform to specimen height requirements of 2.500 ± 0.200 inches.

**Equation 2:** Adjusted Weight of Aggregate = \( \left( \frac{\text{Combined Bulk O.D.}}{\text{Agg. Specific Gravity}} \right) \times 3000 \)

**Note 3:** Requires one sample for each CRA binder content to be tested (minimum of three CRA binder contents, with three Marshall specimens at each CRA binder content).

5.1.4 After the aggregate samples for the Marshall and Rice specimens have been composited, add 1% mineral admixture, by dry weight of the aggregate, and mix thoroughly. Add 3% water, by dry weight of the aggregate, to each sample and mix thoroughly to wet the mineral admixture and aggregate surfaces.

5.1.5 Testing utilizing the prepared virgin aggregate-mineral admixture samples for Marshall and Rice will be performed as specified in Sections 7-9 and 10, respectively.
6. AGGREGATE SPECIFIC GRAVITIES AND ABSORPTION

6.1 Determine the Bulk Oven Dry, S.S.D., Apparent Specific Gravities and Absorption for the fine aggregate (minus No. 4) and the coarse aggregate (plus No. 4) in accordance with ARIZ 211 and ARIZ 210, respectively.

6.2 Determine the combined bulk oven dry specific gravity of the fine aggregate and the coarse aggregate without mineral admixture and also the combined water absorption of the fine aggregate and the coarse aggregate without mineral admixture, in accordance with ARIZ 251.

6.2.1 The combined aggregate bulk oven dry specific gravity without mineral admixture and the combined aggregate water absorption without mineral admixture are used only to determine compliance with specification requirements.

6.3 Using Equation 3, calculate the Combined Bulk Oven Dry (Gsb), S.S.D., and Apparent Specific Gravities of the aggregate-mineral admixture blend.

\[
\frac{P_f + P_c + P_{\text{admix}}}{G_f + G_c + G_{\text{admix}}} = P_{f} + P_{c} + P_{\text{admix}}
\]

Where:
- \( P_f, P_c \) = Weight percent of fine aggregate (minus No. 4) and coarse aggregate (plus No. 4) respectively. Determined from the aggregate composite without mineral admixture.
- \( P_{\text{admix}} \) = Percent mineral admixture by weight of the aggregate.
- \( P_f + P_c = 100 \)
- \( P_f + P_c + P_{\text{admix}} = 100 + \% \) Mineral Admixture
- \( G_f, G_c \) = Bulk Oven Dry, S.S.D., or Apparent specific gravity of the fine aggregate and the coarse aggregate respectively.
- \( G_{\text{admix}} \) = Specific gravity of the mineral admixture.

- Type I or II Cement = 3.14
- Type IP Cement = 3.00
- Hydrated Lime = 2.20
Example [for combined Bulk Oven Dry Specific Gravity (Gsb)]:

\[
P_f = 38 \\
P_c = 62 \\
G_f = 2.732 \\
G_c = 2.772 \\
G_{\text{admix}} = 2.20 \text{ (Hydrated Lime)} \\
P_{\text{admix}} = 1.0
\]

\[
\text{Example of Combined Bulk Oven Dry Specific Gravity (Gsb) of Aggregate and Mineral Admixture} = \frac{38 + 62 + 1.0}{2.732 + 2.772 + 2.20} = 2.750
\]

6.4 Using Equation 4, calculate the Combined Water Absorption of the aggregate-mineral admixture blend.

**Equation 4:**

\[
\left( \frac{\text{Combined Water Absorption}}{\text{of Aggregate and Mineral Admixture Blend}} \right) = \frac{(P_f \times A_f) + (P_c \times A_c) + (P_{\text{admix}} \times A_{\text{admix}})}{P_f + P_c + P_{\text{admix}}}
\]

Where:  
\( P_f, P_c \) = Weight percent of fine aggregate (minus No. 4) and coarse aggregate (plus No. 4) respectively. Determined from the aggregate composite without mineral admixture.  
\( P_{\text{admix}} \) = Percent mineral admixture by weight of the aggregate.  
\( P_f + P_c = 100 \)  
\( P_f + P_c + P_{\text{admix}} = 100 + \% \text{ Mineral Admixture} \)  
\( A_f, A_c \) = Percent water absorption of the coarse aggregate and the fine aggregate respectively.  
\( A_{\text{admix}} \) = Percent water absorption of mineral admixture (assumed to be 0.0%).
Example:  \[\begin{align*}
P_f &= 38 \\
P_c &= 62 \\
A_f &= 2.42 \\
A_c &= 1.68 \\
P_{\text{admix}} &= 1.0 \\
A_{\text{admix}} &= 0.00
\end{align*}\]

\[
\frac{\text{Combined Water Absorption of Aggregate and Mineral Admixture}}{38 + 62 + 1.0} = \frac{(38 \times 2.42) + (62 \times 1.68) + (1.0 \times 0.00)}{38 + 62 + 1.0} = 1.94
\]

7. PREPARATION OF SPECIMENS FOR BULK SPECIFIC GRAVITY/BULK DENSITY AND MARSHALL STABILITY/FLOW DETERMINATION

7.1 Specimens shall be prepared as described herein, utilizing the apparatus specified in ARIZ 410.

Note 4: Normally a range of 3 different CRA binder contents at 1.0% increments will provide sufficient information, although in some cases it may be necessary to prepare additional sets of samples at other CRA binder contents. Two series of CRA binder contents are customarily used: either 6.0%, 7.0%, and 8.0% CRA by total mix weight; or 6.5%, 7.5%, and 8.5% CRA by total mix weight.

7.2 The aggregate-mineral admixture samples, prepared as specified in Section 5, shall be dried to constant weight at 325 ± 5 °F and shall be at this temperature at the time of mixing with the CRA. If necessary, a small amount of proportioned minus No. 8 aggregate make-up material shall be added to bring samples to the desired weight.

7.3 Before each batch of AR-AC is mixed, the CRA shall be heated in a loosely covered container in a forced draft oven for approximately 2 hours or as necessary to reach a temperature of 325 ± 5 °F (See CAUTION below). Upon removal from the oven, the CRA shall be thoroughly stirred to uniformly distribute the rubber particles throughout the CRA before adding the designated proportion to the aggregate-mineral admixture blend. If there is any delay before beginning of mixing the CRA with the aggregate-mineral admixture blend, thoroughly stir the CRA again immediately before pouring.
**CAUTION:** To avoid damage to the CRA, a hot plate or open flame is not used to bring it to the specified temperature. Once the CRA temperature has reached $325 \pm 5 \, ^\circ\mathrm{F}$, the container may briefly be moved to a hot plate for no more than 5 minutes at a time to maintain that temperature. If a hot plate is utilized, a suitable shield, baffle plate, or sand bath shall be used on the surface of the hot plate to minimize localized overheating. The CRA shall be constantly stirred to avoid sticking or scorching. Do not heat the CRA longer than necessary to complete batching and mixing operations (approximately three hours total heating time), or damage may occur.

7.4 Using Equation 5, calculate the weight of CRA to be used for each CRA binder content:

\[
\text{Weight of CRA} = \frac{\text{Weight of Aggregate and Mineral Admixture}}{100 - \text{Percent of CRA}} \times \text{Percent of CRA}
\]

Example (for Marshall specimens with 7.5% CRA) (See Note 4):

\[
\begin{align*}
\text{Weight of CRA} &= \frac{3030}{100 - 7.5} \times 7.5 = 245.7 \text{ grams}
\end{align*}
\]

**Note 5:** Before each batch is mixed, the mixing bowl and whip shall be heated to $325 \pm 5 \, ^\circ\mathrm{F}$.

7.5 The aggregate-mineral admixture blend and the appropriate amount of CRA shall be mixed together for 90 to 120 seconds at the required mixing temperature of $325 \pm 5 \, ^\circ\mathrm{F}$. **Mechanical mixing is required.**

**Note 6:** Although a wide range of mixers may provide the desired well-coated homogeneous mixture, commercial dough mixers with whips are often used. Minimum recommended capacity of the mixing bowl is 12 quarts.
Note 7: After mechanical mixing, hand mixing shall be used as necessary to produce a well-coated homogeneous mixture.

7.6 Immediately after mixing, each batch of AR-AC shall be thoroughly blended and spread according to the procedures described in ARIZ 416. The circular mass shall be cut into 6 equal pie-shaped segments. Take opposite segments for each individual specimen and use up the entire batch.

7.7 Each individual AR-AC specimen shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The specimens shall be oven-cured for 2 hours ± 10 minutes at 325 ± 5 ºF.

7.8 A mold assembly (base plate, mold, and collar) shall be heated to approximately 325 ± 5 ºF. The face of the compaction hammer shall be thoroughly cleaned and heated on a hot plate set at 325 ± 5 ºF. A suitable shield, baffle plate, or sand bath shall be used on the surface of the hot plate to minimize localized overheating of the face of the hammer.

7.9 Place a 4-inch diameter paper disc in the bottom of the mold before the mixture is introduced. Quickly place the mixture into the mold using a transfer bowl or other suitable device. Spade the mixture vigorously with a heated flat metal spatula, with a blade approximately 1-inch wide and 6-inches long and stiff enough to penetrate the entire layer of material, 15 times around the perimeter and 10 times at random into the mixture, penetrating the mixture to the bottom of the mold. Smooth the surface of mix to a slightly rounded shape.

Note 8: To ease removal of the end papers after compaction, they may be sprayed with a light application of aerosol based vegetable oil. PAM® cooking spray has been found to work well for this application.

7.10 Before compaction, put the mold containing the mixture in an oven as necessary to heat the mixture to the proper compaction temperature of 325 ± 5 ºF. The specimen shall not be heated for more than one hour after being placed in the mold. Prior to compaction, it shall be verified that the mixture is at the required temperature. An electronic temperature probe with a digital display or a long-stem metal thermometer with a dial face shall be used. The use of an infrared temperature gun is not allowed.

7.11 Immediately upon removing the mold assembly loaded with mix from the oven, place a paper disc on top of mixture, place the mold assembly on the compaction pedestal in the mold holder, and apply 75 blows with the
compaction hammer. Remove the base plate and collar, and reverse and reassemble the mold. Apply 75 compaction blows to the face of the reversed specimen.

**Note 9:** The compaction hammer shall apply only one blow after each fall, that is, there shall not be a rebound impact. The compaction hammer shall meet the requirements specified in Subsection 2.4 of ARIZ 410.

7.12 Remove the collar and top paper disc. Remove the base plate and remove the bottom paper disc while the specimen is still hot. Replace the base plate immediately, making sure to keep the mold and specimen oriented so that the bottom face of the compacted specimen remains directly in contact with, and is fully supported by, the base plate.

**Note 10:** Paper discs need to be removed while the AR-AC specimen is hot. The discs are very difficult to remove after the specimens have cooled.

7.13 If any part of the top surface of a compacted specimen is visually observed to increase in height (rise or swell in the mold) after compaction, stop testing and discard the prepared specimens. Adjust the gradation of the aggregate-mineral admixture blend to provide additional void space to accommodate the CRA, then batch and compact new trial AR-AC specimens. If no visible increase in height occurs, proceed with Subsections 7.14 through 7.17.

7.14 Allow each compacted specimen to cool in a vertical position in the mold (with the base plate on the bottom and the top surface exposed to air) until they are cool enough to be extruded without damaging the specimen. Rotate the base plate occasionally to prevent sticking.

**Note 11:** Generally specimens can be extruded without damage when they are at a temperature of approximately 77 to 90 ºF.

**Note 12:** Cooling may be accomplished at room temperature, or in a 77 ºF air bath. If more rapid cooling is desired, the mold and specimen may be placed in front of a fan until cool, **but do not turn the mold on its side.**

7.15 Extrude the specimen by orienting the mold so that the ram pushes on the bottom face (base plate face) of the specimen. If any specimen is deformed or damaged during extrusion, the entire set of specimens at that CRA binder content shall be discarded and a new set prepared.
7.16 Immediately upon extrusion, measure and record the height of the specimen to the nearest 0.001 inch and determine and record its weight in air to the nearest 0.1 gram. Compacted specimens shall be 2.500 ± 0.200 inches in height. If this criteria is not met for the specimens at each CRA binder content, the entire set of specimens at that CRA binder content shall be discarded and a new set prepared after necessary adjustments in the weight of aggregate have been made using Equation 2 (see Note 2 in Table 1).

7.17 Repeat the procedures in Subsections 7.4 through 7.16 for the required specimens.

8. BULK SPECIFIC GRAVITY/BULK DENSITY OF SPECIMENS

8.1 Determine the bulk specific gravity and bulk density of the three compacted AR-AC specimens at each CRA binder content in accordance with Method A of ARIZ 415, except the paraffin method shall not be used. The determination of the "Weight in Water" and "S.S.D. Weight" of each specimen will be completed before the next specimen is submerged for its "Weight in Water" determination.

Note 13: Specimens fabricated in the laboratory that have not been exposed to moisture do not require drying after extrusion from the molds. The specimen weight in air obtained in Subsection 8.1 is its dry weight.

8.2 For each CRA binder content, the densities of individual compacted specimens shall not differ by more than 2.0 pcf. If this density requirement is not met, the entire set of specimens at that CRA binder content shall be discarded and a new set of specimens prepared.

8.3 For each set of three specimens, determine the average bulk specific gravity ($G_{mb}$), and using that value calculate the average bulk density ($G_{mb}$ multiplied by 62.3 lb./cu. ft.).
9. **MARSHALL STABILITY AND FLOW DETERMINATION**

9.1 Specimens shall be tested for Marshall Stability and Flow utilizing the apparatus specified in ARIZ 410.

*Note 14:* Marshall Stability and Flow values are determined and recorded for information only.

9.2 For each specimen, determine the uncorrected stability, the stability (corrected for height), and the flow according to ARIZ 410, Subsections 4.6 through 4.11. In addition, determine the average stability (corrected for height) and average flow for each set of specimens prepared at each CRA binder content.

*Note 15:* Flow values may be high compared to conventional asphaltic concrete mixtures.

10. **MAXIMUM THEORETICAL SPECIFIC GRAVITY (RICE TEST)**

10.1 The maximum theoretical specific gravity of the mixture shall be determined in accordance with ARIZ 806 with the following modifications:

10.1.1 Prepare the AR-AC specimens including mineral admixture according to the procedures described in Section 5 and Subsections 7.2 through 7.5 using 6.0% or 6.5% CRA by total mix weight. A liquid anti-stripping agent is not used.

10.1.2 Immediately after mixing, the material shall be thoroughly blended and spread according to the procedures described in ARIZ 416. The circular mass shall be cut into 6 equal pie-shaped segments. Take opposite segments for each individual test sample and use up the entire batch.

10.1.3 Each individual test sample shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The test samples shall be oven-cured for 2 hours ± 10 minutes at 325 ± 5 °F.

10.1.4 After curing, each test sample shall be spread on a sheet of heavy paper or in a large flat bottom pan. Before the samples are completely cooled, separate the particles of the mixture, taking care not to fracture the mineral aggregate particles, so that the particles of the fine aggregate portion are not larger than ¼ inch. Allow the test samples to cool to room temperature.
10.1.5 Using Equation 6, calculate the effective specific gravity of the combined aggregate and mineral admixture (G_{se}).

**Equation 6:**

\[
G_{se} = \frac{100 - P_b}{100 - \frac{P_b}{G_{mm} - \frac{100}{G_b}}}
\]

Where:
- \(G_{se}\) = Effective specific gravity of the combined aggregate and mineral admixture.
- \(P_b\) = CRA binder content at which the Rice test was performed (Rice \(P_b\)).
- \(G_{mm}\) = Measured maximum theoretical specific gravity of the mix at Rice \(P_b\).
- \(G_b\) = Specific gravity of the CRA.

Example:
- \(P_b = 6.0\)
- \(G_{mm} = 2.547\)
- \(G_b = 1.037\)

\[
G_{se} = \frac{100 - 6.0}{100 - \frac{6.0}{2.547 - \frac{100}{1.037}}} = 2.808
\]

10.1.6 Using Equation 7, calculate the maximum theoretical specific gravity (\(G_{mm}\)) for each CRA binder content.

**Note16:** \(G_{se}\) is considered constant regardless of the CRA binder content.

**Equation 7:**

\[
G_{mm} = \frac{100}{P_{sa} + \frac{P_b}{G_{se} + \frac{100}{G_b}}}
\]

Where:
- \(G_{mm}\) = Calculated maximum theoretical specific gravity of the AR-AC at each CRA binder content \(P_b\).
- \(P_{sa}\) = Aggregate and mineral admixture content, percent by total weight of mix (100-\(P_b\)).
- \(P_b\) = Percent CRA binder content, by total weight of mix, for each CRA binder content.
G_{se} = \text{Effective specific gravity of the combined aggregate and mineral admixture.} \\
G_{b} = \text{Specific gravity of the CRA.}

Example (for 7.5\% CRA binder content) (See Note 4):

P_{sa} = 92.5 \\
P_{b} = 7.5 \\
G_{se} = 2.808 \\
G_{b} = 1.037

\[
G_{mm} = \frac{\frac{100}{P_{sa}} + \frac{P_{b}}{G_{se}}}{\frac{G_{se}}{G_{b}}} = 2.489
\]

11. DETERMINATION OF DESIGN PERCENT CRA BINDER CONTENT

11.1 The design percent CRA binder content is determined as follows in Subsections 11.2 through 11.3.

11.2 For each CRA binder content used, calculate effective voids (V_{a}), percent absorbed CRA (P_{ba}), and voids in mineral aggregate (VMA) using the following equations.

11.2.1 Using Equation 8, calculate the effective voids (V_{a}). The calculated G_{mm} values for the respective CRA binder contents are used to determine the corresponding effective voids content of the compacted Marshall specimens at each CRA binder content.

\[
V_{a} = \left( \frac{G_{mm} - G_{mb}}{G_{mm}} \right) \times 100
\]

Where: \(V_{a}\) = Effective voids in the compacted mixture, percent of total volume. 
\(G_{mm}\) = Calculated maximum theoretical specific gravity of the AC-AR at each CRA binder content \(P_{b}\). 
\(G_{mb}\) = Bulk specific gravity of compacted mixture specimens.
Example (for 7.5% CRA binder content) (See Note 4):

\[
G_{mm} = 2.489 \\
G_{mb} = 2.263
\]

\[
V_a = \frac{2.489 - 2.263}{2.489} \times 100 = 9.1
\]

11.2.2 Using Equation 9, calculate the percent absorbed CRA (P_{ba}).

**Note17:** The percent absorbed CRA (P_{ba}) is the same for all CRA binder contents.

**Equation 9:**

\[
P_{ba} = \left( \frac{G_{se} - G_{sb}}{G_{sb} \times G_{se}} \right) \times G_b \times 100
\]

Where: 
- \( P_{ba} \) = Absorbed CRA, percent by total weight of mix.
- \( G_{se} \) = Effective specific gravity of the combined aggregate and mineral admixture.
- \( G_{sb} \) = Bulk oven dry specific gravity of the combined aggregate and mineral admixture.
- \( G_b \) = Specific gravity of the CRA.

Example:
- \( G_{se} = 2.808 \)
- \( G_{sb} = 2.750 \)
- \( G_b = 1.037 \)

\[
P_{ba} = \frac{2.808 - 2.750}{2.750 \times 2.808} \times 1.037 \times 100 = 0.78
\]

11.2.3 Using Equation 10, calculate voids in mineral aggregate (VMA) of the compacted Marshall specimens at each CRA binder content.

**Equation 10:**

\[
VMA = 100 - \left( \frac{G_{mb} \times P_{sa}}{G_{sb}} \right)
\]
Where:  

\[ VMA = \text{voids in the mineral aggregate, percent of bulk volume.} \] 
\[ G_{mb} = \text{bulk specific gravity of compacted mixture specimens.} \] 
\[ P_{sa} = \text{aggregate and mineral admixture content, percent by total weight of mix (100-P_{b}).} \] 
\[ G_{sb} = \text{bulk oven dry specific gravity of the combined aggregate and mineral admixture.} \]

Example (for 7.5% CRA binder content) (See Note 4):

\[ G_{mb} = 2.263 \] 
\[ P_{sa} = 92.5 \] 
\[ G_{sb} = 2.750 \]

\[ VMA = 100 - \frac{2.263 \times 92.5}{2.750} = 23.9 \]

11.3 If the specified value for effective voids \((V_a)\) falls between the resultant \(V_a\) values determined for two of the CRA binder contents used, straight-line interpolation shall be used to calculate the bulk specific gravity \((G_{mb})\) at each 0.1% CRA binder content increment between the two corresponding measured values of \(G_{mb}\). Using equations 7 and 8, determine the values for \(G_{mm}\) and \(V_a\) at each interpolated CRA binder content. Select the CRA binder content which results in \(V_a\) being as close as possible to the specified value.

11.3.1 Using equation 10, calculate \(VMA\) at the selected CRA binder content. Using straight-line interpolation, determine the values for stability and flow that correspond to the selected CRA binder content. (Stability and flow values are determined for information only.) If the value for \(VMA\) is within the limits of the specifications, the selected CRA binder content shall be the design CRA binder content.

11.3.2 On separate graphs (using the same scale for percent CRA binder for each graph), plot the values for bulk specific gravity, effective voids, \(VMA\), stability, and flow for each set of three specimens and at the design CRA binder content versus the percent CRA binder. Draw a smooth “best fit” curve based on the plotted points, as shown in Figure 4. The graphs in Figure 4 are for illustrative purposes only. Straight-line interpolation, as specified in Subsections 11.3 and 11.3.1, is used for mix design calculations. (Values for stability and flow are plotted for information only.)
11.3.3 If it is not possible to obtain specification compliance within the range of CRA binder contents used, a determination must be made to either redesign the mix (different aggregate gradation and/or source) or prepare additional specimens at other CRA binder contents for testing and volumetric analyses.

