## Meinzer II<sup>TM</sup>

## **Testing Sieve Shaker**

**Operation & Set-up Manual** 



Model: MEINZ110 MEINZ220

## Advantech Manufacturing, Inc.

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A Product of the United State of America



## Introduction

Thank you for selecting this high-quality piece of testing equipment. We appreciate your support and pledge to assist you in the service of your Advantech testing apparatus.

The Meinzer II is a maintenance free, lightweight and portable vibrating shaker that will provide precise, repeatable results time after time.

By utilizing an electromagnetic drive, fixed amplitude and rubber spring mounts, this unit produces the simultaneous vertical and horizontal movement needed for basic dry particle sizing analysis. This unit is ideal for use with aggregates, cements, chemicals, powdered metals, cosmetics, grains, seeds, coal, soils, pharmaceuticals, tobacco, coffee and many other dry components in ground, granular or powder form. Each unit is "burned in" and run continuously for over half a day, guaranteeing performance right out of the box. *This unit is not recommended for wet sieving operation.* 

Besides the physical nuts and bolts, this device is backed by a company with decades of experience in the dedicated service of users in the powder and particulate industries. We look forward to servicing you as well.

"The Leader in Sieving Technology""

## **Specifications**

### Model Designations and Power Requirements

- Model MEINZ110 . . . . . . 110VAC/60Hz operation. Accepts 8"(203.2mm), 200mm or 3" (76.2mm) diameter sieves
- Model MEINZ220 . . . . . 220VAC/50/60Hz operation. Accepts 8" (203.2mm), 200mm or 3" (76.2mm) diameter sieves

#### Timer

• 60 minute variable, automatic shut-off, or continuous run setting

### **Dimensional Specifications**

## Weight

• 37 lbs (17Kg)

## **General Specifications**

- Durable, powder coated finish
- Unit Capacity . . . . . . . 8" (203.2mm) or 200mm 8 full height or 15 half height, plus pan and cover

### Unit ships with

- Operation and Set-up Manual
- 6' (2 meter) Power Cable
- Top Plate Assembly
- Meinzer II Shaker fitted with Clamping Straps and Buckles
- Sieves, pans and covers must be ordered separately.

## Installation & Set-up Instructions

The Meinzer II Testing Sieve Shaker is designed to provide years of trouble-free service. To ensure this device delivers optimum performance, several points must be observed prior to operation.

#### 1) Installation

Position the machine on a level, sturdy surface to ensure the even distribution of the sample over the sieves during operation.

#### 2) Electrical Connections

Verify that the voltage and frequency on the Rating Label at the back of the shaker corresponds with the electrical power supply being used. If any discrepancy occurs, please consult your supplier or contact Advantech Tech Support for assistance. Do not connect any power supply other than that stated on the Rating Label. <a href="Important!">Important!</a> This equipment must be grounded.

When the power is properly connected, the red rocker switch will illuminate when switched to the "on" position. If the switch does not light, the connector may not be fully inserted or the fuse may be blown. Replacement fuses must be of identical rating.

#### 3) Process Timer

The Meinzer II features a mechanical, 60 minute variable timer with automatic shut-off. There is also a continuous running time feature. Set the timer by rotating the dial past the desired interval, then return it to the desired time. Operating periods are increased by rotating clockwise and decreased by rotating counter-clockwise. The timer will commence timing down as soon as the knob is released, regardless of an electrical connection. When the knob is turned counter-clockwise from the "OFF" position to the continuous running mark, "I", the shaker will continue running until the knob is manually returned to the "OFF" position.

#### 4) Maintenance

The Meinzer II Sieve Shaker is maintenance free aside from keeping surfaces clean. Simply wipe with a soft damp cloth using a solution of warm water and a mild liquid detergent. **Do not clean your sieve shaker or sieves with solvents.** 

### 6) General Advice

The Meinzer II Sieve Shaker is constructed and factory tested to ensure correct operation when connected to the specific electrical power supply indicated on the Rating Label of the machine. Under typical usage, no lubrication or resetting is required. Use of unapproved parts or any alteration to the machine voids the warranty.