11.4 Using Equation 11, calculate the percent effective CRA binder content ($P_{be}$) at each percent CRA binder content, including the design percent CRA binder content.

**Equation 11:**\[ P_{be} = P_b - \left( \frac{P_{ba} \times P_{sa}}{100} \right) \]

Where:
- $P_{be}$ = Percent effective CRA binder content of the mixture (free binder not absorbed).
- $P_b$ = CRA binder content, percent by total weight of mix.
- $P_{ba}$ = Absorbed CRA, percent by total weight of mix.
- $P_{sa}$ = Aggregate and mineral admixture content, percent by total weight of mix (100-$P_b$).

Example (for 9.3% CRA design binder content):
- $P_b = 9.3$
- $P_{ba} = 0.78$
- $P_{sa} = 90.7$

\[ P_{be} = 9.3 - \frac{0.78 \times 90.7}{100} = 8.59 \]

12. **MIX DESIGN GRADATION TARGET VALUES**

12.1 The target values for the aggregate composite without mineral admixture, and the aggregate-mineral admixture blend composite, for the AR-AC mixture shall be expressed as percent passing particular sieve sizes as required by the specifications for the project. An example mix design composite, with and without mineral admixture, is shown in Figure 1.
13. REPORT

13.1 Report the test results and data obtained on the appropriate form. Liberal use of the remarks area to clarify and/or emphasize any element of the design is strongly recommended. Information required in the mix design report includes the items listed below:

13.1.1 Name and address of the testing organization and the signature and seal of the mix design engineer responsible for the mix design.

13.1.2 Aggregate:
- Source and identification
- Individual stockpile or bin gradations
- Aggregate blend proportions and composite gradation, with and without mineral admixture
- Fine and coarse aggregate specific gravities (Bulk Oven Dry, SSD, Apparent) and absorption
- Combined aggregate specific gravities (Bulk Oven Dry, SSD, Apparent)
- Combined aggregate absorption
- Aggregate quality
  - LA Abrasion
  - Sand Equivalent
  - Fractured Coarse Aggregate Particles (percentage with one fractured face and percentage with two fractured faces)
  - Uncompacted Void Content
  - Carbonates (when applicable)

13.1.3 Mineral Admixture:
- Type and source
- Percentage used
- Specific gravity

13.1.4 Combined Mineral Aggregate and Mineral Admixture
- Combined specific gravities [Bulk Oven Dry ($G_{sb}$), SSD, Apparent] and absorption of the combined aggregate and mineral admixture
- Effective specific gravity of the combined aggregate and mineral admixture ($G_{se}$)
13.1.5 CRA Binder Design (from supplier), including:
- Source and grade of base asphalt cement
- Source and type of crumb rubber
- Crumb rubber gradation
- Proportions of asphalt cement and crumb rubber
  - Percentage of crumb rubber, by weight of asphalt cement
  - Percentages of crumb rubber and asphalt cement, by weight of total binder
- CRA binder properties, in compliance with Section 1009 of the ADOT Specifications
- CRA binder specific gravity ($G_b$) (at 77 °F)
- Asphalt cement specific gravity (at 77 °F)

13.1.6 Maximum theoretical specific gravity ($G_{mm}$) and density (pcf) at the CRA binder content at which the Rice test was performed ($Rice P_b$)

13.1.7 Mixture Compaction Trials:
- Percent CRA binder content ($P_b$), by weight of the bituminous mixture
- Aggregate and mineral admixture content ($P_{sa}$)
- Calculated maximum theoretical specific gravity ($G_{mm}$) and density (pcf)
- Bulk specific gravity ($G_{mb}$) and bulk density (pcf) of Marshall specimens
- Percent effective voids ($V_a$)
- Percent voids in mineral aggregate (VMA)
- Percent absorbed CRA ($P_{ba}$)
- Percent effective CRA binder content ($P_{be}$)
- Marshall stability (nearest 10 pounds) (recorded for information only)
- Marshall flow (0.01 inch) (recorded for information only)

13.1.8 Plots of the following properties versus percent CRA binder content (See examples in Figure 4.):
- Percent effective voids ($V_a$)
- Percent voids in mineral aggregate (VMA)
- Bulk specific gravity ($G_{mb}$)
- Marshall stability (for information only)
- Marshall flow (for information only)
13.1.9 Mixture Properties at Design Percent CRA Binder Content:
- Percent CRA binder content ($P_b$), by weight of the bituminous mixture
- Aggregate and mineral admixture content ($P_{sa}$)
- Calculated maximum theoretical specific gravity ($G_{mm}$) and density (pcf)
- Bulk specific gravity ($G_{mb}$) and bulk density (pcf) of Marshall specimens
- Percent effective voids ($V_a$)
- Percent voids in mineral aggregate (VMA)
- Percent absorbed CRA ($P_{ba}$)
- Percent effective CRA binder content ($P_{be}$)
- Marshall stability (nearest 10 pounds) (recorded for information only)
- Marshall flow (0.01 inch) (recorded for information only)

13.1.10 Mix Design Summary. The mix design summary shall contain the information shown in Figure 5. [Pages 1 and 2 of an example Mix Design Summary for an Asphaltic Concrete (Asphalt-Rubber) Marshall Mix Design.]
## Example Mix Design Composite
(With and Without Mineral Admixture)

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<th>Material</th>
<th>3/16&quot; Washed Crusher Fines</th>
<th>1/4&quot; Washed Crusher Fines</th>
<th>3/8&quot; Aggregate</th>
<th>1/2&quot; Aggregate</th>
<th>Composite</th>
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<tr>
<td>Percent used in composite</td>
<td>15</td>
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<table>
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<th>With Admix</th>
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<table>
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* 1.0 percent admixture, by weight of dry mineral aggregate, is used.

FIGURE 1
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<th>INDIV. % RET.</th>
<th>ACCUM. % RET.</th>
<th>FRACTURED COARSE AGGREGATE PARTICLES (Plus #4) (500 grams)</th>
<th>PERCENT CARBONATES (Plus #4) (300 grams)</th>
<th>COARSE AGGREGATE SPECIFIC GRAVITY/ABSORPTION (Plus #4) (4000 grams)</th>
<th>FINE AGGREGATE SPECIFIC GRAVITY/ABSORPTION (Minus #4) (1200 grams)</th>
<th>SAND EQUIVALENT (Minus #4) (600 grams)</th>
<th>UNCOMPACTED VOID CONTENT (Minus #8) (500 grams)</th>
<th>ABRASION [Type C] (5000 grams)</th>
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<td>0.890</td>
<td>98.666</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1157</td>
<td>579</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>1.334</td>
<td>100.00</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1200</td>
<td>600</td>
<td>500</td>
</tr>
</tbody>
</table>
### Example Weigh Up Sheet #2

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SIEVE</th>
<th>INDIVIDUAL PERCENT RETAINED</th>
<th>ACCUMULATIVE PERCENT RETAINED</th>
<th>RICE (3000 grams)</th>
<th>MARSHALL (3000 grams)</th>
<th>MINUS NO. 8 MAKE-UP MATERIAL (500 grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>ACCUMULATIVE WEIGHT (grams)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/16&quot; Washed Crusher Fines</td>
<td>#4</td>
<td>0.030</td>
<td>0.030</td>
<td>1</td>
<td>1</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>3.720</td>
<td>3.750</td>
<td>113</td>
<td>113</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>11.250</td>
<td>15.000</td>
<td>450</td>
<td>450</td>
<td>294</td>
</tr>
<tr>
<td>1/4&quot; Washed Crusher Fines</td>
<td>#4</td>
<td>0.702</td>
<td>15.702</td>
<td>471</td>
<td>471</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>3.717</td>
<td>19.419</td>
<td>583</td>
<td>583</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>4.581</td>
<td>24.000</td>
<td>720</td>
<td>720</td>
<td>413</td>
</tr>
<tr>
<td>3/8&quot; Aggregate</td>
<td>3/8&quot;</td>
<td>1.380</td>
<td>25.380</td>
<td>761</td>
<td>761</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>1/4&quot;</td>
<td>6.600</td>
<td>31.980</td>
<td>959</td>
<td>959</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>9.690</td>
<td>41.670</td>
<td>1250</td>
<td>1250</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>10.350</td>
<td>52.020</td>
<td>1561</td>
<td>1561</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>1.980</td>
<td>54.000</td>
<td>1620</td>
<td>1620</td>
<td>465</td>
</tr>
<tr>
<td>1/2&quot; Aggregate</td>
<td>1/2&quot;</td>
<td>3.634</td>
<td>57.634</td>
<td>1729</td>
<td>1729</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>3/8&quot;</td>
<td>20.240</td>
<td>77.874</td>
<td>2336</td>
<td>2336</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>1/4&quot;</td>
<td>18.630</td>
<td>96.504</td>
<td>2895</td>
<td>2895</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>1.472</td>
<td>97.976</td>
<td>2939</td>
<td>2939</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.690</td>
<td>98.666</td>
<td>2960</td>
<td>2960</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>- #8</td>
<td>1.334</td>
<td>100.000</td>
<td>3000</td>
<td>3000</td>
<td>500</td>
</tr>
</tbody>
</table>

Weight of Mineral Admixture (Hydrated Lime) [1.0%, by weight of Dry Mineral Aggregate] 30 30 ---

Total weight of Mineral Aggregate and Mineral Admixture 3030 3030 ---

**FIGURE 3**
The above graphs are for illustrative purposes only. Straight-line interpolation, as specified in Subsections 11.3 and 11.3.1, is used for mix design calculations.

FIGURE 4
Asphaltic Concrete (Asphalt-Rubber) Marshall Mix Design

**Example Mix Design Summary (Page 1 of 2)**

**Date:** February 22, 2013

**Contractor:** Best Paving Contractors, Inc.

**ADOT Project Number:** STP-777-Z(777)A

**ADOT TRACS Number:** H77701C

**Project Name:** Uptown - Downtown

**Mix Designation:** ADOT 415 [Asphaltic Concrete (Asphalt-Rubber)]

**ADOT Project Number:** STP-777-Z(777)A

**Mix Design Laboratory:** XYZ Testing Laboratories

**Mix Design Engineer:** Frank Dogood

**Grade of Asphalt-Rubber Binder:** CRA Type 2

### COMPOSITE GRADATION:

<table>
<thead>
<tr>
<th>Mineral Aggregate</th>
<th>Percent Used</th>
</tr>
</thead>
<tbody>
<tr>
<td>3/16&quot; Washed Crusher Fines</td>
<td>15</td>
</tr>
<tr>
<td>1/4&quot; Washed Crusher Fines</td>
<td>9</td>
</tr>
<tr>
<td>3/8&quot; Aggregate</td>
<td>30</td>
</tr>
<tr>
<td>1/2&quot; Aggregate</td>
<td>45</td>
</tr>
</tbody>
</table>

| Hydrated Lime | 1.0 |

<table>
<thead>
<tr>
<th>Mineral Admixture</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieve Size</td>
<td>Percent Passing w/o Admix</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>100</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>96</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>75</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>50</td>
</tr>
<tr>
<td>#4</td>
<td>38</td>
</tr>
<tr>
<td>#8</td>
<td>19</td>
</tr>
<tr>
<td>#10</td>
<td>17</td>
</tr>
<tr>
<td>#16</td>
<td>12</td>
</tr>
<tr>
<td>#30</td>
<td>7</td>
</tr>
<tr>
<td>#40</td>
<td>5</td>
</tr>
<tr>
<td>#50</td>
<td>4</td>
</tr>
<tr>
<td>#100</td>
<td>3</td>
</tr>
<tr>
<td>#200</td>
<td>2.0</td>
</tr>
</tbody>
</table>

### DESIGN DATA:

<table>
<thead>
<tr>
<th>% CRA</th>
<th>7.5</th>
<th>8.5</th>
<th>9.3</th>
<th>9.5</th>
<th>Spec.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Specific Gravity</td>
<td>2.263</td>
<td>2.282</td>
<td>2.290</td>
<td>2.292</td>
<td></td>
</tr>
<tr>
<td>Bulk Density (lb./cu. ft.)</td>
<td>141.0</td>
<td>142.2</td>
<td>142.7</td>
<td>142.8</td>
<td></td>
</tr>
<tr>
<td>Stability (lb.)</td>
<td>1780</td>
<td>1790</td>
<td>1930</td>
<td>1970</td>
<td></td>
</tr>
<tr>
<td>Flow (0.01 inch)</td>
<td>25</td>
<td>27</td>
<td>29</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>Effective Voids (%)</td>
<td>9.1</td>
<td>6.9</td>
<td>5.5</td>
<td>5.1</td>
<td>5.5 ± 0.5</td>
</tr>
<tr>
<td>VMA (%)</td>
<td>23.9</td>
<td>24.1</td>
<td>24.5</td>
<td>24.6</td>
<td>19.0 Min.</td>
</tr>
<tr>
<td>Effective CRA, Total Mix (%)</td>
<td>6.78</td>
<td>7.79</td>
<td>8.59</td>
<td>8.79</td>
<td></td>
</tr>
<tr>
<td>Absorbed CRA (%)</td>
<td>0.78</td>
<td>0 – 1.0</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Max. Theoretical Specific Gravity at 9.3% CRA = 2.423
Max. Theoretical Density at 9.3% CRA = 151.0 lb/cu. ft.

### AGGREGATE PROPERTIES:

<table>
<thead>
<tr>
<th>Test Characteristic</th>
<th>Fine</th>
<th>Coarse</th>
<th>Combined</th>
<th>Spec.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk O.D. Specific Gravity</td>
<td>2.732</td>
<td>2.772</td>
<td>2.757</td>
<td>2.350 – 2.850</td>
</tr>
<tr>
<td>SSD Specific Gravity</td>
<td>2.798</td>
<td>2.819</td>
<td>2.811</td>
<td>-----</td>
</tr>
<tr>
<td>Apparent Specific Gravity</td>
<td>2.925</td>
<td>2.907</td>
<td>2.914</td>
<td>-----</td>
</tr>
<tr>
<td>Absorption</td>
<td>2.42</td>
<td>1.68</td>
<td>1.96</td>
<td>0 – 2.5%</td>
</tr>
<tr>
<td>Sand Equivalent</td>
<td>97</td>
<td>Min. 55</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Uncompacted Void Content</td>
<td>47.8</td>
<td>Min. 45.0%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carbonates</td>
<td>2</td>
<td>Max. 20%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fractured Coarse Aggregate Particles</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At least one fractured face</td>
<td>99</td>
<td>Min. 92%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>At least two fractured faces</td>
<td>97</td>
<td>Min. 85%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abrasion (Type C)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 Rev. % Loss</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>500 Rev. % Loss</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**ADDITIONAL DATA / INFORMATION:**

- Source of Mineral Aggregate: Excellent Aggregate Materials, Inc.
- Mineral Admixture Type/Specific Gravity: Hydrated Lime/2.20

### RECOMMENDED CRA BINDER CONTENT:

(By weight of total mix) 9.3%

### COMBINED AGGREGATE AND MINERAL ADMIXTURE PROPERTIES:

- Bulk O.D. Specific Gravity: 2.750
- SSD Specific Gravity: 2.803
- Apparent Specific Gravity: 2.905
- Water Absorption: 1.94
- Effective Specific Gravity: 2.808

### REMARKS:

- Figure 5 (Continued on Next Page)
Asphaltic Concrete (Asphalt-Rubber) Marshall Mix Design
Example Mix Design Summary (Page 2 of 2)

<table>
<thead>
<tr>
<th>CRA BINDER DESIGN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source of Base Asphalt Cement: Supreme Asphalt Suppliers, Inc.</td>
</tr>
<tr>
<td>Source of Crumb Rubber: Rubber Suppliers, Inc.</td>
</tr>
<tr>
<td>Grade of Base Asphalt Cement: PG 58-22</td>
</tr>
<tr>
<td>Type of Crumb Rubber: Type B</td>
</tr>
<tr>
<td>Grade of Asphalt-Rubber Binder: CRA Type 2</td>
</tr>
<tr>
<td>Specific Gravity of Base Asphalt Cement: 1.009 (at 77 °F)</td>
</tr>
<tr>
<td>Specific Gravity of CRA Binder: 1.037 (at 77 °F)</td>
</tr>
<tr>
<td>Crumb Rubber (by weight of asphalt cement): 20.5% (Minimum 20% required)</td>
</tr>
<tr>
<td>Proportions of Asphalt Cement and Crumb Rubber (by weight of total binder):</td>
</tr>
<tr>
<td>83.0% Asphalt Cement</td>
</tr>
<tr>
<td>17.0% Crumb Rubber</td>
</tr>
<tr>
<td>CRA Binder Properties (ADOT Specifications Section 1009):</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Property</th>
<th>Test Result</th>
<th>Requirements (CRA Type 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rotational Viscosity: 177 °C (350 °F); (Rion Model VT-04, No. 1 Rotor); Pascal seconds</td>
<td>2.1</td>
<td>1.5 – 4.0</td>
</tr>
<tr>
<td>Penetration: 4 °C (39.2 °F), 200 g, 60 sec. (ASTM D 5); 0.1 mm</td>
<td>26</td>
<td>15 Minimum</td>
</tr>
<tr>
<td>Softening Point: (AASHTO T 53); °C</td>
<td>61</td>
<td>54 Minimum</td>
</tr>
<tr>
<td>Resilience: 25 °C (77 °F) (ASTM D 5329); %</td>
<td>47</td>
<td>20 Minimum</td>
</tr>
</tbody>
</table>

Crumb Rubber Gradation (Arizona Test Method 714):

<table>
<thead>
<tr>
<th>Sieve Size</th>
<th>Percent Passing</th>
<th>Requirements (Type B)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 10</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>No. 16</td>
<td>90</td>
<td>65 – 100</td>
</tr>
<tr>
<td>No. 30</td>
<td>30</td>
<td>20 – 100</td>
</tr>
<tr>
<td>No. 50</td>
<td>7</td>
<td>0 – 45</td>
</tr>
<tr>
<td>No. 200</td>
<td>0.3</td>
<td>0 - 5</td>
</tr>
</tbody>
</table>

FIGURE 5 (Continued from Previous Page)
MARSHALL MIX DESIGN METHOD FOR ASPHALTIC CONCRETE WITH RECLAIMED ASPHALT PAVEMENT (RAP)

(An Arizona Method)

1. SCOPE

1.1 This method is used to design Asphaltic Concrete mixes with Reclaimed Asphalt Pavement (RAP) using 4-inch diameter Marshall apparatus.

1.2 This test method involves hazardous material, operations, and equipment. This test method does not purport to address all of the safety concerns associated with its use. It is the responsibility of the user to consult and establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1.3 See Appendix A1 of the Materials Testing Manual for information regarding the procedure to be used for rounding numbers to the required degree of accuracy.

1.4 A listing of subsequent Sections and Figures in this procedure is given below:

<table>
<thead>
<tr>
<th>Section or Figure #</th>
<th>Title</th>
<th>Page #</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section 2</td>
<td>Test Methods and Apparatus</td>
<td>2</td>
</tr>
<tr>
<td>Section 3</td>
<td>Materials</td>
<td>3</td>
</tr>
<tr>
<td>Section 4</td>
<td>Determination of Composite Gradation</td>
<td>5</td>
</tr>
<tr>
<td>Section 5</td>
<td>Preparing Aggregate/RAP/Mineral Admixture Samples for Mix Design Testing</td>
<td>7</td>
</tr>
<tr>
<td>Section 6</td>
<td>Virgin Aggregate and RAP Aggregate Specific Gravities and Absorption</td>
<td>12</td>
</tr>
<tr>
<td>Section 7</td>
<td>Asphalt Binder Selection</td>
<td>20</td>
</tr>
<tr>
<td>Section 8</td>
<td>Laboratory Mixing and Compaction Temperatures</td>
<td>23</td>
</tr>
<tr>
<td>Section 9</td>
<td>Preparation of Specimens for Bulk Specific Gravity/Bulk Density and Marshall Stability/Flow Determination</td>
<td>25</td>
</tr>
<tr>
<td>Section 10</td>
<td>Bulk Specific Gravity/Bulk Density of Specimens</td>
<td>29</td>
</tr>
<tr>
<td>Section 11</td>
<td>Marshall Stability and Flow Determination</td>
<td>30</td>
</tr>
<tr>
<td>Section 12</td>
<td>Maximum Theoretical Specific Gravity (Rice Test)</td>
<td>30</td>
</tr>
<tr>
<td>Section 13</td>
<td>Determination of Design Asphalt Binder Content</td>
<td>32</td>
</tr>
<tr>
<td>Section 14</td>
<td>Immersion Compression (IMC) Test</td>
<td>36</td>
</tr>
<tr>
<td>Section 15</td>
<td>Mix Design Gradation Target Values</td>
<td>37</td>
</tr>
<tr>
<td>Section 16</td>
<td>Report</td>
<td>37</td>
</tr>
<tr>
<td>Figure 1</td>
<td>Example of Composite #1 (Virgin Aggregate and RAP Aggregate)</td>
<td>41</td>
</tr>
<tr>
<td>Figure 2</td>
<td>Example of Batch Sheet #1 (For Composite #1) (Virgin Aggregate and RAP Aggregate)</td>
<td>42</td>
</tr>
</tbody>
</table>
2. TEST METHODS AND APPARATUS

2.1 This test method is used in conjunction with the test methods and standards listed below. Requirements for the frequency of equipment calibration and verification are found in Appendix A3 of the Materials Testing Manual. The required apparatus is shown in the individual test methods and standards listed below, as appropriate.

2.1.1 Arizona Test Methods:

- ARIZ 201 Sieving of Coarse and Fine Graded Soils and Aggregates
- ARIZ 205 Composite Grading
- ARIZ 210 Specific Gravity and Absorption of Coarse Aggregate
- ARIZ 211 Specific Gravity and Absorption of Fine Aggregate
- ARIZ 212 Percentage of Fractured Coarse Aggregate Particles
- ARIZ 238 Percent Carbonates in Aggregate
- ARIZ 240 Sieve Analysis and Separation of Salvaged AC Pavement Particles for Recycled Asphaltic Concrete
- ARIZ 247 Particle Shape and Texture of Fine Aggregate Using Uncompacted Void Content
- ARIZ 248 Alternate Procedures for Sieving of Coarse and Fine Graded Soils and Aggregates
- ARIZ 251 Combined Coarse and Fine Aggregate Specific Gravity and Absorption
- ARIZ 406 Moisture Content of Bituminous Mixtures
- ARIZ 410 Compaction and Testing of Bituminous Mixtures Utilizing Four Inch Marshall Apparatus
- ARIZ 415 Bulk Specific Gravity and Bulk Density of Compacted Bituminous Mixtures
- ARIZ 416 Preparing and Splitting Field Samples of Bituminous Mixtures for Testing
• ARIZ 417 Maximum Theoretical Specific Gravity of Field
Produced Bituminous Mixtures
• ARIZ 802 Effect of Water on Strength of Compacted Treated
and Untreated Bituminous Mixtures (Immersion
Compression Test)
• ARIZ 806 Maximum Theoretical Specific Gravity of Laboratory
Prepared Bituminous Mixtures (Rice Test)

2.1.2 AASHTO Standards:
• AASHTO M 320 Standard Specification for Performance-Graded
Asphalt Binder
• AASHTO R 29 Grading or Verifying the Performance Grade of an
Asphalt Binder
• AASHTO T 96 Resistance to Degradation of Small-Size Coarse
Aggregate by Abrasion and Impact in the
Los Angeles Machine
• AASHTO T 164 Quantitative Extraction of Asphalt Binder from
Hot-Mix Asphalt (HMA) – Method A
• AASHTO T 170 Recovery of Asphalt from Solution by
Abson Method
• AASHTO T 176 Plastic Fines in Graded Aggregates and Soils by
Use of the Sand Equivalent Test
• AASHTO T 228 Specific Gravity of Semi-Solid Bituminous Materials

Note 1: Testing by AASHTO T 228 shall be performed at 77 °F.

• AASHTO R 76 Reducing Samples of Aggregate to Testing Size
• AASHTO T 316 Viscosity Determination of Asphalt Binder Using
Rotational Viscometer

2.1.3 ASTM Standards:
• ASTM D 2493 Standard Viscosity-Temperature Chart for Asphalts
• ASTM D 5404 Standard Practice of Recovery of Asphalt from
Solution Using the Rotary Evaporator

3. MATERIALS

3.1 Mineral Aggregate - The mineral aggregate used in the design shall be
produced material from the source(s) for the project. Mineral aggregate
shall consist of a combination of virgin aggregate and aggregate from
RAP (RAP aggregate). When the terms “mineral aggregate” or
“aggregate” are used without being further described as “virgin” or “RAP”, the intended meaning is the total aggregate material used in the mixture.

3.1.1 RAP aggregate used for mix design purposes shall be extracted aggregate from AASHTO T 164 (Method A), except RAP aggregate used for Abrasion testing may be obtained using an ignition furnace.

3.1.2 The composited gradation of the virgin aggregate and RAP aggregate, and the composited gradation of the virgin aggregate-RAP aggregate-mineral admixture blend, shall comply with the grading limits of the specifications.

3.1.3 The composited virgin mineral aggregate shall conform to the requirements of the specifications for Sand Equivalent (AASHTO T 176), Combined Bulk Oven Dry Specific Gravity (ARIZ 251), Combined Water Absorption (ARIZ 251), and when applicable, Uncompacted Void Content (ARIZ 247).

3.1.4 The composited virgin aggregate and RAP aggregate shall conform to the requirements of the specifications for Fractured Coarse Aggregate Particles (ARIZ 212) and when applicable, Percent Carbonates (ARIZ 238).

3.1.5 Virgin mineral aggregate and RAP aggregate from each source shall be tested separately for compliance to the project requirements for Abrasion (AASHTO T 96). If multiple RAP stockpiles are utilized, and if the stockpiles are from different sources, an abrasion test will be necessary for each source. If the RAP is obtained from one source and is fractionated, one abrasion test will suffice.

3.2 Reclaimed Asphalt Pavement (RAP) - RAP consists of salvaged, milled, pulverized, broken, or crushed asphalt pavement. RAP contains “RAP aggregate” and bituminous materials hereinafter referred to as “RAP Binder”. The gradation of the RAP material shall conform to the requirements of the specifications.

3.3 Bituminous Material - The bituminous material used in the design shall be the total asphalt binder which is to be used in the production of the asphaltic concrete. Bituminous material shall consist of a combination of virgin asphalt binder and asphalt binder from RAP (RAP binder). When the terms “bituminous material” or “asphalt binder” or “binder” are used without being further described as “virgin” or “RAP”, the intended meaning is the total bituminous material used in the mixture.

3.3.1 If ≤15% RAP binder is being used, the virgin binder must conform to the requirements of Section 1005 of the specifications for the specific grade of
binder shown in the Special Provisions. The specific gravity of the virgin asphalt binder used in the asphaltic concrete shall be determined in accordance with AASHTO T 228 (at 77 °F). (See Section 7.)

3.3.2 If >15% RAP binder is being used, the blended RAP binder and virgin binder must conform to the requirements of Section 1005 of the specifications for the specific grade of binder shown in the Special Provisions. The specific gravity of the virgin asphalt binder, the specific gravity of the RAP binder from each RAP stockpile, and the specific gravity of the blended RAP binder and virgin binder shall be determined in accordance with AASHTO T 228 (at 77 °F). (See Section 7.) As an alternate to determining the specific gravity of the recovered RAP binder from each RAP stockpile, the specific gravity of proportionally blended recovered RAP binder may be determined. (See Note 9.)

3.4 Mineral Admixture - Mineral admixture is required. The mineral admixture used in the design shall be the same type of material which is to be used in production of the asphaltic concrete. The mineral admixture shall conform to the requirements of the specifications.

3.5 Batch Plants - Mix designs developed for asphaltic concrete which is to be produced in a batch plant shall be prepared in accordance with this procedure, modified as follows:

3.5.1 Virgin aggregate samples will be obtained from the hot bins for all testing except Sand Equivalent. Testing for Sand Equivalent will be performed on virgin aggregate stockpile samples proportioned and composited to meet Composite #2 without admixture. Any method may be used to combine the virgin aggregate stockpile samples for Sand Equivalent testing as long as the resultant gradation is representative of the composite gradation without admixture shown in Composite #2.

3.5.2 The mix design shall provide component percentages and composite gradations for both stockpile and hot bin materials.

4. DETERMINATION OF COMPOSITE GRADATION

4.1 The gradation of the aggregate from each individual component stockpile, bin, or RAP aggregate, shall be determined in accordance with ARIZ 248 using washed sieve analysis Alternate #3, Alternate #4, or Alternate #5. The RAP aggregate gradation shall be determined using the aggregate from solvent extraction testing of the RAP. For Alternate #5, washing of the coarse aggregate may be performed on the composite plus No. 4 material and applied to the composite percent pass the
minus No. 200 determined from the unwashed coarse sieving and washed fine sieving of the individual stockpiles.

4.2 The composite gradation of the virgin aggregate and RAP aggregate blend is determined using desired percentages of each component based on washed sieve analysis. Mix designs may be developed based on bin or stockpile material, as appropriate for the respective mix production facility to be used.

4.3 The virgin aggregate and RAP aggregate blend composite shall be determined in accordance with ARIZ 205, paragraph 2(e).

4.4 The virgin aggregate-RAP aggregate-mineral admixture blend composite is determined by adjusting the virgin aggregate and RAP aggregate blend composite (percent passing) for mineral admixture by performing the calculation in Equation 1 for each sieve:

\[
\text{Equation 1:} \quad \left( \frac{\text{% passing each sieve in the virgin aggregate and RAP aggregate blend composite}}{(100 + \text{% Mineral Admixture})} \right) \times 100
\]

4.5 The composite gradations of the components and the percentage of each will be included in the design report for each of the composite types listed below.

4.5.1 Composite #1 - Virgin Aggregate and RAP Aggregate (with and without mineral admixture).

4.5.2 Composite #2 - Virgin Aggregate (with and without mineral admixture)

4.5.3 Composite #3 - Virgin Aggregate and Dry Screened RAP (without mineral admixture)
5. PREPARING AGGREGATE/RAP/MINERAL ADMIXTURE SAMPLES FOR MIX DESIGN TESTING

5.1 Preparation of Mineral Aggregate:

5.1.1 Dry the mineral aggregate from each individual stockpile at a temperature not exceeding any temperature restrictions specified in subsequent test procedures. Drying shall be performed until no further weight loss is obtained from continued drying.

5.1.2 Based on the virgin mineral aggregate stockpile or bin gradations and composite percentages, virgin mineral aggregate material which is retained on the individual No. 8 and larger sieve sizes, and the minus No. 8 material, is used to prepare samples for mix design testing.

5.1.3 Based on the RAP aggregate gradation(s) and composite percentage(s), RAP aggregate material which is retained on the individual No. 8 and larger sieve sizes, and the minus No. 8 material, is used to prepare samples for mix design testing.

5.2 Preparation of Reclaimed Asphalt Pavement (RAP):

5.2.1 Drying/Processing of RAP:

5.2.1.1 Based on the amount of RAP used in the composite gradation, determine the minimum weight of RAP from each stockpile needed for mix design, then increase it by 100% to be sure enough is available for possible retesting.

5.2.1.2 Spread the material from each individual RAP stockpile to be used for mix design testing in separate pans at 1 to 1½ inches in depth and dry in an oven at 140 ± 5 °F such that no further weight loss is obtained from continued drying at 140 ± 5 °F. Allow the material to cool. Split or quarter samples of RAP material from each stockpile, in accordance with AASHTO R 76, as necessary to perform the remainder of the testing and batching required in Section 5.

5.2.1.3 Determine the RAP binder content of material from each RAP stockpile, by weight of RAP, in accordance with AASHTO T 164 – Method A (Centrifuge Extraction) and the gradation of the extracted RAP aggregate in accordance with ARIZ 248. Determine the moisture content in accordance with ARIZ 406.
Note 2: For mixes utilizing >15% RAP binder, the extract from the solvent extraction is saved for testing as specified in Subsection 7.2. Specific time frames related to binder recovery must be followed. (See Subsection 7.4.1.)

5.2.1.4 The binder content of RAP material from each individual stockpile is recorded as "P_{br}\" ("P_{br1}\" and "P_{br2}\", if two RAP stockpiles are used).

5.2.2 Dry Gradation of RAP (into individual sizes for No. 8 and larger sieves, and minus No. 8 material):

5.2.2.1 Determine the appropriate size test sample of each individual RAP stockpile as shown in Table 1.

<table>
<thead>
<tr>
<th>Maximum Size of Particle</th>
<th>Minimum Weight of Sample, grams (lbs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1¼&quot;</td>
<td>10000 (22)</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>5000 (11)</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>2000 (4.4)</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>1000 (2.2)</td>
</tr>
</tbody>
</table>

5.2.2.2 Dry sieve each test sample in accordance with ARIZ 240, but utilizing the No. 8 sieve as the smallest sieve. Compute the unwashed gradation in accordance with ARIZ 248, Alternate #2. This gradation is used only for batching test specimens for Batch Sheet #3. (See Subsection 5.3.3.)

5.2.3 Screening of dry RAP into individual sizes for use in mix design batching:

5.2.3.1 Screen the RAP material from the individual stockpiles and separate into individual sizes for No. 8 and larger sieves, and minus No. 8 material. The RAP shall be screened for 5 minutes ± 15 seconds per ARIZ 240 to prevent excessive breakdown of the RAP agglomerations. The screened RAP will be used in Batch Sheet #3.

5.3 Batching of Test Samples: Multiple batch sheets, as described below, will be necessary due to the use of RAP. Batch Sheet #1 is for samples which include virgin aggregate and RAP aggregate. Batch Sheet #2 is for samples where only the virgin aggregate is used. Batch Sheet #3 is for samples that include virgin aggregate, RAP, and mineral admixture; where the virgin aggregate and mineral admixture must be wet treated prior to incorporating the RAP. Each virgin mineral aggregate, RAP aggregate, and RAP component which has been screened into individual sizes for No. 8 and larger sieves, and minus No. 8 material, will be used in the batching process.
5.3.1 Batch Sheet #1:

5.3.1.1 This batch sheet will utilize the individual virgin aggregate gradations and the RAP aggregate gradation(s) determined from solvent extraction (except RAP aggregate used for abrasion testing may be obtained using an ignition furnace). The component proportions will be based on Composite #1.

5.3.1.2 Samples for Fractured Coarse Aggregate Particles, Percent Carbonates (if required), and Abrasion (each separate source of virgin aggregate and RAP aggregate) will be prepared using Batch Sheet #1.

5.3.1.3 Examples of Composite #1 and corresponding Batch Sheet #1 are shown in Figures 1 and 2, respectively.

5.3.2 Batch Sheet #2:

5.3.2.1 This batch sheet will utilize the individual virgin aggregate gradations. The individual virgin stockpile proportions will be based on Composite #2. The percentage of material from each virgin aggregate stockpile is adjusted, as shown in Equations 18 and 19 (prior to Figure 3), so that the sum of the virgin aggregate stockpile percentages equal 100 percent.

5.3.2.2 Samples for Coarse and Fine Aggregate Specific Gravity and Absorption, Sand Equivalent, and Uncompacted Void Content (when required) will be prepared using Batch Sheet #2. In addition, minus No. 8 virgin aggregate make-up material is prepared using Batch Sheet #2.

5.3.2.3 Examples of Composite #2 and corresponding Batch Sheet #2 are shown in Figures 3 and 4, respectively.

5.3.3 Batch Sheet #3:

5.3.3.1 This batch sheet will utilize the individual virgin aggregate gradations and the gradation of the dry screened RAP material from each stockpile. The individual component stockpile proportions will be based on Composite #3. The percent of RAP material from each stockpile is adjusted to account for the RAP binder contained in the RAP material, as shown in Section 18. Marshall, Maximum Theoretical Specific Gravity (Rice Test), and Immersion Compression (IMC) samples will be prepared using Batch Sheet #3. Also using Batch Sheet #3, samples of dry screened RAP material are prepared for performing the Rice Test for each individual RAP stockpile. (See Subsection 6.3.1.4.)
5.3.3.2 Examples of Composite #3 and corresponding Batch Sheet #3 are shown in Figures 5 and 6, respectively.

5.3.3.3 The virgin aggregate samples for the Marshall, Rice, and Immersion Compression (IMC) specimens are batched separately from the RAP. Add the required percentage of mineral admixture to the virgin aggregate samples, by dry weight of the total aggregate (virgin aggregate and RAP aggregate), and mix thoroughly. Add 3% water, by dry weight of virgin aggregate, to each batched virgin aggregate/mineral admixture sample and mix thoroughly to wet the mineral admixture and virgin aggregate surfaces.

5.3.3.3.1 After mixing, place the wet treated virgin aggregate in a suitable pan, form a shallow crater in the center of the aggregate, and place the RAP material from each stockpile in the crater (do not allow the RAP material to touch the pan).

5.3.3.4 Testing utilizing the prepared virgin aggregate-RAP-mineral admixture samples for Marshall, Rice, and Immersion Compression will be performed as specified in Sections 9-11, 12, and 14, respectively.

5.3.4 Table 2 summarizes the test samples required for each batch sheet.

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Sample Size</th>
<th>Number of Samples</th>
<th>Batch Sheet #</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fractured Coarse Aggregate Particles (ARIZ 212)</td>
<td>Required grams of Virgin Mineral Aggregate and RAP Aggregate as determined by test method [No mineral admixture]</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Percent Carbonates (ARIZ 238)</td>
<td>300 grams of Virgin Mineral Aggregate and RAP Aggregate [No mineral admixture]</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Abrasion (AASHTO T 96)</td>
<td>5000 grams of Virgin Mineral Aggregate as per test method for grading type [No mineral admixture]</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Abrasion (AASHTO T 96)</td>
<td>5000 grams of RAP Aggregate as per test method for grading type [No mineral admixture]</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Fine Aggregate Specific Gravity/Absorption (ARIZ 211)</td>
<td>1200 grams of Virgin Mineral Aggregate [No mineral admixture]</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Table 2 (Continued)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coarse Aggregate Specific Gravity/Absorption (ARIZ 210)</td>
<td>Required grams of Virgin Mineral Aggregate as determined by the</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nominal Maximum Aggregate Size</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[No mineral admixture]</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sand Equivalent (AASHTO T 176)</td>
<td>500 to 750 grams of Virgin Mineral Aggregate</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[No mineral admixture]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Uncompacted Void Content (ARIZ 247)</td>
<td>Minimum 500 grams of Virgin Mineral Aggregate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(When required)</td>
<td>[No mineral admixture]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minus No. 8 Make-Up Material</td>
<td>An adequate amount (normally 500 grams) of Virgin Mineral Aggregate</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[No mineral admixture]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum Theoretical Specific Gravity (Rice Test) (ARIZ 806, as modified in Section 12)</td>
<td>3000 grams of Virgin Mineral Aggregate and RAP Aggregate</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(See Note 3.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Plus mineral admixture, by percent required by ARIZ 802 (as modified in Section 14)]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Yields 3 test specimens]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum Theoretical Specific Gravity (Rice Test) [ARIZ 417, as modified in Subsection 6.3.1.4]</td>
<td>3150 grams of Dry Screened RAP Material</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[No mineral admixture]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(for each RAP stockpile)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Yields 3 test specimens]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Marshall Density-Stability/Flow (ARIZ 415 and ARIZ 410, as modified in Sections 9 through 11)</td>
<td>3300 grams of Virgin Mineral Aggregate and RAP Aggregate</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(See Notes 3 and 4.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Plus mineral admixture, by percent required by ARIZ 802 (as modified in Section 14)]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(See Note 5.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Each sample yields 1 set of 3 Marshall Specimens]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immersion Compression (ARIZ 802, as modified in Section 14)</td>
<td>3400 grams of Virgin Mineral Aggregate and RAP Aggregate</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(See Notes 3 and 4.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Plus required percent of mineral admixture]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[Each sample yields 1 set of 2 IMC Specimens]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 2 (Continued)

| Note 3: | Samples are prepared with Virgin Mineral Aggregate and Dry Screened RAP material, however the amount shown for each sample is actually the amount of Virgin Mineral Aggregate and RAP Aggregate in the sample. Due to the asphalt binder in the RAP material, the required amount of RAP material necessary to provide the required samples of Virgin Mineral Aggregate and RAP Aggregate will be greater than that shown, as illustrated in Figure 6. The adjusted percent of RAP material from each stockpile to be used is determined as shown in Section 18. |

| Note 4: | Generally the weight of virgin aggregate and RAP shown will provide specimens of acceptable heights, but adjustments may be necessary in some cases. Use Equation 2 to adjust sample size as necessary to conform to specimen height requirements of 2.500 ± 0.200 inches for Marshall Density-Stability/Flow specimens and 4.000 ± 0.100 inches for IMC specimens. |

| Equation 2: | \[
\text{Adjusted Sample Size} = \frac{\text{Combined Bulk O.D.}}{2.650} \times \begin{cases} 3300 \text{ grams (for Marshall Density-Stability/Flow)} \\ 3400 \text{ grams (for Immersion Compression)} \end{cases}
\]

| Note 5: | Requires one (1) sample for each asphalt binder content to be tested (minimum of 3 binder contents, with 3 Marshall specimens at each binder content). |

6. **VIRGIN AGGREGATE AND RAP AGGREGATE SPECIFIC GRAVITIES AND ABSORPTION**

6.1 Determine the Bulk Oven Dry, S.S.D., Apparent Specific Gravities and Absorption for the virgin fine aggregate (minus No. 4) and the virgin coarse aggregate (plus No. 4) in accordance with ARIZ 211 and ARIZ 210, respectively.

6.2 Determine the combined bulk oven dry specific gravity of the virgin fine aggregate and the virgin coarse aggregate without mineral admixture and also the combined water absorption of the virgin fine aggregate and the virgin coarse aggregate without mineral admixture, in accordance with ARIZ 251. The composite of virgin aggregate materials is determined by Composite #2.

6.2.1 The combined virgin aggregate bulk oven dry specific gravity without mineral admixture and the combined virgin aggregate water absorption
without mineral admixture are used to determine compliance with specification requirements.

6.2.2 The combined virgin aggregate water absorption without mineral admixture is recorded as "A_v" and used in Equations 4b, 4c, and 6.

6.3 Estimate the Bulk Oven Dry Specific Gravity of the RAP aggregate from each RAP stockpile as follows.

6.3.1 For each RAP stockpile, the following items are used in the calculations.

6.3.1.1 The estimated binder absorption of the RAP aggregate.

**Note 6:** The binder absorption of the RAP aggregate \( (P_{bar}) \) is normally estimated to be 0.50 percent. An exception is made when the binder content of the RAP material is less than 1.00 percent, in which case the binder absorption of the RAP aggregate is estimated to be one-half of the binder content of the RAP material.

6.3.1.2 The tested specific gravity of the recovered RAP binder (when >15% RAP binder is used) or an estimated specific gravity of 1.050 for the RAP binder (when ≤15% RAP binder is used).

6.3.1.3 The binder content of the RAP material.

6.3.1.4 The maximum theoretical specific gravity and the effective specific gravity of the RAP aggregate determined in accordance with ARIZ 417, modified as follows:

6.3.1.4.1 From each RAP stockpile, weigh up a 3150 gram sample of dry screened material for individual No. 8 and larger sieves, and minus No. 8 material per the "As-Received" gradation. For an example, see the right side of Figure 6. Each 3000 gram sample is split into three individual approximate 1000 gram samples. Alternatively, individual 1000 gram samples may be weighed up separately.

6.3.1.4.2 Spread each sample in a large pan at nominal single stone thickness.

6.3.1.4.3 Place the sample in an oven at 230 ± 5 °F for 30 ± 5 minutes.

6.3.1.4.4 Immediately upon removal from the oven, break up fine particle agglomerations so that they are not larger than 1/4-inch, taking care not to fracture the aggregate particles.
6.3.1.4.5 Care must be taken not to lose minus No. 200 material while removing the sample from the flask following the evacuation of air from the sample. A recovery container must be provided to capture the water and minus No. 200 material while the contents of the flask are poured into a nest of No. 40 and No. 200 sieves.

6.3.1.4.6 After the minus No. 200 material has settled in the recovery container, carefully decant the water so that no minus No. 200 material is lost.

6.3.1.4.7 Dry the recovered minus No. 200 material to constant weight at a temperature that will not cause material to be lost due to splattering. Determine and record the weight of the recovered minus No. 200 material to the nearest 0.1 gram.

6.3.1.4.8 Combine the material retained on the No. 40 and No. 200 sieves and perform fan drying.

6.3.1.4.9 The weight of the recovered minus No. 200 material is added to the $W_{sd}$ weight.

6.3.1.4.10 Using Equation 3, calculate the effective specific gravity of each RAP aggregate ($G_{ser}$).

\[
G_{ser} = \frac{100 - P_{brn}}{100 - \left( \frac{P_{brn}}{G_{mmr}} \right)} - \frac{G_{brn}}{G_{brn}}
\]

Where:  
- $G_{ser}$ = Effective specific gravity of each RAP aggregate.
- $P_{brn}$ = Binder content of material from each RAP stockpile.
- $G_{mmr}$ = Maximum theoretical specific gravity of material from each RAP stockpile.
- $G_{brn}$ = When >15% RAP binder is used, the tested specific gravity of the recovered RAP binder from each RAP stockpile. Alternatively, blended RAP binder from each RAP stockpile may be tested. (See Note 9.) When ≤15% RAP binder is used, an estimated specific gravity of 1.050 is used for the RAP binder.

Example [for RAP Aggregate 1 (Fine RAP)]:

- $P_{br1} = 5.82$
- $G_{mmr} = 2.469$
- $G_{br1} = 1.049$
Example [for RAP Aggregate 2 (Coarse RAP)]:

\[
G_{sbr} = \frac{100 - 5.82}{100 - 5.82} = 2.694
\]

\[
= \frac{2.469 - 1.049}{2.431 - 1.052} = 2.431
\]

6.3.2 Using Equation 4a, calculate the Bulk Oven Dry Specific Gravity (G_{sbr}) of each RAP aggregate as a function of the estimated Binder Absorption of the RAP aggregate and the calculated Effective Specific Gravity.

**Equation 4a:**

\[
G_{sbr} = \frac{G_{ser}}{P_{bar} \times G_{ser} + 1}
\]

Where: 
- \( G_{sbr} \) = Bulk oven dry specific gravity of each RAP aggregate.
- \( P_{bar} \) = Estimated binder absorption of the RAP aggregate.
(See Note 6.)

Example [for RAP Aggregate 1 (Fine RAP)]:

\[
G_{ser} = 2.694
\]
\[
P_{bar} = 0.50
\]
\[
G_{br1} = 1.049
\]

\[
G_{sbr} = \frac{2.694}{0.50 \times 2.694 + 1} = 2.660
\]

Example [for RAP Aggregate 2 (Coarse RAP)]:

\[
G_{ser} = 2.431
\]
\[
P_{bar} = 0.50
\]
\[
G_{br2} = 1.052
\]
6.4 Using Equation 4b, calculate the corresponding estimated S.S.D. specific gravity of each RAP aggregate.

\[
G_{sbr} = \frac{2.431}{0.50 \times 2.431 + 1} = 2.403
\]

**Equation 4b:**

\[
\left[\text{S.S.D. Specific Gravity of each RAP aggregate}\right] = G_{sbr} \times \left(1 + \frac{A_v}{100}\right)
\]

Where: \(G_{sbr}\) = Bulk oven dry specific gravity of each RAP aggregate.

\(A_v\) = Combined virgin aggregate water absorption without mineral admixture.

**Note 7:** The water absorption of the RAP aggregates is assumed to be the same as the combined virgin aggregate water absorption.

Example [for RAP Aggregate 1 (Fine RAP)]:

\[
G_{sbr} = 2.660
\]

\[
A_v = 1.13
\]

\[
\left[\text{S.S.D. Specific Gravity of RAP aggregate 1 (Fine RAP)}\right] = 2.660 \times \left(1 + \frac{1.13}{100}\right) = 2.690
\]

Example [for RAP Aggregate 2 (Coarse RAP)]:

\[
G_{sbr} = 2.403
\]

\[
A_v = 1.13
\]

\[
\left[\text{S.S.D. Specific Gravity of RAP aggregate 2 (Coarse RAP)}\right] = 2.403 \times \left(1 + \frac{1.13}{100}\right) = 2.430
\]

6.5 Using Equation 4c, calculate the corresponding estimated Apparent specific gravity of each RAP aggregate.

\[
\left[\text{Apparent Specific Gravity of each RAP aggregate}\right] = \frac{1}{\frac{1}{G_{sbr}} - \frac{A_v}{100}}
\]
Where: $G_{sbr}$ = Bulk oven dry specific gravity of each RAP aggregate.
$A_v$ = Combined virgin aggregate water absorption without mineral admixture. (See Note 7.)

Example [for RAP Aggregate 1 (Fine RAP)]:

$$
G_{sbr} = 2.660 \\
A_v = 1.13
$$

$$
\left[\frac{\text{Apparent Specific Gravity of RAP aggregate 1 (Fine RAP)}}{1} \right] = \frac{1}{1 - \frac{1.13}{2.660}} = 2.742
$$

Example [for RAP Aggregate 2 (Coarse RAP)]:

$$
G_{sbr} = 2.403 \\
A_v = 1.13
$$

$$
\left[\frac{\text{Apparent Specific Gravity of RAP aggregate 2 (Coarse RAP)}}{1} \right] = \frac{1}{1 - \frac{1.13}{2.403}} = 2.470
$$

6.6 Using Equation 5, calculate the combined Bulk Oven Dry ($G_{sb}$) specific gravity, combined S.S.D. specific gravity, and combined Apparent specific gravity of the virgin aggregate, RAP aggregate, and mineral admixture.

**Equation 5:**

$$
\left[\frac{\text{Combined Specific Gravity of Virgin Aggregate, RAP Aggregate, and Mineral Admixture}}{1} \right] = \frac{P_f + P_c + P_{ra1} + P_{ra2} + P_{admix}}{G_f + G_{c} + G_{ra1} + G_{ra2} + G_{admix}}
$$

Where: $P_f$ = Weight percent of total virgin fine aggregate (minus No. 4), determined from Composite #1. [Calculated by multiplying the percentage of each virgin aggregate material used in the composite by the respective percent passing No. 4, and summing the individual results.]
\[ P_c = \text{Weight percent of total virgin coarse aggregate (plus No. 4), determined from Composite #1. [Calculated by subtracting } P_f \text{ from the total percent virgin aggregate used in the composite.]} \]

\[ P_{ra1}, P_{ra2} = \text{Weight percent of each RAP aggregate, determined from Composite #1.} \]

\[ P_{admix} = \text{Percent mineral admixture by weight of the aggregate.} \]

\[ P_f + P_c + P_{ra1} + P_{ra2} = 100 \]

\[ P_f + P_c + P_{ra1} + P_{ra2} + P_{admix} = 100 + \% \text{ Mineral Admixture} \]

\[ G_f, G_c, G_{ra1}, G_{ra2} = \text{Bulk Oven Dry, SSD, or Apparent specific gravity of the virgin fine aggregate, virgin coarse aggregate, and RAP aggregate(s) respectively.} \]

\[ G_{admix} = \text{Specific gravity of the mineral admixture.} \]

- Type I or II Cement = 3.14
- Type IP Cement = 3.00
- Hydrated Lime = 2.20

Example [for combined Bulk Oven Dry Specific Gravity (G_{sb})]:

\[ G_f = 2.797 \]
\[ G_c = 2.789 \]
\[ G_{ra1} = G_{sbr} \text{ [for RAP Aggregate 1 (Fine RAP)]} = 2.660 \]
\[ G_{ra2} = G_{sbr} \text{ [for RAP Aggregate 2 (Coarse RAP)]} = 2.403 \]
\[ G_{admix} = 3.00 \text{ (Type IP Cement)} \]
\[ P_f = (0.35 \times 97.1) + (0.13 \times 98.8) + (0.08 \times 22.5) + (0.19 \times 1.6) = 48.93 \]
\[ P_c = (75 – 48.93) = 26.07 \]
\[ P_{ra1} = 15.0 \text{ [for RAP Aggregate 1 (Fine RAP)]} \]
\[ P_{ra2} = 10.0 \text{ [for RAP Aggregate 2 (Coarse RAP)]} \]
\[ P_{admix} = 1.0 \]

Example of Combined Bulk Oven Dry Specific Gravity (G_{sb}) of Virgin Aggregate, RAP Aggregate, and Mineral Admixture:

\[
\begin{bmatrix}
\text{Example of Combined Bulk Oven Dry Specific Gravity (G}_{sb}\text{) of Virgin Aggregate, RAP Aggregate, and Mineral Admixture} \\
\text{= } & \\
\frac{48.93 + 26.07 + 15.0 + 10.0 + 1.0}{48.93 + 26.07 + 15.0 + 10.0 + 1.0} & = 2.732 \\
\text{= } & \\
\frac{2.797 + 2.789 + 2.660 + 2.403 + 3.00}{2.797 + 2.789 + 2.660 + 2.403 + 3.00} & = 2.732 \\
\end{bmatrix}
\]

6.7 Using Equation 6, calculate the combined water absorption of the virgin aggregate, RAP aggregate, and mineral admixture.
**Equation 6:**

\[
\text{Combined Water Absorption of Virgin Aggregate, RAP Aggregate, and Mineral Admixture} = \frac{(P_f \times A_f) + (P_c \times A_c) + (P_{ra} \times A_v) + (P_{admix} \times A_{admix})}{P_f + P_c + P_{ra} + P_{admix}}
\]

Where: $P_f, P_c,$ = Weight percent of virgin fine aggregate (minus No. 4) and virgin coarse aggregate (plus No. 4), respectively, as determined for $P_f$ and $P_c$ for use in Equation 5.

$P_{ra} =$ Total percent of RAP aggregate (Determined from Composite #1).

$A_v =$ Combined virgin aggregate water absorption without mineral admixture. (See Note 7.)

$P_{admix} =$ Percent mineral admixture by weight of the aggregate.

$P_f + P_c + P_{ra} = 100$

$P_f + P_c + P_{ra} + P_{admix} = 100 + \% \text{Mineral Admixture}$

$A_f, A_c =$ Percent water absorption of the virgin fine aggregate and the virgin coarse aggregate respectively.

$A_{admix} =$ Percent water absorption of mineral admixture (assumed to be 0.0%).

Example:

\[
\begin{align*}
P_f &= 48.93 \\
P_c &= 26.07 \\
A_f &= 1.11 \\
A_c &= 1.17 \\
P_{ra} &= 25.0 \\
A_v &= 1.13 \\
P_{admix} &= 1.0 \\
A_{admix} &= 0.00
\end{align*}
\]
7. ASPHALT BINDER SELECTION

7.1 Estimate the total binder content that will be needed in the mix design and determine whether the RAP binder content is lower or higher than the estimated total binder content.

Note 8: Estimating the total binder content relative to the RAP binder content may assist the designer in estimating the amount of RAP material which can be used in the mix.

7.2 For mixes utilizing >15% RAP binder, the extract obtained from the solvent extraction of material from each RAP stockpile, performed in Subsection 5.2.1.3, is used for additional testing. Determine and remove mineral matter from the extract using AASHTO T 164, Annex Method A1.2 (Centrifuge).

7.3 For mixes utilizing ≤15% RAP binder, the virgin binder used in the mix design shall be the grade called for in the Special Provisions. Determine the specific gravity [AASHTO T 228 (at 77 °F)] of the virgin binder.

7.4 For mixes utilizing >15% RAP binder, the virgin binder used in the mix design will be the grade determined as follows:

7.4.1 Recover the RAP binder from the extract solvent from each RAP stockpile using either AASHTO T 170 or ASTM D 5404. The solvent used in the extraction process shall not contain stabilizers that act as acid scavengers. The entire procedure, from the start of the extraction to the final recovery, must be completed within eight hours.

7.4.2 Perform PG grading (AASHTO R 29) on the recovered RAP binder from each RAP stockpile. Rotational viscosity, flash point, and mass loss tests are not required. Use PAV aging temperature for grade specified in the Special Provisions. Determine the specific gravity [AASHTO T 228 (at 77 °F)] of the recovered RAP binder from each RAP stockpile.
Note 9: Alternatively, the recovered binder from each RAP stockpile may be proportionally blended and the PG grading and specific gravity determined on the blended RAP binder.

7.4.3 Perform PG verification (AASHTO R 29) and determine the specific gravity [AASHTO T 228 (at 77 °F)] of the virgin binder, or use PG verification and specific gravity provided by the virgin asphalt binder supplier.

7.4.4 Blend RAP binder and virgin binder using the following procedure.

7.4.4.1 Using Equation 7, calculate the percent of RAP binder to blend with virgin binder. The percent of RAP binder, by weight of the total binder in the mix, shall not exceed the allowable maximum specified.

\[
P_{rb} = \left( \frac{P_{ra1}}{100 - P_{br1}} \times P_{br1} \right) + \left( \frac{P_{ra2}}{100 - P_{br2}} \times P_{br2} \right) \times 100 \\
\text{Equation 7:}
\]

Where: 
- \( P_{rb} \) = Percent of RAP Binder to blend with Virgin binder (Percent of RAP binder, by weight of total binder, in the mix).
- \( P_{ra1}, P_{ra2} \) = Percent of RAP aggregate from each RAP stockpile (from Composite #1).
- \( P_{br1}, P_{br2} \) = Binder content of the material from each RAP stockpile.
- \( P_{admix} \) = Percent mineral admixture used.
- \( P_b \) = Total asphalt binder content (percent by total weight of the bituminous mixture).

Note 10: The total asphalt binder content estimated in Subsection 7.1 is initially used for the value of \( P_b \) in Equations 7, 7a, and 7b. Subsequently, the actual total asphalt binder content from the mix design shall be used for the value of \( P_b \).
Example:

\[ \frac{15.0}{100 - 5.82} \times 5.82 + \frac{10.0}{100 - 3.43} \times 3.43 \]

\[ \frac{1 + \frac{1.0}{100}}{100 - 5.40} \times 5.40 \]

\[ P_{rb} = \frac{15.0}{100 - 5.82} \times 5.82 + \frac{10.0}{100 - 3.43} \times 3.43 \times 100 = 22.24\% \]

7.4.4.2 The proportional amount of RAP binder from each RAP stockpile can be calculated using Equations 7a and 7b.

**Equation 7a:**

\[ \frac{P_{r1}}{100} \left( \frac{100 - P_{br1}}{1 + \frac{P_{admix}}{100}} \right) \times P_{br1} \times 100 \]

Example (for Equation 7a) [for RAP Stockpile 1 (Fine RAP)]:

\[ \frac{15.0}{100 - 5.82} \times 5.82 \times 100 = 16.08\% \]

**Equation 7b:**

\[ \frac{P_{r2}}{100} \left( \frac{100 - P_{br2}}{1 + \frac{P_{admix}}{100}} \right) \times P_{br2} \times 100 \]

Example (for Equation 7b) [for RAP Stockpile 2 (Coarse RAP)]:
Example (for Equation 7b) [for RAP Stockpile 2 (Coarse RAP)]:

\[
P_{rb2} = \frac{10.0}{100} - \frac{3.43}{100 - 5.4} \times 3.43 \times 100 = 6.16\%
\]

7.4.4.3 Blend the required amount of RAP binder (proportional amounts from each RAP stockpile if necessary) with the required amount of virgin binder. The RAP binder and the virgin binder shall be heated sufficiently to pour and thoroughly stirred prior to blending. The blended material shall be thoroughly stirred prior to testing.

7.4.5 Perform PG verification (AASHTO R 29) on the blend of RAP and virgin binder. Flash point and mass loss tests are not required.

Note 11: If the total binder content, estimated in Subsection 7.1, deviates by more than 0.30 percent from the total binder content established by mix design, the PG grading must be verified on a binder blended at the blend percentages established by the mix design.

7.4.6 If the blended binder does not meet the specified grade, a different virgin binder grade, or a different proportion of RAP and virgin binder, must be selected. An iterative process may be required in order to obtain a final blend of RAP and virgin binder that meets the specified binder grade.

Note 12: The virgin binder grade may only be changed one PG grade (6 °C on either or both the high and low temperatures) from that shown in the bid documents.

7.4.7 Determine the specific gravity [AASHTO T 228 (at 77 °F)] of the blended binder. Develop a viscosity - temperature curve (ASTM D 2493) for the blended binder.

8. LABORATORY MIXING AND COMPACITION TEMPERATURES

8.1 For mixes containing ≤15% RAP binder, the laboratory mixing and compaction temperatures shall be determined based on the specified virgin binder utilized. These temperatures are determined in accordance with Subsections 8.1.1 through 8.1.4 below.
8.1.1 The rotational viscosity of the asphalt binder at 275 °F and 350 °F shall be determined in accordance with AASHTO T 316, and a viscosity-temperature curve developed in accordance with ASTM D 2493.

8.1.2 The laboratory mixing temperature range is defined as the range of temperatures where the un-aged asphalt binder has a rotational viscosity of 0.17 ± 0.02 Pascal·seconds. The actual laboratory mixing temperature used is normally selected at or near the mid-point of the range.

8.1.3 The laboratory compaction temperature range is defined as the range of temperatures where the un-aged asphalt binder has a rotational viscosity of 0.28 ± 0.03 Pascal·seconds. The actual laboratory compaction temperature used is normally selected at or near the mid-point of the range.

8.1.4 The viscosity-temperature curve shall be included in the mix design report. For PG asphalt binders that have a maximum laboratory mixing temperature exceeding 325 °F or a maximum laboratory compaction temperature exceeding 300 °F, the laboratory mixing and compaction temperature ranges shall be specified in writing by the asphalt binder supplier. A viscosity-temperature curve will meet this requirement for written documentation if the viscosity-temperature curve is developed and submitted by the binder supplier and includes language that the recommended laboratory mixing and compaction temperatures are within acceptable ranges, and shall include a statement indicating the maximum laboratory mixing temperature to which the binder can be heated without damage. The laboratory mixing and compaction temperature ranges, as well as the actual laboratory mixing and compaction temperatures used, shall be reported on the mix design.

8.2 For mixes containing >15% RAP binder, the laboratory mixing and compaction temperatures shall be determined from a viscosity-temperature curve developed for the blended binder. These temperatures are determined in accordance with Subsections 8.2.1 through 8.2.5 below.

8.2.1 The rotational viscosity of the blended binder at 275 °F and 350 °F shall be determined in accordance with AASHTO T 316, and a viscosity-temperature curve developed in accordance with ASTM D 2493.

8.2.2 Determine the laboratory mixing temperature range and the laboratory compaction temperature range, along with the actual laboratory mixing
and compaction temperatures, in accordance with Subsections 8.1.2 and 8.1.3.

8.2.3 If the maximum laboratory mixing temperature exceeds 350 °F, a temperature of 350 °F shall be used.

8.2.4 If the maximum laboratory compaction temperature exceeds 325 °F, a temperature of 325 °F shall be used.

8.2.5 The viscosity-temperature curve shall be included in the mix design report. The laboratory mixing and compaction temperature ranges, as well as the actual laboratory mixing and compaction temperatures used, shall be reported on the mix design.

9. PREPARATION OF SPECIMENS FOR BULK SPECIFIC GRAVITY/BULK DENSITY AND MARSHALL STABILITY/FLOW DETERMINATION

9.1 Specimens shall be prepared as described herein, utilizing the apparatus specified in ARIZ 410.

Note 13: A minimum of three different asphalt binder contents at 0.5% increments is required, although in some cases it may be necessary to prepare additional sets of samples at other asphalt binder contents.

9.2 The virgin aggregate-RAP-mineral admixture samples, prepared as specified in Section 5, shall be dried to constant weight at the laboratory mixing temperature ± 5 °F and shall be at this temperature at the time of mixing with the virgin asphalt binder. If necessary, a small amount of proportioned minus No. 8 virgin aggregate make-up material shall be added to bring samples to the desired weight.

9.3 Before each batch of asphaltic concrete is mixed, the virgin asphalt binder shall be heated in a loosely covered container in a forced draft oven for approximately 2 hours or as necessary to bring the asphalt binder to the laboratory mixing temperature ± 5 °F.

9.4 The weight of virgin asphalt binder required to provide the desired total binder content shall be calculated as follows.

9.4.1 The total binder content of a mix design sample is the total of the added virgin asphalt binder plus the RAP binder. Using Equations 8 through 10, calculate the weight of virgin asphalt binder ($W_{vb}$) to be added to the virgin aggregate-dry screened RAP-mineral admixture sample.
Equation 8: \[ W_b = \frac{W_s + W_a}{100 - P_b} \times P_b \]

Equation 9: (Calculated for each RAP stockpile)
\[ W_{rbn} = \frac{P_{ran}}{100 - P_{brn}} \times W_s \times P_{brn} \]

Equation 10: \[ W_{vb} = W_b - \left[ W_{rb1} + W_{rb2} \right] \]

Where:
- \( W_b \) = Weight of Total Binder (virgin asphalt binder + RAP binder).
- \( W_s \) = Weight of Total Aggregate (virgin aggregate + RAP aggregate).
- \( W_a \) = Weight of Mineral Admixture.
- \( P_b \) = Percent Total Binder (virgin asphalt binder + RAP binder) for each asphalt binder content, by total weight of mix.
- \( W_{rbn} \) = Weight of RAP binder for each RAP stockpile.
- \( P_{ran} \) = Percent of each RAP aggregate used (Determined from Composite #1).
- \( P_{brn} \) = Binder content of the material from each RAP stockpile.
- \( W_{vb} \) = Weight of virgin asphalt binder.

Example (for 5.0% asphalt binder content) (See Note 13.):
- \( W_s = 3300 \text{ g} \)
- \( W_a = 33 \text{ g} \)
- \( P_b = 5.0\% \)
- \( P_{ra1} = 15.0\% \)
- \( P_{ra2} = 10.0\% \)
- \( P_{br1} = 5.82\% \)
- \( P_{br2} = 3.43\% \)

\[ W_b = \frac{3300 + 33}{100 - 5.0} \times 5.0 = 175.4 \text{ grams} \]

For RAP Stockpile 1 (Fine RAP):
\[ W_{rb} = \frac{15.0}{100 - 5.82} \times 3300 \times 5.82 = 30.6 \text{ grams} \]
For RAP Stockpile 2 (Coarse RAP): \[ W_{vb} = \frac{10.0}{100} \times 3300 \times 3.43 = 11.7 \text{ grams} \]

\[ W_{vb} = 175.4 - [30.6 + 11.7] = 133.1 \text{ grams} \]

**Note 14:** Before each batch is mixed, the mixing bowl and whip shall be heated to the laboratory mixing temperature ± 5 ºF.

9.5 The virgin aggregate-RAP-mineral admixture blend and the appropriate amount of virgin asphalt binder shall be mixed together for 90 to 120 seconds at the required laboratory mixing temperature ± 5 ºF. **Mechanical mixing is required.**

**Note 15:** Although a wide range of mixers may provide the desired well-coated homogeneous mixture, commercial dough mixers with whips are often used. Minimum recommended capacity of the mixing bowl is 12 quarts.

**Note 16:** After mechanical mixing, hand mixing shall be used as necessary to produce a well-coated homogeneous mixture.

9.6 Immediately after mixing, each batch shall be thoroughly blended and spread according to ARIZ 416. The circular mass shall be cut into 6 equal pie-shaped segments. Take opposite segments for each individual specimen and use up the entire batch.

9.7 Each individual specimen shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The specimens shall be oven-cured for 2 hours ± 10 minutes at the laboratory compaction temperature ± 5 ºF.

9.8 A mold assembly (base plate, mold, and collar) shall be heated to the laboratory compaction temperature ± 5 ºF. The face of the compaction hammer shall be thoroughly cleaned and heated on a hot plate set at the laboratory compaction temperature ± 5 ºF. A suitable shield, baffle plate, or sand bath shall be used on the surface of the hot plate to minimize localized overheating of the face of the hammer.

9.9 Place a 4-inch diameter paper disc in the bottom of the mold before the mixture is introduced. Quickly place the mixture into the mold using a
transfer bowl or other suitable device. Spade the mixture vigorously with a heated flat metal spatula, with a blade approximately 1-inch wide and 6-inches long and stiff enough to penetrate the entire layer of material, 15 times around the perimeter and 10 times at random into the mixture, penetrating the mixture to the bottom of the mold. Smooth the surface of mix to a slightly rounded shape.

9.10 Before compaction, put the mold containing the mixture in an oven as necessary to heat the mixture to the laboratory compaction temperature ± 5 °F. The specimen shall not be heated for more than one hour after being placed in the mold. Prior to compaction, it shall be verified that the mixture is at the required temperature. An electronic temperature probe with a digital display or a long-stem metal thermometer with a dial face shall be used. The use of an infrared temperature gun is not allowed.

9.11 Immediately upon removing the mold assembly loaded with mix from the oven, place a paper disc on top of mixture, place the mold assembly on the compaction pedestal in the mold holder, and apply 75 blows with the compaction hammer. Remove the base plate and collar, and reverse and reassemble the mold. Apply 75 compaction blows to the face of the reversed specimen.

**Note 17:** The compaction hammer shall apply only one blow after each fall, that is, there shall not be a rebound impact. The compaction hammer shall meet the requirements specified in Subsection 2.4 of ARIZ 410.

9.12 Remove the collar and top paper disc. Remove the base plate and remove the bottom paper disc while the specimen is still hot.

**Note 18:** Paper discs need to be removed while the specimen is hot. The discs are difficult to remove after the specimens have cooled.

9.13 Allow each compacted specimen to cool until they can be extruded without damaging the specimen.

**Note 19:** Generally, specimens can be extruded without damage when they are at a temperature of approximately 77 to 90 °F.
Note 20: Cooling may be accomplished at room temperature, or in a 77 °F air bath. If more rapid cooling is desired, the mold and specimen may be placed in front of a fan until cool.

9.14 Extrude the specimen by orienting the mold so that the ram pushes on the bottom face (base plate face) of the specimen. If any specimen is deformed or damaged during extrusion, the entire set of specimens at that asphalt binder content shall be discarded and a new set prepared.

9.15 Immediately upon extrusion, measure and record the height of the specimen to the nearest 0.001 inch and determine and record its weight in air to the nearest 0.1 gram. Compacted specimens shall be 2.500 ± 0.200 inches in height. If this criteria is not met for the specimens at each asphalt binder content, the entire set of specimens at that asphalt binder content shall be discarded and a new set prepared after necessary adjustments in the sample size have been made using Equation 2. (See Note 4 in Table 2.)

9.16 Repeat the procedures in Subsections 9.4 through 9.15 for the required specimens.

10. BULK SPECIFIC GRAVITY/BULK DENSITY OF SPECIMENS

10.1 Determine the bulk specific gravity and bulk density of the three compacted specimens at each asphalt binder content in accordance with ARIZ 415, Method A. The determination of the “Weight in Water” and “S.S.D. Weight” of each specimen will be completed before the next specimen is submerged for its “Weight in Water” determination.

Note 21: Specimens fabricated in the laboratory that have not been exposed to moisture do not require drying after extrusion from the molds. The specimen weight in air obtained in Subsection 10.1 is its dry weight.

10.2 For each asphalt binder content, the densities of individual compacted specimens shall not differ by more than 2.0 pcf. If this density requirement is not met, the entire set of specimens at that asphalt binder content shall be discarded and a new set of specimens prepared.

10.3 For each set of three specimens, determine the average bulk specific gravity ($G_{mb}$), and using that value calculate the average bulk density ($G_{mb}$ multiplied by 62.3 lb./cu. ft.).
11. MARSHALL STABILITY AND FLOW DETERMINATION

11.1 Specimens shall be tested for Marshall Stability and Flow utilizing the apparatus specified in ARIZ 410.

11.2 For each specimen, determine the uncorrected stability, the stability (corrected for height), and the flow according to ARIZ 410, Subsections 4.6 through 4.11. In addition, determine the average stability (corrected for height) and average flow for each set of specimens prepared at each asphalt binder content.

12. MAXIMUM THEORETICAL SPECIFIC GRAVITY (RICE TEST)

12.1 The maximum theoretical specific gravity of the mixture shall be determined in accordance with ARIZ 806, with the following modifications:

12.1.1 Prepare the specimens including mineral admixture according to the procedures described in Section 5 and Subsections 9.2 through 9.5 using 5.5% or 6.0% total asphalt binder (virgin binder plus RAP binder) by total mix weight. A liquid anti-stripping agent is not used.

12.1.2 Immediately after mixing, the material shall be thoroughly blended and spread according to the procedures described in ARIZ 416. The circular mass shall be cut into 6 equal pie-shaped segments. Take opposite segments for each individual test sample and use up the entire batch.

12.1.3 Each individual test sample shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The test samples shall be oven-cured for 2 hours ± 10 minutes at the laboratory compaction temperature ± 5 °F.

12.1.4 After curing, each test sample shall be spread on a sheet of heavy paper or in a large flat bottom pan. Before the samples are completely cooled, separate the particles of the mixture, taking care not to fracture the mineral aggregate particles, so that the particles of the fine aggregate portion are not larger than ¼ inch. Allow the test samples to cool to room temperature.

12.1.5 Using Equation 11, calculate the effective specific gravity of the combined virgin aggregate, RAP aggregate, and mineral admixture ($G_{se}$).
Equation 11: \[ G_{se} = \frac{100 - P_b}{100} \frac{G_{mm} - G_b}{P_b} \]

Where:  
- \( G_{se} \) = Effective specific gravity of the combined virgin aggregate, RAP aggregate, and mineral admixture.  
- \( P_b \) = Total asphalt binder content (virgin binder + RAP binder) at which the Rice test was performed (Rice \( P_b \)).  
- \( G_{mm} \) = Measured maximum theoretical specific gravity of the mix at Rice \( P_b \).  
- \( G_b \) = When >15% RAP binder is used, the specific gravity of the blended virgin and RAP binder. (When \( \leq 15\% \) RAP binder is used, the specific gravity of the virgin binder is used for the value of \( G_b \).)

Example:

\[
\begin{align*}
P_b &= 6.0 \\
G_{mm} &= 2.526 \\
G_b &= 1.031
\end{align*}
\]

\[
G_{se} = \frac{100 - 6.0}{100} \frac{2.526 - 1.031}{6.0} = 2.784
\]

12.1.6 Using Equation 12, calculate the maximum theoretical specific gravity (\( G_{mm} \)) for each asphalt binder content.

Note 22: \( G_{se} \) is considered constant regardless of asphalt binder content.

Equation 12: \[ G_{mm} = \frac{100}{G_{se}} \frac{100 - P_b}{P_{sa}} \frac{P_b}{G_b} \]

Where:  
- \( G_{mm} \) = Calculated maximum theoretical specific gravity of the mix at each asphalt binder content \( P_b \).  
- \( P_{sa} \) = Aggregate and mineral admixture content, percent by total weight of mix (100-\( P_b \)).
Pb = Percent asphalt binder, by total weight of mix, for each asphalt binder content.

Gse = Effective specific gravity of the combined virgin aggregate, RAP aggregate, and mineral admixture.

Gb = When >15% RAP binder is used, the specific gravity of the blended virgin and RAP binder.
(When ≤15% RAP binder is used, the specific gravity of the virgin binder is used for the value of Gb.)

Example (for 5.0% asphalt binder content) (See Note 13.):

\[ P_{sa} = 95.0 \]
\[ P_b = 5.0 \]
\[ G_{se} = 2.784 \]
\[ G_b = 1.031 \]

\[ G_{mm} = \frac{100}{\frac{95.0}{2.784} + \frac{5.0}{1.031}} = 2.566 \]

13. DETERMINATION OF DESIGN ASPHALT BINDER CONTENT

13.1 The design percent asphalt binder content is determined as follows in Subsections 13.2 and 13.3.

13.2 For each asphalt binder content used, calculate effective voids (V_a), percent absorbed asphalt (P_{ba}), voids in mineral aggregate (VMA), percent effective asphalt binder content (P_{be}), and the ratio of the mix design composite gradation target for the No. 200 sieve, including mineral admixture, to the effective asphalt binder content (Dust/Bitumen Ratio) using the following equations.

13.2.1 Using Equation 13, calculate the effective voids (V_a). The calculated G_{mm} values for the respective asphalt binder contents are used to determine the corresponding effective voids content of the compacted Marshall specimens at each asphalt binder content.

**Equation 13:**

\[ V_a = \frac{G_{mm} - G_{mb}}{G_{mm}} \times 100 \]
Where: \( V_a \) = Effective voids in the compacted mixture, percent of total volume.

\( G_{mm} \) = Calculated maximum theoretical specific gravity of the mix at each asphalt binder content \( P_b \).

\( G_{mb} \) = Bulk specific gravity of compacted mixture specimens.

Example (for 5.0% asphalt binder content) (See Note 13.):

\[
G_{mm} = 2.566 \\
G_{mb} = 2.400 \\
V_a = \frac{2.566 - 2.400}{2.566} \times 100 = 6.5
\]

13.2.2 Using Equation 14, calculate the percent absorbed asphalt (\( P_{ba} \)).

**Note 23:** The percent absorbed asphalt (\( P_{ba} \)) is the same for all binder contents.

**Equation 14:**

\[
P_{ba} = \frac{G_{se} - G_{sb}}{G_{sb} \times G_{se}} \times G_b \times 100
\]

Where: \( P_{ba} \) = Absorbed asphalt, percent by total weight of mix.

\( G_{se} \) = Effective specific gravity of the combined virgin aggregate, RAP aggregate, and mineral admixture.

\( G_{sb} \) = Bulk oven dry specific gravity of the combined virgin aggregate, RAP aggregate, and mineral admixture.

\( G_b \) = When >15% RAP binder is used, the specific gravity of the blended virgin and RAP binder. (When \( \leq 15\% \) RAP binder is used, the specific gravity of the virgin binder is used for the value of \( G_b \).)

Example:

\[
G_{se} = 2.784 \\
G_{sb} = 2.732 \\
G_b = 1.031
\]
Using Equation 15, calculate voids in mineral aggregate (VMA) of the compacted Marshall specimens at each asphalt binder content.

**Equation 15:** \[ \text{VMA} = 100 - \frac{G_{mb} \times P_{sa}}{G_{sb}} \]

Where: 
- VMA = Voids in the mineral aggregate, percent of bulk volume.
- \( G_{mb} \) = Bulk specific gravity of compacted mixture specimens.
- \( P_{sa} \) = Aggregate and mineral admixture content, percent by total weight of mix (100-\( P_b \)).
- \( G_{sb} \) = Bulk oven dry specific gravity of the combined virgin aggregate, RAP aggregate, and mineral admixture.

Example (for 5.0% asphalt binder content) (See Note 13.):

\[
G_{mb} = 2.400 \\
P_{sa} = 95.0 \\
G_{sb} = 2.732
\]

\[ \text{VMA} = 100 - \frac{2.400 \times 95.0}{2.732} = 16.5 \]

Using Equation 16, calculate the percent effective asphalt binder content (\( P_{be} \)) of the compacted Marshall specimens at each asphalt binder content.

**Equation 16:** \[ P_{be} = P_b - \frac{P_{ba} \times P_{sa}}{100} \]

Where: 
- \( P_{be} \) = Percent effective asphalt binder content of the mixture (free binder not absorbed).
- \( P_b \) = Asphalt binder content, percent by total weight of mix.
- \( P_{ba} \) = Absorbed asphalt, percent by total weight of mix.
- \( P_{sa} \) = Aggregate and mineral admixture content, percent by total weight of mix (100-\( P_b \)).
Example (for 5.0% asphalt binder content) (See Note 13.):

\[
P_b = 5.0 \\
P_{ba} = 0.70 \\
P_{sa} = 95.0
\]

\[
P_{be} = 5.0 - \frac{0.70 \times 95.0}{100} = 4.34
\]

13.2.5 Using Equation 17, calculate the the ratio of the mix design composite gradation target for the No. 200 sieve, including mineral admixture, to the effective asphalt binder content (Dust/Bitumen Ratio) of the compacted Marshall specimens at each asphalt binder content.

**Equation 17:**

\[
\text{Dust / Bitumen Ratio} = \frac{\text{Mix Design}}{P_{be}}
\]

Where: \( P_{be} \) = Percent effective asphalt binder content of the mixture (free binder not absorbed).

Example (for 5.0% asphalt binder content) (See Note 13.):

\[
P_{be} = 4.34
\]

Dust / Bitumen Ratio = \( \frac{5.2}{4.34} = 1.20 \)

13.3 If the specified value for effective voids (\( V_a \)) falls between the resultant \( V_a \) values determined for two of the asphalt binder contents used, line straight-line interpolation shall be used to calculate the bulk specific gravity (\( G_{mb} \)) at each 0.1% asphalt binder content increment between the two corresponding measured values of \( G_{mb} \). Using equations 12 and 13, determine the values for \( G_{mm} \) and \( V_a \) at each interpolated asphalt binder content. Select the total asphalt binder content which results in \( V_a \) being as close as possible to the specified value.

13.3.1 Using equations 15, 16, and 17, calculate VMA, \( P_{be} \), and the Dust/Bitumen Ratio at the selected asphalt binder content. Using straight-line interpolation, determine the values for stability and flow that correspond to the selected asphalt binder content. If the values for VMA, Dust/Bitumen Ratio, stability, and flow are within the limits of the
specifications, the selected asphalt binder content shall be the design asphalt binder content.

13.3.2 On separate graphs (using the same scale for percent total asphalt binder for each graph), plot the values for bulk specific gravity, effective voids, VMA, Dust/Bitumen Ratio, stability, and flow for each set of three specimens and at the design asphalt binder content, versus the percent total asphalt binder. Draw a smooth “best fit” based on the plotted points, as shown in Figure 7. The graphs in Figure 7 are for illustrative purposes only. Straight-line interpolation, as specified in Subsections 13.3 and 13.3.1, is used for mix design calculations.

13.3.3 If it is not possible to obtain specification compliance within the range of asphalt binder contents used, a determination must be made to either redesign the mix (different aggregate gradation and/or source) or prepare additional specimens at other asphalt binder contents for testing and volumetric analyses.

14. IMMERSION COMPRESSION (IMC) TEST

14.1 The Immersion Compression Test shall be performed in accordance with ARIZ 802, modified as follows:

14.1.1 Prepare the asphaltic concrete sample including mineral admixture according to the procedures described in Section 5 and Subsections 9.2 through 9.5 at the design asphalt binder content.

14.1.2 Immediately after mixing, each batch shall be thoroughly blended and spread according to ARIZ 416. The circular mass shall be cut into 4 equal pie-shaped segments. Take opposite segments for each individual specimen and use up the entire batch.

14.1.3 Each individual sample shall be spread in a large pan at nominal single-stone thickness. Avoid stacking particles as feasible. The samples shall be oven-cured for 2 hours ± 10 minutes at 255 ± 5 ºF.

14.1.4 Compacted specimens shall be 4.000 ± 0.100 inches in height.
15. **MIX DESIGN GRADATION TARGET VALUES**

15.1 The target values for the combined virgin aggregate and RAP aggregate without mineral admixture (Composite #1), and the combined virgin aggregate-RAP aggregate-mineral admixture blend (Composite #1), in the mixture shall be expressed as percent passing particular sieve sizes as required by the specifications for the project.

16. **REPORT**

16.1 Report the test results and data obtained on the appropriate form. Liberal use of the remarks area to clarify and/or emphasize any element of the design is strongly recommended. Information required in the mix design report includes the items listed below. In addition, all composites and batch sheets/weigh-ups (examples are given in Figures 1 through 6) are to be included in the mix design report.

16.1.1 Name and address of the testing organization and the signature and seal of the mix design engineer responsible for the mix design.

16.1.2 Virgin Aggregate:
- Source and identification
- Individual stockpile or bin gradations
- Aggregate blend proportions and composite gradation with and without mineral admixture (See example in Figure 3.)
- Fine and coarse aggregate specific gravities (Bulk Oven Dry, SSD, Apparent) and absorption
- Combined Bulk Oven Dry Specific Gravity
- Combined Absorption
- Aggregate quality:
  - LA Abrasion
  - Sand Equivalent
  - Uncompacted Void Content (when applicable)

16.1.3 Mineral Admixture:
- Type and source
- Percentage used
- Specific gravity
16.1.4 RAP (each stockpile):
- Source and identification
- Anticipated gradation
- Binder content \((P_{bm})\)
- Maximum theoretical specific gravity

16.1.5 RAP Aggregate (each stockpile):
- Percentage used
- Gradation
- Estimated binder absorption \((P_{bar})\) (normally 0.50 percent) (See Note 6.)
- Estimated Bulk Oven Dry, S.S.D., and Apparent specific gravities
- Aggregate quality:
  - LA Abrasion

16.1.6 Combined Virgin Aggregate and RAP Aggregate:
- Composite gradation with and without mineral admixture (See example in Figure 1.)
- Aggregate quality:
  - Fractured Coarse Aggregate Particles (percentage with one fractured face and when required, percentage with two fractured faces)
  - Carbonates (when applicable)

16.1.7 Virgin Asphalt Binder:
- Source and grade
- Specific gravity (at 77 °F)
- Viscosity - Temperature Curve and the following:
  - Laboratory mixing temperature range and actual laboratory mixing temperature used.
  - Laboratory compaction temperature range and actual laboratory compaction temperature used.
- Virgin asphalt binder content (by weight of total binder and by weight of total mix)
- PG verification results (if RAP binder content is greater than 15 percent)

16.1.8 RAP Binder:
- RAP binder content (by weight of total binder and by weight of total mix)
- PG grading results (Standard PG Grade and Actual Grade), including specific gravity \((G_{bm})\) (at 77 °F), on recovered RAP binder from each RAP stockpile (if RAP binder content is greater than 15 percent)
16.1.9 Virgin Asphalt Binder and RAP Binder Blend:
   - Total Binder Content \( (P_b) \)
   - PG verification results, including specific gravity \( (G_b) \) (at 77 °F), (if RAP binder content is greater than 15 percent)

16.1.10 Combined Virgin Aggregate and Dry Screened RAP:
   - Individual stockpile or bin gradations
   - Percentage of virgin aggregates used
   - Percentage of RAP materials used (adjusted for RAP binder content)

16.1.11 Combined Aggregate (virgin aggregate and RAP aggregate) and Mineral Admixture:
   - Combined specific gravities [Bulk Oven Dry \( (G_{sb}) \), SSD, and Apparent]
   - Combined absorption

16.1.12 Maximum theoretical specific gravity \( (G_{mm}) \) and density (pcf) at the total asphalt binder content at which the Rice test was performed \( (Rice \ P_b) \).

16.1.13 Mixture Compaction Trials:
   - Percent virgin asphalt binder, by weight of total binder in the mix
   - Percent RAP binder, by weight of total binder in the mix
   - Total asphalt binder content \( (P_b) \), by weight of the bituminous mixture
   - Combined aggregate (virgin aggregate and RAP aggregate) and mineral admixture content \( (P_{sa}) \)
   - Calculated maximum theoretical specific gravity \( (G_{mm}) \) and density (pcf)
   - Bulk specific gravity \( (G_{mb}) \) and bulk density (pcf) of Marshall specimens
   - Percent effective voids \( (V_a) \)
   - Percent voids in mineral aggregate \( (VMA) \)
   - Dust/Bitumen Ratio
   - Percent absorbed asphalt \( (P_{ba}) \)
   - Effective specific gravity of the combined aggregate (virgin aggregate and RAP aggregate) and mineral admixture \( (G_{se}) \)
   - Percent effective asphalt binder content \( (P_{be}) \)
   - Marshall stability (nearest 10 pounds)
   - Marshall flow (0.01 inch)
16.1.14 Plots of the following properties versus percent total asphalt binder (See examples in Figure 7.):
- Percent effective voids ($V_a$)
- Percent voids in mineral aggregate (VMA)
- Dust/Bitumen Ratio
- Bulk specific gravity ($G_{mb}$)
- Marshall stability
- Marshall flow

16.1.15 Mixture Properties at Design Asphalt Binder Content:
- Percent virgin asphalt binder, by weight of total binder in the mix
- Percent RAP binder, by weight of total binder in the mix
- Total asphalt binder content ($P_b$), by weight of the bituminous mixture
- Combined aggregate (virgin aggregate and RAP aggregate) and mineral admixture content ($P_{sa}$)
- Calculated maximum theoretical specific gravity ($G_{mm}$) and density (pcf)
- Bulk specific gravity ($G_{mb}$) and bulk density (pcf) of Marshall specimens
- Percent effective voids ($V_a$)
- Percent voids in mineral aggregate (VMA)
- Dust/Bitumen Ratio
- Percent absorbed asphalt ($P_{ba}$)
- Effective specific gravity of the combined aggregate (virgin aggregate and RAP aggregate) and mineral admixture ($G_{se}$)
- Percent effective asphalt binder content ($P_{be}$)
- Marshall stability (nearest 10 pounds)
- Marshall flow (0.01 inch)
- Immersion Compression wet strength (psi)
- Immersion Compression dry strength (psi)
- Index of retained strength

16.1.16 Mix Design Summary [The Mix Design Summary shall contain the information shown in Figure 8. (Pages 1 and 2 of an example Mix Design Summary for an Asphaltic Concrete Marshall Mix Design with RAP.)]
## Example of Composite #1
**(Virgin Aggregate and RAP Aggregate)**

<table>
<thead>
<tr>
<th>Material</th>
<th>Virgin Aggregate (75.0%)</th>
<th>RAP Aggregate (25.0%)</th>
<th>Composite</th>
<th>Specifications ADOT 416 (3/4 Inch Special Mix)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Washed Crusher Fines</td>
<td>Crusher Fines 3/8&quot;</td>
<td>3/4&quot;</td>
<td>RAP 1 (Fine) RAP 2 (Coarse)</td>
</tr>
<tr>
<td>Percent used in composite</td>
<td>35.0 13.0 8.0 19.0 15.0 10.0</td>
<td>35.0 13.0 8.0 19.0 15.0 10.0</td>
<td>Without Admix  With Admix</td>
<td>Without Admix  With Admix</td>
</tr>
<tr>
<td>Sieve</td>
<td>Gradation – Percent Passing</td>
<td>Without Admix  With Admix</td>
<td>Without Admix  With Admix</td>
<td></td>
</tr>
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<td>1&quot;</td>
<td>100.0 100.0 100.0 100.0 100.0 100.0</td>
<td>100 100</td>
<td>100 100</td>
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</tr>
<tr>
<td>3/4&quot;</td>
<td>100.0 100.0 100.0 82.0 100.0 100.0</td>
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<td>90 - 100 90 - 100</td>
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<tr>
<td>1/2&quot;</td>
<td>100.0 100.0 100.0 13.6 100.0 63.0</td>
<td>80 80</td>
<td>62 - 77 62 - 77</td>
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</tr>
<tr>
<td>3/8&quot;</td>
<td>100.0 100.0 100.0 4.0 100.0 10.5</td>
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<td>62 - 77 62 - 77</td>
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</tr>
<tr>
<td>1/4&quot;</td>
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<td>68 68</td>
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</tr>
<tr>
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<td>97.1 98.8 22.5 1.6 77.6 6.0</td>
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<td>62 62</td>
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</tr>
<tr>
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<td>4.3 5.2</td>
<td>1.5 - 4.5 2.5 - 6.0</td>
<td></td>
</tr>
</tbody>
</table>

* 1.0 percent admixture, by weight of the total aggregate (virgin aggregate and RAP aggregate), is used in this example.

**FIGURE 1**
Example of Batch Sheet #1  
(For Composite #1)  
(Virgin Aggregate and RAP Aggregate)

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SIEVE</th>
<th>INDIV. % RET.</th>
<th>ACCUM. % RET.</th>
<th>FRACTURED COARSE AGGREGATE PARTICLES [Plus #4] (Virgin Agg. and RAP Agg.) (500 grams)</th>
<th>PERCENT CARBONATES [Plus #4] (Virgin Agg. and RAP Agg.) (300 grams)</th>
<th>ABRASION [Type B] (Virgin Agg.) (5000 grams)</th>
<th>ABRASION [Type B] (RAP Agg.) (5000 grams)</th>
<th>ACCUMULATIVE WEIGHT (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Virgin Aggregate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Washed Crusher Fines</td>
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<td>1.015</td>
<td>1.015</td>
<td>13</td>
<td>8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(35.0%)</td>
<td>#8</td>
<td>12.705</td>
<td>13.720</td>
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<td>35.000</td>
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<tr>
<td>Virgin Aggregate</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>35.156</td>
<td>15</td>
<td>9</td>
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<tr>
<td>(13.0%)</td>
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<td>3.731</td>
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<td></td>
</tr>
<tr>
<td>3/8 Inch (8.0%)</td>
<td>1/4&quot;</td>
<td>3.280</td>
<td>51.280</td>
<td>57</td>
<td>34</td>
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<td>57</td>
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<td>56.000</td>
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<td>Virgin Aggregate</td>
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<td></td>
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</tr>
<tr>
<td>3/4 Inch (19.0%)</td>
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<td>1.824</td>
<td>74.240</td>
<td>330</td>
<td>198</td>
<td>5000</td>
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<td>0.456</td>
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<td>RAP Aggregate 1</td>
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<td></td>
<td></td>
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<td></td>
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<tr>
<td>(Fine RAP) (15.0%)</td>
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<td>1.005</td>
<td>76.005</td>
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<td>209</td>
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<td>(Coarse RAP) (10.0%)</td>
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<td>93.700</td>
<td>427</td>
<td>256</td>
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<td>3/8&quot;</td>
<td>5.250</td>
<td>98.950</td>
<td>494</td>
<td>297</td>
<td>5000</td>
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<tr>
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<td>100.000</td>
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</tbody>
</table>

FIGURE 2
17. ADJUSTED PERCENTAGE FOR EACH VIRGIN AGGREGATE STOCKPILE

17.1 In order to develop the data for Composite #2, Equations 18 and 19 are used to calculate the adjusted percentage for each virgin aggregate stockpile, as shown in Figure 3.

Equation 18: \[ P_{sv} = P_{sv1} + P_{sv2} + ... + P_{svn} \]

Where: \( P_{sv} \) = Sum of the original individual virgin aggregate stockpile percentages from Composite #1.

\( P_{sv1}, P_{sv2}, P_{svn} \) = The original individual virgin aggregate stockpile percentages from Composite #1.

Example:

\[
\begin{align*}
P_{sv1} & = 35.0 \\
P_{sv2} & = 13.0 \\
P_{sv3} & = 8.0 \\
P_{sv4} & = 19.0 \\
P_{sv} & = 35.0 + 13.0 + 8.0 + 19.0 = 75.0
\end{align*}
\]

Equation 19: \[ P_{svcn} = \frac{P_{svn}}{P_{sv}} \times 100 \]

Where: \( P_{svcn} \) = Adjusted individual virgin aggregate stockpile percentages in Composite #2 (as a percentage of virgin aggregates only).

\( P_{sv} \) = Sum of the original individual virgin aggregate stockpile percentages from Composite #1.

Example: (for Washed Crusher Fines)

\[
\begin{align*}
P_{sv} & = 75.0 \\
P_{sv1} & = 35.0 \\
P_{svcn} & = \frac{35.0}{75.0} \times 100 = 46.667
\end{align*}
\]
Example: (for Crusher Fines)

\[ P_{sv} = 75.0 \]
\[ P_{sv2} = 13.0 \]

\[ P_{svc2} = \frac{13.0}{75.0} \times 100 = 17.333 \]

Example: (for 3/8 Inch)

\[ P_{sv} = 75.0 \]
\[ P_{sv3} = 8.0 \]

\[ P_{svc3} = \frac{8.0}{75.0} \times 100 = 10.677 \]

Example: (for 3/4 Inch)

\[ P_{sv} = 75.0 \]
\[ P_{sv4} = 19.0 \]

\[ P_{svc4} = \frac{19.0}{75.0} \times 100 = 25.333 \]
# Example of Composite #2
(Virgin Aggregate)

<table>
<thead>
<tr>
<th>Material</th>
<th>Washed</th>
<th>Crusher</th>
<th>3/8 Inch</th>
<th>3/4 inch</th>
<th>Composite</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fines</td>
<td>Fines</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Original individual virgin aggregate stockpile percentages used in Composite #1.</td>
<td>35.0</td>
<td>13.0</td>
<td>8.0</td>
<td>19.0</td>
<td>(Total = 75.0%)</td>
</tr>
<tr>
<td>Adjusted individual virgin aggregate stockpile percentages.</td>
<td>46.667</td>
<td>17.333</td>
<td>10.667</td>
<td>25.333</td>
<td>Without Admix With Admix *</td>
</tr>
<tr>
<td>(Total = 100%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sieve</th>
<th>Gradation – Percent Passing</th>
</tr>
</thead>
<tbody>
<tr>
<td>1&quot;</td>
<td>100.0 100.0 100.0 100.0 100 100</td>
</tr>
<tr>
<td>3/4&quot;</td>
<td>100.0 100.0 100.0 82.0 95 95</td>
</tr>
<tr>
<td>1/2&quot;</td>
<td>100.0 100.0 100.0 13.6 78 78</td>
</tr>
<tr>
<td>3/8&quot;</td>
<td>100.0 100.0 100.0 4.0 76 76</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>100.0 100.0 59.0 1.6 71 71</td>
</tr>
<tr>
<td>#4</td>
<td>97.1 98.8 22.5 1.6 65 65</td>
</tr>
<tr>
<td>#8</td>
<td>60.8 70.1 1.3 1.5 41 42</td>
</tr>
<tr>
<td>#10</td>
<td>58.8 68.1 1.2 1.5 40 41</td>
</tr>
<tr>
<td>#16</td>
<td>33.3 45.1 1.0 1.5 24 25</td>
</tr>
<tr>
<td>#30</td>
<td>19.4 31.3 1.0 1.5 15 16</td>
</tr>
<tr>
<td>#40</td>
<td>13.9 25.9 0.9 1.4 11 12</td>
</tr>
<tr>
<td>#50</td>
<td>10.0 22.0 0.9 1.4 9 10</td>
</tr>
<tr>
<td>#100</td>
<td>5.5 17.0 0.8 1.4 6 7</td>
</tr>
<tr>
<td>#200</td>
<td>3.3 8.4 0.7 1.2 3.4 4.7</td>
</tr>
</tbody>
</table>

* The weight of admixture used in the mix is based on the weight of the total aggregate (virgin aggregate and RAP aggregate). As shown in Composite #1, the percent admixture, by weight of the total aggregate, used in this example is 1.0 percent. However, since Composite #2 is for virgin aggregate only (but will contain all of the admixture), to correctly calculate the composite gradation with admixture the percent of admixture must be adjusted for the percent of RAP aggregate used in the mix (25.0% in this example). The adjusted percent of admixture is determined by dividing “100” by the percent of virgin aggregate (75.0 in this example). The resultant value (1.33 for this example) is the adjusted percent of admixture used to calculate the composite gradation with admixture.

\[
%\text{passing (with admix)} = \left(\frac{\% \text{passing without admix}}{100} + \text{adjusted percent of admixture}\right) \times 100
\]

Example for the No. 200 sieve:

\[
%\text{passing (with admix)} = \left[\frac{3.4 + 1.33}{100 + 1.33}\right] \times 100 = 4.7
\]

FIGURE 3
## Example of Batch Sheet #2 (For Composite #2) (Virgin Aggregate)

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SIEVE</th>
<th>INDIV. % RET.</th>
<th>ACCUM. % RET.</th>
<th>COARSE SPECIFIC GRAVITY (Plus #4) (4000 grams)</th>
<th>FINE SPECIFIC GRAVITY (Minus #4) (1200 grams)</th>
<th>SAND EQUIVALENT [Minus #4] (600 grams)</th>
<th>UNCOMPACTED VOID CONTENT [Minus #8] (500 grams)</th>
<th>MINUS NO. 8 MAKE-UP MATERIAL (500 grams)</th>
<th>ACCUMULATIVE WEIGHT (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Virgin Aggregate</td>
<td>#4</td>
<td>1.353</td>
<td>1.353</td>
<td>156</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Washed Crusher Fines (Adjusted = 46.667%)</td>
<td>#8</td>
<td>16.940</td>
<td>18.293</td>
<td>---</td>
<td>312</td>
<td>156</td>
<td>---</td>
<td>---</td>
<td>346</td>
</tr>
<tr>
<td>#8</td>
<td>28.374</td>
<td>46.667</td>
<td>---</td>
<td>833</td>
<td>417</td>
<td>346</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>#8</td>
<td>4.975</td>
<td>51.850</td>
<td>---</td>
<td>925</td>
<td>463</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>#8</td>
<td>12.150</td>
<td>64.000</td>
<td>---</td>
<td>1148</td>
<td>574</td>
<td>494</td>
<td>494</td>
<td>494</td>
<td></td>
</tr>
<tr>
<td>Virgin Aggregate</td>
<td>1/4&quot;</td>
<td>4.374</td>
<td>68.374</td>
<td>683</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Crusher Fines (Adjusted = 17.333%)</td>
<td>#4</td>
<td>3.893</td>
<td>72.267</td>
<td>1131</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>#8</td>
<td>2.261</td>
<td>74.528</td>
<td>---</td>
<td>1190</td>
<td>595</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>#8</td>
<td>0.139</td>
<td>74.667</td>
<td>---</td>
<td>1193</td>
<td>596</td>
<td>495</td>
<td>495</td>
<td>495</td>
<td>495</td>
</tr>
<tr>
<td></td>
<td>1/2&quot;</td>
<td>17.328</td>
<td>96.555</td>
<td>3650</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>3/8&quot;</td>
<td>2.432</td>
<td>98.987</td>
<td>3930</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>1/4&quot;</td>
<td>0.608</td>
<td>99.595</td>
<td>4000</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.025</td>
<td>99.620</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.380</td>
<td>100.000</td>
<td>---</td>
<td>1200</td>
<td>600</td>
<td>500</td>
<td>500</td>
<td>500</td>
</tr>
</tbody>
</table>

**FIGURE 4**
18. ADJUSTED PERCENTAGE FOR EACH RAP STOCKPILE

18.1 In order to develop the data for Composite #3, Equation 20 is used to calculate the adjusted percentage for each RAP stockpile, as shown in Figure 5.

\[ P_{rcn} = \frac{P_{ran}}{1 - \frac{P_{brn}}{100}} \]

Where:
- \( P_{rcn} \) = Adjusted individual RAP stockpile percentages (includes RAP binder).
- \( P_{ran} \) = Percent of each RAP aggregate used (Determined from Composite #1).
- \( P_{brn} \) = Binder content of the material from each RAP stockpile.

Example: For RAP Stockpile 1 (Fine RAP)

\[
P_{ra1} = 15.0 \\
P_{br1} = 5.82 \\
\]

\[
P_{rc1} = \frac{15.0}{1 - \frac{5.82}{100}} = 15.927
\]

Example: For RAP Stockpile 2 (Coarse RAP)

\[
P_{ra2} = 10.0 \\
P_{br2} = 3.43 \\
\]

\[
P_{rc2} = \frac{10.0}{1 - \frac{3.43}{100}} = 10.355
\]
### Example of Composite #3
**(Virgin Aggregate and Dry Screened RAP)**

<table>
<thead>
<tr>
<th>Material</th>
<th>Virgin Aggregate (75.0%)</th>
<th>Dry Screened RAP (26.282%)</th>
<th>Composite Without Admix **</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Washed Crusher Fines</td>
<td>Crusher Fines</td>
<td>RAP 1 (Fine)</td>
</tr>
<tr>
<td></td>
<td>3/8 Inch</td>
<td>3/4 inch</td>
<td>(% Binder = 5.82) *</td>
</tr>
<tr>
<td>Percent</td>
<td>35.0</td>
<td>13.0</td>
<td>8.0</td>
</tr>
<tr>
<td>used in</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>composite</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sieve</td>
<td>Gradation - Percent Passing</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1”</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>3/4”</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>1/2”</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>3/8”</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>1/4”</td>
<td>100.0</td>
<td>100.0</td>
<td>59.0</td>
</tr>
<tr>
<td>#4</td>
<td>97.1</td>
<td>98.8</td>
<td>22.5</td>
</tr>
<tr>
<td>#8</td>
<td>60.8</td>
<td>70.1</td>
<td>1.3</td>
</tr>
</tbody>
</table>

* By weight of Dry Screened RAP

** The composite gradation without admixture includes the contribution of binder in the Dry Screened RAP [as indicated by the total percent of Dry Screened RAP (26.282%), the total percent of Virgin Aggregate and Dry Screened RAP used in the composite (101.282%), and the percent passing the 1 inch sieve (101%)].
### Example of Batch Sheet #3
(For Composite #3)
(Virgin Aggregate and Dry Screened RAP)

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SIEVE</th>
<th>INDIVIDUAL PERCENT RETAINED</th>
<th>ACCUMULATIVE PERCENT RETAINED</th>
<th>RICE (3000 grams)</th>
<th>MARSHALL (3300 grams)</th>
<th>IMC (3400 grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Washed Crusher Fines (35%)</td>
<td>#4</td>
<td>1.015</td>
<td>1.015</td>
<td>30</td>
<td>33</td>
<td>35</td>
</tr>
<tr>
<td></td>
<td>#6</td>
<td>12.705</td>
<td>13.720</td>
<td>412</td>
<td>453</td>
<td>466</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>21.280</td>
<td>35.000</td>
<td>1050</td>
<td>1155</td>
<td>1190</td>
</tr>
<tr>
<td>Crusher Fines (13%)</td>
<td>#4</td>
<td>0.156</td>
<td>35.156</td>
<td>1055</td>
<td>1160</td>
<td>1195</td>
</tr>
<tr>
<td></td>
<td>#6</td>
<td>3.731</td>
<td>38.887</td>
<td>1167</td>
<td>1283</td>
<td>1322</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>9.113</td>
<td>48.000</td>
<td>1440</td>
<td>1584</td>
<td>1632</td>
</tr>
<tr>
<td>#4</td>
<td>2.290</td>
<td>51.280</td>
<td>1538</td>
<td>1692</td>
<td>1744</td>
<td></td>
</tr>
<tr>
<td>3/8 Inch (8%)</td>
<td>#6</td>
<td>1.696</td>
<td>55.896</td>
<td>1677</td>
<td>1845</td>
<td>1900</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.104</td>
<td>56.000</td>
<td>1680</td>
<td>1848</td>
<td>1904</td>
</tr>
<tr>
<td>1/4 Inch</td>
<td>3.260</td>
<td>59.520</td>
<td>1763</td>
<td>1911</td>
<td>2000</td>
<td></td>
</tr>
<tr>
<td>RAP 2</td>
<td>3/8 Inch (19%)</td>
<td>1.2996</td>
<td>72.416</td>
<td>2172</td>
<td>2350</td>
<td>2462</td>
</tr>
<tr>
<td></td>
<td>3/4 Inch</td>
<td>1.824</td>
<td>74.240</td>
<td>2227</td>
<td>2450</td>
<td>2524</td>
</tr>
<tr>
<td></td>
<td>0.456</td>
<td>74.715</td>
<td>--</td>
<td>2466</td>
<td>--</td>
<td></td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>0.019</td>
<td>74.735</td>
<td>--</td>
<td>2466</td>
<td>--</td>
</tr>
<tr>
<td>Weight of Mineral Admixture (Type IIP Cement)</td>
<td>75.000</td>
<td>75.000</td>
<td>2250</td>
<td>2475</td>
<td>2550</td>
<td></td>
</tr>
<tr>
<td>Total weight of Virgin Aggregate and Mineral Admixture</td>
<td>2280</td>
<td>2508</td>
<td>2584</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SIEVE</th>
<th>INDIVIDUAL PERCENT RETAINED</th>
<th>ACCUMULATIVE PERCENT RETAINED</th>
<th>RICE (1150 grams from each Dry Screened RAP stockpile)</th>
<th>Accumulative Percent Retained (Adjusted to 100%)</th>
<th>Accumulative Weight (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRY SCREENED RAP (26.282%)</td>
<td>1/4 Inch</td>
<td>5.081</td>
<td>5.081</td>
<td>152</td>
<td>168</td>
<td>173</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>2.106</td>
<td>7.197</td>
<td>217</td>
<td>239</td>
<td>246</td>
</tr>
<tr>
<td></td>
<td>#6</td>
<td>4.730</td>
<td>11.977</td>
<td>359</td>
<td>395</td>
<td>407</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>3.950</td>
<td>15.927</td>
<td>478</td>
<td>526</td>
<td>542</td>
</tr>
<tr>
<td></td>
<td>#4</td>
<td>3.852</td>
<td>20.369</td>
<td>611</td>
<td>672</td>
<td>693</td>
</tr>
<tr>
<td></td>
<td>#6</td>
<td>4.494</td>
<td>24.863</td>
<td>746</td>
<td>820</td>
<td>845</td>
</tr>
<tr>
<td></td>
<td>#8</td>
<td>2.820</td>
<td>26.054</td>
<td>782</td>
<td>860</td>
<td>886</td>
</tr>
<tr>
<td></td>
<td>1/2 Inch</td>
<td>1.2996</td>
<td>72.416</td>
<td>2172</td>
<td>2350</td>
<td>2462</td>
</tr>
<tr>
<td>Total weight of Virgin Aggregate and Dry Screened RAP</td>
<td>3038</td>
<td>3342</td>
<td>3444</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total weight of Virgin Aggregate, Dry Screened RAP, and Mineral Admixture</td>
<td>3068</td>
<td>3375</td>
<td>3478</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


**[2]** Accumulative Wt. for each sieve (Virgin Agg.) = Desired Weight of Virgin Agg. and RAP Agg. x Accumulative % Retained for each sieve (Virgin Agg.)

**[3]** Accumulative Wt. for each sieve (Dry Screened RAP) = Desired Weight of Virgin Agg. and RAP Agg. x Accumulative % Retained for each sieve (Dry Screened RAP)

**[4]** Accumulative Wt. for each sieve for Rice from each Dry Screened RAP Stockpile = Accumulative % Retained (Adjusted to 100%) x Accumulative Wt. from each sieve for Rice from each Dry Screened RAP Stockpile x 3000

---

**FIGURE 6**
Example Plots of Effective Voids, VMA, Dust/Bitumen Ratio, Bulk Specific Gravity, Marshall Stability, and Marshall Flow versus Percent Total Asphalt Binder Content

The above graphs are for illustrative purposes only. Straight-line interpolation, as specified in Subsections 13.3 and 13.3.1, is used for mix design calculations.

FIGURE 7
Asphaltic Concrete Marshall Mix Design with RAP
Example Mix Design Summary (Page 1 of 2)

CONTRACTOR: Best Paving Contractors, Inc.
ADOT Project Number: STP-999-Z(999A)
ADOT TRACS Number: H999901C
Project Name: Nowhere - Somewhere
Mix Designation: ADOT 416 (3/4 Inch Special Mix with RAP)

Date: February 22, 2013
Mix Design Laboratory: XYZ Testing Laboratories
Mix Design Engineer: Raul "RAP MAN" Raponi
Asphalt Binder Specified in Special Provisions: PG 76-16
Virgin Asphalt Binder Required by Mix Design: PG 76-22

COMPOSITE GRADATION:

Virgin Aggregate (75.0%) Percent Used
Washed Crusher Fines 35.0
Crusher Fines 13.0
3/8 Inch 8.0
3/4 Inch 19.0
RAP Aggregate (25.0 %)
RAP 1 (Fine) 15.0
RAP 2 (Coarse) 10.0
Mineral Admixture (1.0%)
Type IP Cement 1.0

IMMERSION COMPRESSION – ARIZ 802:
5.4% Total Binder, 1.0% Type IP Cement
Dry Strength (psi) 269
Wet Strength (psi) 197
Index of Retained Strength (%) 73

RAP 1 (FINE) RAP BINDER:
5.82%
RAP 2 (COARSE) RAP BINDER:
3.43%

RAP 1 (FINE) RAP BINDER GRADE (ACTUAL): 93-8
RAP 2 (COARSE) RAP BINDER GRADE (ACTUAL): 91-9

RAP 1 (FINE) RAP BINDER SPECIFIC GRAVITY: 1.049 @ 77°F
RAP 2 (COARSE) RAP BINDER SPECIFIC GRAVITY: 1.052 @ 77°F

AGGREGATE PROPERTIES: (See page 2.)

REMARKS:

FIGURE 8 (Continued on Next Page)
### FIGURE 8 (Continued from Previous Page)

**AGGREGATE PROPERTIES:**

<table>
<thead>
<tr>
<th>Test Characteristic</th>
<th>Virgin Aggregate</th>
<th>RAP Aggregate</th>
<th>Combined Virgin Aggregate and RAP Aggregate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Od Specific Gravity</td>
<td>2.797</td>
<td>2.789</td>
<td>2.794</td>
</tr>
<tr>
<td>SSD Specific Gravity</td>
<td>2.828</td>
<td>2.821</td>
<td>2.826</td>
</tr>
<tr>
<td>Apparent Specific Gravity</td>
<td>2.887</td>
<td>2.883</td>
<td>2.885</td>
</tr>
<tr>
<td>Absorption</td>
<td>1.11</td>
<td>1.17</td>
<td>1.13</td>
</tr>
<tr>
<td>Effective Specific Gravity</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Sand Equivalent</td>
<td>76</td>
<td>55 Min.</td>
<td>---</td>
</tr>
<tr>
<td>Uncompacted Void Content</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Carbonates</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Fractured Coarse Aggregate Particles</td>
<td>At least one fractured face</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>At least two fractured faces</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Abrasion</td>
<td>(Type B)</td>
<td>(Type B)</td>
<td>100 Rev. % loss</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>500 Rev. % loss</td>
</tr>
</tbody>
</table>
APPENDIX
ROUNDING PROCEDURE

The following describes the rounding procedure which is to be used for rounding numbers to the required degree of accuracy:

1. Except as specified in Section 2 below, the following procedure will apply. This procedure correlates with the "built-in" rounding method normally utilized by calculators and computers.

   1.1 When the figure next beyond the last figure or place to be retained is less than 5, the figure in the last place retained is left unchanged.

   Examples: Rounding 2.6324 to the nearest thousandth is 2.632
               Rounding 7843.343 to the nearest hundredth is 7843.34
               Rounding 4928.22 to the nearest tenth is 4928.2
               Rounding 7293.1 to the nearest whole number is 7293
               Rounding 4928.22 to the nearest whole number is 4928
               Rounding 2042 to the nearest units of 10 is 2040
               Rounding 3548 to the nearest units of 100 is 3500
               Rounding 8436 to the nearest units of 1000 is 8000

   1.2 When the figure next beyond the last figure or place to be retained is 5 or larger, the figure in the last place retained is increased by 1.

   Examples: Rounding 4839.4575 to the nearest thousandth is 4839.458
               Rounding 9347.215 to the nearest hundredth is 9347.22
               Rounding 8420.35 to the nearest tenth is 8420.4
               Rounding 1728.5 to the nearest whole number is 1729
               Rounding 3685 to the nearest units of 10 is 3690
               Rounding 6650 to the nearest units of 100 is 6700
               Rounding 2500 to the nearest units of 1000 is 3000

               Rounding 2.6326 to the nearest thousandth is 2.633
               Rounding 7843.347 to the nearest hundredth is 7843.35
               Rounding 4928.28 to the nearest tenth is 4928.3
               Rounding 7293.9 to the nearest whole number is 7294
               Rounding 2046 to the nearest units of 10 is 2050
               Rounding 3572 to the nearest units of 100 is 3600
               Rounding 8634 to the nearest units of 1000 is 9000
1.3 No result shall be rounded more than once.

Example: 3024.5 rounded to the nearest units of 10 will be 3020;

not

3024.5 rounded to 3025, and then rounded again to 3030.

2. The rounding procedure specified in Section 1 above shall be used for all calculations and recording of data in performing materials testing, except when a specific test method cites a method of rounding which differs from this procedure, for example, the sand equivalent test (AASHTO T 176 or Arizona Test Method 242).

3. Compliance will be based upon interpreting the reported results as though they are rounded to the terms (whole numbers, decimals, or fractions reduced to decimals) of the specifications. For example, a value reported as 8.4% shall be considered as having no deviation from specifications that require 4 – 8%. It would however be a deviation for specifications requiring 4.0 – 8.0%.

4. Computers and most electronic calculators automatically carry several decimal places beyond the point of desired accuracy. At times, results of calculations utilizing these values are different than that achieved when calculations are performed utilizing values that have been rounded to the desired degree of accuracy by the above rules. The user is cautioned that the use of a computer or electronic calculator without re-entry of values after rounding, and discarding any figures beyond those needed, may cause unacceptable variations in final results.
METRIC GUIDE

The following information is provided as a guide for utilizing the International System of Units (SI), generally referenced as "metric units". Related information can be found in AASHTO R1 "Metric Practice Guide", and ASTM E380, "Use of the International System of Units (SI) (The Modernized Metric System)"

Included herein are commonly used equivalents and conversions for U.S. Customary Units and Metric Units. It is not the intention of this guide to provide a detailed compilation of such equivalents and conversions. Such lists are available in many publications, including those referenced above.

One common conversion, which is found in many test procedures, is to determine an equivalent temperature in units of either degrees Celsius or degrees Fahrenheit.

Convert degrees Fahrenheit to degrees Celsius by:

\[ ^{\circ}C = \frac{5}{9} \times (^{\circ}F - 32) \]

Convert degrees Celsius to degrees Fahrenheit by:

\[ ^{\circ}F = \left[ \frac{9}{5} \times ^{\circ}C \right] + 32 \]

Under the SI (Metric) system, the base unit for mass is the "kilogram". (Although not technically correct, "weight" is often used in common practice to mean "mass".) The base unit for length is the "meter". The base unit for time is the "second". Primary metric units for area and volume are the "square meter" and the "cubic meter", respectively.

In addition to expressing values in the base or primary metric units, other associated metric units are identified and determined by varying the magnitude of the base metric unit by powers of 10. Metric values are commonly shown in scientific notation form, (for example, \(1 \times 10^4 = 10,000\); \(1 \times 10^{-4} = 0.0001\)).
Table 1 below gives a listing of prefixes used in the metric system, with their associated powers of ten, and their symbol.

<table>
<thead>
<tr>
<th>Prefix</th>
<th>Power of ten</th>
<th>Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>*deci</td>
<td>$10^{-1}$</td>
<td>d</td>
</tr>
<tr>
<td>*centi</td>
<td>$10^{-2}$</td>
<td>c</td>
</tr>
<tr>
<td>milli</td>
<td>$10^{-3}$</td>
<td>m</td>
</tr>
<tr>
<td>micro</td>
<td>$10^{-6}$</td>
<td>μ</td>
</tr>
<tr>
<td>nano</td>
<td>$10^{-9}$</td>
<td>n</td>
</tr>
<tr>
<td>pico</td>
<td>$10^{-12}$</td>
<td>p</td>
</tr>
<tr>
<td>femto</td>
<td>$10^{-15}$</td>
<td>f</td>
</tr>
<tr>
<td>atto</td>
<td>$10^{-18}$</td>
<td>a</td>
</tr>
<tr>
<td>*deka</td>
<td>$10^{1}$</td>
<td>da</td>
</tr>
<tr>
<td>*hecto</td>
<td>$10^{2}$</td>
<td>h</td>
</tr>
<tr>
<td>kilo</td>
<td>$10^{3}$</td>
<td>k</td>
</tr>
<tr>
<td>mega</td>
<td>$10^{6}$</td>
<td>M</td>
</tr>
<tr>
<td>giga</td>
<td>$10^{9}$</td>
<td>G</td>
</tr>
<tr>
<td>tera</td>
<td>$10^{12}$</td>
<td>T</td>
</tr>
<tr>
<td>peta</td>
<td>$10^{15}$</td>
<td>P</td>
</tr>
<tr>
<td>exa</td>
<td>$10^{18}$</td>
<td>E</td>
</tr>
</tbody>
</table>

* Use is to be avoided where practical. When expressing a quantity by a numerical value and a unit, a prefix should preferably be chosen so that the numerical value lies between 0.1 and 1000. In expressing area and volume, the prefixes hecto, deka, deci, and centi may be required, for example, square hectometer, cubic centimeter.
Table 2 below gives the symbols commonly used for various metric units.

<table>
<thead>
<tr>
<th>TABLE 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>kg = kilogram</td>
</tr>
<tr>
<td>g = gram</td>
</tr>
<tr>
<td>mg = milligram</td>
</tr>
<tr>
<td>m = meter</td>
</tr>
<tr>
<td>km = kilometer</td>
</tr>
<tr>
<td>cm = centimeter</td>
</tr>
<tr>
<td>mm = millimeter</td>
</tr>
<tr>
<td>( \mu m = ) micrometer (micron)</td>
</tr>
<tr>
<td>s = second</td>
</tr>
<tr>
<td>( m^2 = ) square meter</td>
</tr>
<tr>
<td>( cm^2 = ) square centimeter</td>
</tr>
<tr>
<td>( mm^2 = ) square millimeter</td>
</tr>
<tr>
<td>( m^3 = ) cubic meter</td>
</tr>
<tr>
<td>( cm^3 ) or ( cc = ) cubic centimeter</td>
</tr>
<tr>
<td>( mm^3 = ) cubic millimeter</td>
</tr>
<tr>
<td>L = liter</td>
</tr>
<tr>
<td>mL = milliliter</td>
</tr>
<tr>
<td>Pa = pascal</td>
</tr>
<tr>
<td>N = newton</td>
</tr>
<tr>
<td>kPa = kilopascal</td>
</tr>
<tr>
<td>MPa = megapascal</td>
</tr>
</tbody>
</table>
Table 3 below includes common conversions from the base and primary metric units (kilogram, meter, square meter, and cubic meter) to other associated metric units. Also listed are some common derived metric units.

**TABLE 3**

<table>
<thead>
<tr>
<th>Metric Unit Conversion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 gram = 0.001 kilogram</td>
</tr>
<tr>
<td>1 milligram = 1 x 10^-6 kilogram</td>
</tr>
<tr>
<td>1 milligram = 0.001 gram</td>
</tr>
<tr>
<td>1 kilogram = 1000 grams</td>
</tr>
<tr>
<td>1 metric ton = 1000 kilograms</td>
</tr>
<tr>
<td>1 kilometer = 1000 meters</td>
</tr>
<tr>
<td>1 centimeter = 0.01 meter</td>
</tr>
<tr>
<td>1 millimeter = 0.001 meter</td>
</tr>
<tr>
<td>1 micron (micrometer) = 1 x 10^-6 meter</td>
</tr>
<tr>
<td>1 square kilometer = 1 x 10^6 square meters</td>
</tr>
<tr>
<td>1 square centimeter = 1 x 10^-4 square meter</td>
</tr>
<tr>
<td>1 square millimeter = 1 x 10^-6 square meter</td>
</tr>
<tr>
<td>1 cubic centimeter = 1 x 10^-6 cubic meter</td>
</tr>
<tr>
<td>1 cubic millimeter = 1 x 10^-9 cubic meter</td>
</tr>
<tr>
<td>1 liter = 0.001 cubic meter</td>
</tr>
<tr>
<td>1 milliliter = 1 x 10^-6 cubic meter</td>
</tr>
<tr>
<td>1 milliliter = 1 cubic centimeter</td>
</tr>
<tr>
<td>1 newton = 1 kg·m/s^2</td>
</tr>
<tr>
<td>1 pascal = 1 N/m^2</td>
</tr>
<tr>
<td>1 kilopascal = 1000 pascals</td>
</tr>
<tr>
<td>1 megapascal = 1 x 10^6 pascals</td>
</tr>
<tr>
<td>1 poise (absolute viscosity) = 0.10 Pa·s</td>
</tr>
<tr>
<td>1 centistoke (kinematic viscosity) = 1 mm^2/s or 1 x 10^-6 m^2/s</td>
</tr>
</tbody>
</table>
Some common U.S Customary units, with their corresponding base and primary metric unit equivalents, are given below in Table 4.

| TABLE 4 |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| 1 pound (avoirdupois) = 0.453 5924 kilogram |
| 1 ton (2000 lbs.) = 907.1847 kilograms |
| * 1 inch = 0.0254 meter |
| * 1 foot = 0.3048 meter |
| * 1 yard = 0.9144 meter |
| * 1 mile (5280 feet) = 1609.344 meters |
| * 1 square inch = 6.4516 x 10^{-4} square meter |
| * 1 square foot = 0.092 903 04 square meter |
| 1 square yard = 0.836 1274 square meter |
| 1 cubic inch = 1.638 706 x 10^{-5} cubic meter |
| 1 cubic foot = 0.028 316 85 cubic meter |
| 1 cubic yard = 0.764 5549 cubic meter |
| 1 pint (U.S. liquid) = 4.731 765 x 10^{-4} cubic meter |
| 1 quart (U.S. liquid) = 9.463 529 x 10^{-4} cubic meter |
| 1 gallon (U.S. liquid) = 0.003 785 412 cubic meter |

(Exact equivalents are noted with an asterisk.)
Table 5 below lists commonly used conversions for U.S. Customary Units and metric units. Values are shown to the degree of accuracy which generally may be used to achieve satisfactory results. If more accuracy is desired, the values may be derived by using Tables 3 and 4.

**TABLE 5**

<table>
<thead>
<tr>
<th>Conversion</th>
<th>Equivalence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 kilogram = 2.205 pounds</td>
<td></td>
</tr>
<tr>
<td>1 pound = 453.6 grams</td>
<td></td>
</tr>
<tr>
<td>1 ounce (avoirdupois) = 28.35 grams</td>
<td></td>
</tr>
<tr>
<td>1 ton (2000 lbs) = 0.9072 metric ton</td>
<td></td>
</tr>
<tr>
<td>1 meter = 39.37 inches or 3.281 feet</td>
<td></td>
</tr>
<tr>
<td>1 kilometer = 0.62 miles</td>
<td></td>
</tr>
<tr>
<td>* 1 mil = 0.0254 millimeters or 25.4 micrometers</td>
<td></td>
</tr>
<tr>
<td>* 1 inch = 2.54 centimeters or 25.4 millimeters</td>
<td></td>
</tr>
<tr>
<td>* 1 foot = 0.3048 meters</td>
<td></td>
</tr>
<tr>
<td>* 1 yard = 0.9144 meters</td>
<td></td>
</tr>
<tr>
<td>1 mile = 1.61 kilometers</td>
<td></td>
</tr>
<tr>
<td>1 square inch = 6.452 cm$^2$ or 645.16 mm$^2$</td>
<td></td>
</tr>
<tr>
<td>1 square foot = 0.0929 square meters</td>
<td></td>
</tr>
<tr>
<td>1 square yard = 0.836 square meters</td>
<td></td>
</tr>
<tr>
<td>1 cubic inch = 16.39 cm$^3$ or 16386 mm$^3$</td>
<td></td>
</tr>
<tr>
<td>1 cubic foot = 0.028 m$^3$ or 28317 cm$^3$</td>
<td></td>
</tr>
<tr>
<td>1 cubic yard = 0.765 cubic meters</td>
<td></td>
</tr>
<tr>
<td>1 liter = 1.06 quarts (U.S. liquid)</td>
<td></td>
</tr>
<tr>
<td>1 ounce (U.S. fluid) = 29.574 milliliter</td>
<td></td>
</tr>
<tr>
<td>1 pint (U.S. liquid) = 0.47 liter</td>
<td></td>
</tr>
<tr>
<td>1 quart (U.S. liquid) = 0.95 liter</td>
<td></td>
</tr>
<tr>
<td>1 gallon (U.S. liquid) = 3.79 liters</td>
<td></td>
</tr>
<tr>
<td>1 lb/ft$^3$ = 16.02 kg/m$^3$</td>
<td></td>
</tr>
<tr>
<td>1 kilometer/hour = 0.62 mile/hour</td>
<td></td>
</tr>
<tr>
<td>1 mile/hour = 1.61 km/hour</td>
<td></td>
</tr>
<tr>
<td>1 pound/square inch = 6.895 kPa</td>
<td></td>
</tr>
<tr>
<td>1 pound force = 4.448 newton</td>
<td></td>
</tr>
<tr>
<td>1 gallon/square yard = 4.527 liters/m$^2$</td>
<td></td>
</tr>
<tr>
<td>1 gallon/ton (2000 lbs.) = 4.173 liters/metric ton</td>
<td></td>
</tr>
<tr>
<td>1 gallon/cubic yard = 4.951 liters/m$^3$</td>
<td></td>
</tr>
<tr>
<td>1 pound/square yard = 0.542 kg/m$^2$</td>
<td></td>
</tr>
<tr>
<td>1 pound/cubic yard = 0.593 kg/m$^3$</td>
<td></td>
</tr>
<tr>
<td>1 pound/gallon = 0.120 kg/liter</td>
<td></td>
</tr>
<tr>
<td>1 cubic yard/square yard = 0.914 m$^3$/m$^2$</td>
<td></td>
</tr>
<tr>
<td>1 inch/mile = 0.0158 meter/kilometer</td>
<td></td>
</tr>
</tbody>
</table>

(Exact equivalents are noted with an asterisk.)
Table 6 below is from information contained in AASHTO M92 and ASTM E11 "Wire-Cloth Sieves for Testing Purposes", and shows Standard (Metric) and Alternative (U.S. Customary) sieve size designations. As shown, metric size designations are given in mm or μm. (1,000 μm = 1 millimeter)

8 inch diameter sieve = 203.2 mm diameter sieve
12 inch diameter sieve = 304.8 mm diameter sieve

<table>
<thead>
<tr>
<th>Sieve Designation</th>
<th>Standard (mm)</th>
<th>Alternate</th>
<th>Sieve Designation</th>
<th>Standard (μm)</th>
<th>Alternate</th>
</tr>
</thead>
<tbody>
<tr>
<td>125 mm</td>
<td>5 in.</td>
<td>2.36 mm</td>
<td>No. 8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>106 mm</td>
<td>4.24 in.</td>
<td>2.00 mm</td>
<td>No. 10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 mm</td>
<td>4 in.</td>
<td>1.70 mm</td>
<td>No. 12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>90 mm</td>
<td>3-1/2 in.</td>
<td>1.40 mm</td>
<td>No. 14</td>
<td></td>
<td></td>
</tr>
<tr>
<td>75 mm</td>
<td>3 in.</td>
<td>1.18 mm</td>
<td>No. 16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>63 mm</td>
<td>2-1/2 in.</td>
<td>1.00 mm</td>
<td>No. 18</td>
<td></td>
<td></td>
</tr>
<tr>
<td>53 mm</td>
<td>2.12 in.</td>
<td>850 μm</td>
<td>No. 20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>50 mm</td>
<td>2 in.</td>
<td>710 μm</td>
<td>No. 25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>45 mm</td>
<td>1-3/4 in.</td>
<td>600 μm</td>
<td>No. 30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>37.5 mm</td>
<td>1-1/2 in.</td>
<td>500 μm</td>
<td>No. 35</td>
<td></td>
<td></td>
</tr>
<tr>
<td>31.5 mm</td>
<td>1-1/4 in.</td>
<td>425 μm</td>
<td>No. 40</td>
<td></td>
<td></td>
</tr>
<tr>
<td>26.5 mm</td>
<td>1.06 in.</td>
<td>355 μm</td>
<td>No. 45</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25.0 mm</td>
<td>1 in.</td>
<td>300 μm</td>
<td>No. 50</td>
<td></td>
<td></td>
</tr>
<tr>
<td>22.4 mm</td>
<td>7/8 in.</td>
<td>250 μm</td>
<td>No. 60</td>
<td></td>
<td></td>
</tr>
<tr>
<td>19.0 mm</td>
<td>3/4 in.</td>
<td>212 μm</td>
<td>No. 70</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.0 mm</td>
<td>5/8 in.</td>
<td>180 μm</td>
<td>No. 80</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13.2 mm</td>
<td>0.530 in.</td>
<td>150 μm</td>
<td>No. 100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12.5 mm</td>
<td>1/2 in.</td>
<td>125 μm</td>
<td>No. 120</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11.2 mm</td>
<td>7/16 in.</td>
<td>106 μm</td>
<td>No. 140</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9.5 mm</td>
<td>3/8 in.</td>
<td>90 μm</td>
<td>No. 170</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8.0 mm</td>
<td>5/16 in.</td>
<td>75 μm</td>
<td>No. 200</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.7 mm</td>
<td>0.265 in.</td>
<td>63 μm</td>
<td>No. 230</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.3 mm</td>
<td>1/4 in.</td>
<td>53 μm</td>
<td>No. 270</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6 mm</td>
<td>No. 3-1/2</td>
<td>45 μm</td>
<td>No. 325</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.75 mm</td>
<td>No. 4</td>
<td>38 μm</td>
<td>No. 400</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.00 mm</td>
<td>No. 5</td>
<td>32 μm</td>
<td>No. 450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.35 mm</td>
<td>No. 6</td>
<td>25 μm</td>
<td>No. 500</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.80 mm</td>
<td>No. 7</td>
<td>20 μm</td>
<td>No. 635</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### TABLE 7

**SI* (METRIC) CONVERSION FACTORS**

(Approximate equivalents, except as noted**)

<table>
<thead>
<tr>
<th>CONVERSIONS TO SI UNITS</th>
<th>CONVERSIONS FROM SI UNITS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Abbrev./Symbol</strong></td>
<td><strong>When you know</strong></td>
</tr>
<tr>
<td>LENGTH</td>
<td>LENGTH</td>
</tr>
<tr>
<td>in.</td>
<td>inches</td>
</tr>
<tr>
<td>ft.</td>
<td>feet</td>
</tr>
<tr>
<td>yd.</td>
<td>yards</td>
</tr>
<tr>
<td>mi.</td>
<td>miles</td>
</tr>
<tr>
<td>AREA</td>
<td>AREA</td>
</tr>
<tr>
<td>sq. in. or in²</td>
<td>square inches</td>
</tr>
<tr>
<td>sq. ft. or ft²</td>
<td>square feet</td>
</tr>
<tr>
<td>sq. yd. or yd²</td>
<td>square yards</td>
</tr>
<tr>
<td>acres</td>
<td>0.405</td>
</tr>
<tr>
<td>sq. mi. or mi²</td>
<td>square miles</td>
</tr>
<tr>
<td>VOLUME</td>
<td>VOLUME</td>
</tr>
<tr>
<td>fl. oz.</td>
<td>fluid ounces</td>
</tr>
<tr>
<td>gal.</td>
<td>gallons (liquid)</td>
</tr>
<tr>
<td>cu. ft. or ft³</td>
<td>cubic feet</td>
</tr>
<tr>
<td>cu. yd. or yd³</td>
<td>cubic yards</td>
</tr>
<tr>
<td>MASS</td>
<td>MASS</td>
</tr>
<tr>
<td>oz.</td>
<td>ounces</td>
</tr>
<tr>
<td>lb.</td>
<td>pounds</td>
</tr>
<tr>
<td>T</td>
<td>short tons (2000 lb)</td>
</tr>
<tr>
<td>T</td>
<td>short tons (2000 lb)</td>
</tr>
</tbody>
</table>

* SI is the symbol for the International System of Units.
** Exact equivalent.
*** Metric volumes greater than 1000 liters should be shown in m³. The following conversion factors are helpful in making necessary conversions:
   1 liter = 0.001 cubic meter; 1 cubic meter = 264.17 gallons (liquid).
**** 1 metric ton = 1000 kilograms = 1,000,000 grams = 1 Mg.
EQUIPMENT CALIBRATION
AND VERIFICATION

Equipment and apparatus used in testing materials must be calibrated or verified for accuracy on a regular basis to assure the equipment is giving reliable results.

Materials Group, Quality Assurance Section, has developed a listing of requirements, frequency, and the applicable test methods for the calibration and verification of various types of equipment. The listing provided is not intended to address the calibration or verification of all equipment. Appendix A3 contains the following figures:

<table>
<thead>
<tr>
<th>Figure Number</th>
<th>Figure Title</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>General Laboratory Testing Equipment</td>
</tr>
<tr>
<td>2</td>
<td>Aggregates Testing Equipment</td>
</tr>
<tr>
<td>3</td>
<td>Asphalt Binder/Cutback Asphalt/Emulsified Asphalt Testing Equipment</td>
</tr>
<tr>
<td>4</td>
<td>Asphalt Mixtures Testing Equipment</td>
</tr>
<tr>
<td>5</td>
<td>Soils Testing Equipment</td>
</tr>
<tr>
<td>6</td>
<td>Soils Testing Equipment (Continued)</td>
</tr>
<tr>
<td>7</td>
<td>Portland Cement Concrete Testing Equipment</td>
</tr>
<tr>
<td>8</td>
<td>Metal Testing Equipment</td>
</tr>
<tr>
<td>9</td>
<td>Equipment Maintenance</td>
</tr>
</tbody>
</table>

Test methods and standards referenced in the figures contained in Appendix A3 are identified as follows:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AZ</td>
<td>Arizona Test Methods</td>
</tr>
<tr>
<td>T, M, and R</td>
<td>AASHTO Test Methods and Standards</td>
</tr>
<tr>
<td>C, D, and E</td>
<td>ASTM Test Methods and Standards</td>
</tr>
</tbody>
</table>

Related information on the calibration and verification of apparatus can be found in AASHTO R 18, "Establishing and Implementing a Quality System for Construction Materials Testing Laboratories".

Maximum frequency intervals shown in the following figures may be reduced based on specific laboratory conditions, deterioration of equipment, frequency of use, and other contributing factors.

Equipment and apparatus that may be affected by movement must be recalibrated after relocation.
## GENERAL LABORATORY TESTING EQUIPMENT

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical Shakers</td>
<td>Check Sieving Thoroughness.</td>
<td>12</td>
</tr>
<tr>
<td>Ovens</td>
<td>Standardize Thermometric Device.</td>
<td>12</td>
</tr>
<tr>
<td>Sieves</td>
<td>Fine Sieves (openings &lt; No.4) Check Physical Condition</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>Coarse Sieves (openings ≥ No.4) Check Physical Condition and Dimensions of Openings.</td>
<td>12</td>
</tr>
<tr>
<td>Specimen Molds</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
</tr>
<tr>
<td>General Purpose Balances and Masses</td>
<td>Standardize</td>
<td>12</td>
</tr>
<tr>
<td>Thermometers, including Digital Thermistors</td>
<td>Standardize</td>
<td>12</td>
</tr>
<tr>
<td>Analytical Balances and Masses</td>
<td>Calibrate</td>
<td>12</td>
</tr>
<tr>
<td>Calipers</td>
<td>Standardize</td>
<td>12</td>
</tr>
<tr>
<td>Vacuum/Pressure Measurement Devices (Bourdon Gauges, Pressure Manometer, and Electronic Pressure Transducers)</td>
<td>Standardize</td>
<td>12</td>
</tr>
<tr>
<td>Length Measurement Devices (Dial Indicators, LDT's, LVDT's, and Extensometers)</td>
<td>Standardize</td>
<td>12</td>
</tr>
<tr>
<td>Compression, Loading, or Tensile Testing Devices</td>
<td>Standardize</td>
<td>12</td>
</tr>
<tr>
<td>Constant Temperature Baths</td>
<td>Check Temperature Settings.</td>
<td>12</td>
</tr>
</tbody>
</table>

**FIGURE 1**
<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit Weight Apparatus</td>
<td>Standardize</td>
<td>12</td>
<td>T 19, C 29</td>
</tr>
<tr>
<td>Sulfate Oven</td>
<td>Check Rate of Evaporation.</td>
<td>12</td>
<td>T 104, C 88</td>
</tr>
<tr>
<td>Sulfate Soundness Sample Containers</td>
<td>Check Physical Condition.</td>
<td>12</td>
<td>T 104, C 88</td>
</tr>
<tr>
<td>L.A. Abrasion Machine</td>
<td>Check RPM and Critical Dimensions.</td>
<td>24</td>
<td>T 96, C 131</td>
</tr>
<tr>
<td>Steel Balls for Abrasion Machine</td>
<td>Check Individual Weight and Charge Weight.</td>
<td>24</td>
<td>T 96, C 731</td>
</tr>
<tr>
<td>Conical Mold and Tamper</td>
<td>Check Critical Dimensions.</td>
<td>24</td>
<td>AZ 211, T 84, C 128</td>
</tr>
</tbody>
</table>

**FIGURE 2**
<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saybolt Viscometer</td>
<td>Standardize</td>
<td>36</td>
<td>T 59, D 244, T 72</td>
</tr>
<tr>
<td>Timing Devices</td>
<td>Standardize</td>
<td>12</td>
<td>T 49, D 5</td>
</tr>
<tr>
<td>Penetrometer Needle</td>
<td>Check Condition and Critical Dimensions.</td>
<td>12</td>
<td>T 49, D 5</td>
</tr>
<tr>
<td>Penetrometer</td>
<td>Standardize for Penetration Depth.</td>
<td>12</td>
<td>T 49, D 5</td>
</tr>
<tr>
<td>Ductility Apparatus</td>
<td>Check Speed of Travel.</td>
<td>12</td>
<td>T 51, D 113</td>
</tr>
<tr>
<td>Elastic Recovery Apparatus</td>
<td>Check Speed of Travel.</td>
<td>12</td>
<td>T 391, D 6084</td>
</tr>
<tr>
<td>RTFO Carriage</td>
<td>Check Rotation Speed.</td>
<td>12</td>
<td>T 240, D 2872</td>
</tr>
<tr>
<td>Brass Rings and Assembly</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>T 53, D 36</td>
</tr>
<tr>
<td>Pycnometers</td>
<td>Check Physical Condition and Standardize Volume.</td>
<td>12</td>
<td>T 228, D 70</td>
</tr>
<tr>
<td>Collars and Floats</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>T 59, D 139</td>
</tr>
<tr>
<td>Pressurized Aging Vessel</td>
<td>Standardize Temperature and Pressure.</td>
<td>6</td>
<td>R 28, D 6521</td>
</tr>
<tr>
<td>Rotational Viscometer</td>
<td>Standardize with Reference Fluid.</td>
<td>6</td>
<td>T 316, D 4402</td>
</tr>
<tr>
<td>Dynamic Shear Rheometer (DSR)</td>
<td>Standardize with Reference Fluid.</td>
<td>6</td>
<td>T 315, D 7175</td>
</tr>
<tr>
<td>Bending Beam Rheometer (BBR)</td>
<td>Calibrate Masses.</td>
<td>12</td>
<td>T 313, D 6648</td>
</tr>
<tr>
<td>Kinematic Viscometer Tubes</td>
<td>Calibrate with Reference Fluid.</td>
<td>36</td>
<td>T 201, D 2170</td>
</tr>
</tbody>
</table>

**FIGURE 3**
## APPENDIX A3

### September 28, 2012

**ARIZONA DEPARTMENT OF TRANSPORTATION**  
**MATERIALS GROUP**  
**EQUIPMENT CALIBRATION AND VERIFICATION**

**ASPHALT MIXTURES TESTING EQUIPMENT**

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical Compaction Hammers, Breaking Heads</td>
<td>Check Critical Dimensions, Check Mass of Hammer.</td>
<td>12</td>
<td>AZ 410, AZ 815, T 245, T 283, D 4867, D 6926, D 6927</td>
</tr>
<tr>
<td></td>
<td>Monitor Through Proficiency Sample Program.</td>
<td>Per proficiency sample frequency.</td>
<td></td>
</tr>
<tr>
<td>Marshall Stability/Flow Testing Machine</td>
<td>Check Speed of Travel, Verify Load Cell.</td>
<td>12</td>
<td>AZ 410, AZ 422, AZ 815, T 245, T 283, D 4867, D 5581, D 6926, D 6927</td>
</tr>
<tr>
<td>Plungers</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>AZ 802, T 165, T 176, D 1074</td>
</tr>
<tr>
<td>Gyratory Compactor</td>
<td>Standardize Ram Pressure, Frequency of Gyration, LVDT.</td>
<td>12</td>
<td>T 312, D 6925</td>
</tr>
<tr>
<td>Gyratory Compactor</td>
<td>Standardize External or Internal Angle of Gyration.</td>
<td>12</td>
<td>T 312, D 6925</td>
</tr>
<tr>
<td>Ram Face, Base Plate Face</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>T 312, D 7115</td>
</tr>
<tr>
<td>Ignition Oven Internal Balance</td>
<td>Standardize</td>
<td>12</td>
<td>AZ 427, T 308, D 6307</td>
</tr>
<tr>
<td>Rice Flasks</td>
<td>Calibrate Volume.</td>
<td>12</td>
<td>AZ 417, AZ 806, T 209</td>
</tr>
<tr>
<td>Nuclear Asphalt Content Gauge</td>
<td>Check Variability using 3 - Point Calibration.</td>
<td>12</td>
<td>AZ 421</td>
</tr>
</tbody>
</table>

**FIGURE 4**
## SOILS TESTING EQUIPMENT

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical Compactor</td>
<td>Check Rammer Face Diameter and Height of Drop.</td>
<td>12</td>
<td>AZ 225, AZ 226, AZ 232, AZ 245, AZ 246, T 99, T 134, T 135, T 180, D 558, D 559, D 698, D 1557</td>
</tr>
<tr>
<td></td>
<td>Monitor Through Proficiency Sample Program.</td>
<td>Per proficiency sample frequency.</td>
<td></td>
</tr>
<tr>
<td>Liquid Limit Device</td>
<td>Check Wear and Critical Dimensions.</td>
<td>12</td>
<td>T 89, D 4318</td>
</tr>
<tr>
<td>Grooving Tool</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>T 89, D 4318</td>
</tr>
<tr>
<td>Hydrometers</td>
<td>Check Critical Dimensions.</td>
<td>24</td>
<td>T 88, D 422</td>
</tr>
<tr>
<td>Straightedge</td>
<td>Check Dimensions and Planeness of Edge.</td>
<td>12</td>
<td>AZ 225, AZ 245, T 99, T 134, T 135, T 136, T 180, D 558, D 559, D 560, D 698, D 1557</td>
</tr>
<tr>
<td>Weighted Foot Assembly</td>
<td>Check Mass.</td>
<td>12</td>
<td>AZ 242, T 176, D 2419</td>
</tr>
<tr>
<td>CA Kneading Compactor</td>
<td>Standardize</td>
<td>24</td>
<td>T 190, D 2844</td>
</tr>
<tr>
<td>Standard Metal Specimen</td>
<td>Check Outside Diameter.</td>
<td>12</td>
<td>T 190, D 2844</td>
</tr>
<tr>
<td>Metal Follower</td>
<td>Check Diameter.</td>
<td>12</td>
<td>T 190, D 2844</td>
</tr>
</tbody>
</table>

**FIGURE 5**
## Soils Testing Equipment (Continued)

<table>
<thead>
<tr>
<th>Equipment</th>
<th>Requirement</th>
<th>Max. Interval (Months)</th>
<th>Test Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vertical Load</td>
<td>Standardize</td>
<td>12</td>
<td>AZ 249, T 216, T 236, D 2435, D 3080, D 4829</td>
</tr>
<tr>
<td>Uncompacted Void Content Apparatus</td>
<td>Check Critical Dimensions and Calibrate Volume.</td>
<td>12</td>
<td>AZ 247, C 1252</td>
</tr>
<tr>
<td>Resistivity Apparatus</td>
<td>Check Resistivity.</td>
<td>12</td>
<td>AZ 236</td>
</tr>
<tr>
<td>pH Meter</td>
<td>Standardize</td>
<td>12</td>
<td>AZ 236, AZ 237</td>
</tr>
<tr>
<td>Flat and Elongated Proportional Caliper Device</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>D 4791</td>
</tr>
<tr>
<td>Flakiness Index Slotted Sieves</td>
<td>Check Critical Dimensions.</td>
<td>12</td>
<td>AZ 233</td>
</tr>
<tr>
<td>Fine Aggregate Specific Gravity Flask</td>
<td>Calibrate Volume.</td>
<td>12</td>
<td>AZ 211, T 84, C 128</td>
</tr>
<tr>
<td>Coarse Aggregate Specific Gravity Apparatus</td>
<td>Check Wire Diameter, Basket Type, and Water Tank.</td>
<td>12</td>
<td>AZ 210, T 85, C 127</td>
</tr>
<tr>
<td>Speedie Moisture Tester</td>
<td>Verify Scale and Gauge.</td>
<td>6</td>
<td>T 217</td>
</tr>
<tr>
<td>Standard Sand</td>
<td>Check Each New Shipment for Conformance to C 778.</td>
<td>- - -</td>
<td>C 778</td>
</tr>
<tr>
<td>Sand Cone Density Apparatus</td>
<td>Standardize</td>
<td>12</td>
<td>AZ 229, AZ 230</td>
</tr>
</tbody>
</table>

**Figure 6**
# ARIZONA DEPARTMENT OF TRANSPORTATION
MATERIALS GROUP
EQUIPMENT CALIBRATION AND VERIFICATION

September 28, 2012

## PORTLAND CEMENT CONCRETE TESTING EQUIPMENT

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit Weight Measures</td>
<td>Standardize</td>
<td>12</td>
<td>T 121, C 138</td>
</tr>
<tr>
<td>Air Meters (Pressure Type)</td>
<td>Standardize</td>
<td>3 (See Note)</td>
<td>T 152, C 231</td>
</tr>
<tr>
<td>Air Meters (Volumetric Type)</td>
<td>Standardize</td>
<td>12 (See Note)</td>
<td>C 173</td>
</tr>
</tbody>
</table>

Note: The individual test methods specify conditions that require restandardization, such as changes in elevation and rough handling.

Capping Material
- Check Strength
  - 3
  - T 231, C 617

Slump Cones and Rods
- Check Critical Dimensions
  - 12
  - T 119, C 143

Single-Use Concrete Cylinder Molds
- Check Critical Dimensions of each Shipment
  - - -
  - T 22, T 23, C 31, C 39

Cylinder Capping Plates
- Check Critical Dimensions
  - 12
  - T 231, C 617

Recording Thermometer
- Standardize
  - 6
  - C 31, C 39

Autoclave Apparatus
- Check Critical Dimensions, Calibrate Valve and Gauge
  - 12
  - T 107, M 210, C 151

Rebound Hammer (Swiss Hammer)
- Standardize
  - 6
  - C 805

Moist Room (Fog Room)
- Verify Temperature and Humidity with Recording Thermometer
  - 6
  - M 201, C 511

Cube Molds and Tampers
- Check Critical Dimensions and Physical Condition
  - 30
  - T 106, C 109

**FIGURE 7**
### METAL TESTING EQUIPMENT

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>REQUIREMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rockwell Hardness Tester</td>
<td>Standardize</td>
<td>12</td>
<td>E 18</td>
</tr>
</tbody>
</table>

**FIGURE 8**
## EQUIPMENT MAINTENANCE

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>MAX. INTERVAL (Months)</th>
<th>TEST METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Performance-Graded Binder Equipment</td>
<td>12</td>
<td>R 58, T 240, T 313, T 314, T 315, D 2872, D 6521, D 6648, D 6722, D 7175</td>
</tr>
<tr>
<td>Ductimeters</td>
<td>12</td>
<td>T 51, T 306, T 301, D 113, D 6084</td>
</tr>
<tr>
<td>Mechanical Marshall Compactors</td>
<td>12</td>
<td>T 245</td>
</tr>
<tr>
<td>California Kneading Compactors</td>
<td>12</td>
<td>T 247, T 190, D 1561, D 2844</td>
</tr>
<tr>
<td>Gyratory Compactors</td>
<td>12</td>
<td>T 312, D 6925</td>
</tr>
<tr>
<td>Mechanical Compactors</td>
<td>12</td>
<td>T 99, T 180, D 698, D 1557</td>
</tr>
<tr>
<td>Mechanical Shakers</td>
<td>12</td>
<td>Where Applicable.</td>
</tr>
</tbody>
</table>

**NOTE:** There may be more items added to the laboratory’s list of equipment that require maintenance. Maintenance activities will typically involve lubricating, tightening fittings, cleaning, replacing fluids, checking and replacing damaged or worn parts, etc. These activities will vary based on the type of equipment, how often the equipment is used, the manufacturer’s recommendations, etc.

**FIGURE 9**