For replacement parts, please contact Advantech Mfg.'s Tech Support Team at 262.786.1600 or <a href="mailto:sales@advantechmfg.com">sales@advantechmfg.com</a>. Please have your serial number ready for prompt service.

Advantech Mfg. does not accept any responsibility if the operating instructions contained in this manual are not strictly followed.

# Performing a Sieve Analysis using the Meinzer II Testing Sieve Shaker

- 1) Complete set up of the Meinzer II Testing Sieve Shaker per instructions under "Installation and Set-Up Instructions".
- 2) Plug the device into the proper power source (be sure that voltage and cycle requirements are observed).
- 3) Prepare the material sample to be tested using industry-specified sampling and preparation procedures.
- 4) Select the sieves for the analysis.
- 5) Assemble the sieve stack, (coarsest sieve at the top, finest at the bottom) with bottom pan. An extended rim pan may be inserted within the stack to run multiple samples. See **Figure 7 A** in the FAQ's for an example of the extended rim pan. Bear in mind the overall height of the sieve stack may not exceed the capacities shown in **Chart 4A** of the FAQ's.
- 6) Pour the sample to be tested onto the top sieve. Install a standard sieve cover to prevent sample loss.
- 7) Place the sieve stack on the Bottom Plate.
- 8) Cover the completed stack with the Top Plate and secure as follows:
  - Raise the Clamping Latch Lever upwards to expose the Latch Hook.
  - While holding the Clamping Latch with one hand, press the lever on the Cam Buckle with the other.
  - Slide the Cam Buckle along the Strap until it can be engaged into the Clamping Latch hook.
  - Release the Cam Buckle and pull the loose end of the Strap downwards to partially tension. Do NOT over tension! The Clamping Latch lever will remain in the raised position, 20 30 degrees from the vertical when partially tensioned. Repeat these steps on the other side.

 Press both levers down, closing the Clamping Latches to secure the stack. Do not use excessive force. It may be necessary to loosen the straps slightly to secure. Repeat the previous action to release or increase the tension in the strap as necessary to ensure a firm grip.

Warning: Do not run the machine with unsecured sieves.

- 9) Set the timer for the desired test interval. For timer setting instructions please refer to item three in the Installation & Set-up Instructions at the front of this manual.
- 10) Once the test is complete, allow the sieve stack to come to a stop.
- 11) Remove the sieve stack and proceed to weigh-up the retained fractions.

  <u>Warning</u>: Do not attempt to release the Clamping Latches and remove the sieves while the stack is still in motion.

#### For More Information...

For recommendations on sampling procedures, sample size, sieve selection, calibration, test intervals, sieve care and cleaning and related topics, please see Advantech Manufacturing publication R1986AS, *Test Sieving: Principles and Procedures* at the back of this manual. Please contact your local Advantech Manufacturing representative, Advantech Manufacturing or order directly from our website www.advantechmfg.com.



# Test Sieving: Principles and Procedures

A Discussion of the Uses, Capabilities, and Limitations of Testing Sieves as Analytical Tools



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THE LEADER IN SIEVING TECHNOLOGY®

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## **Foreword**

Through ASTM and many industry organizations, standards have been established for particle size for powder, granular and larger sized materials. This manual has been prepared to help guide users of test sieves through the proper procedures as well as provide many additional tips that can enhance the existing procedures.

Our aim is to provide assistance to both the experienced and nonexperienced particle technologist in developing comprehensive particle size test results, reduce test variations and enable the user to isolate and identify sources of error or variations in the data.

Advantech Test Sieves, manufactured in the U.S.A., are the most accurate test sieves available in the world today. The use of Advantech Test Sieves will provide more precise and reproducible data, resulting in better product control and a possible reduction of variables.

In preparing this manual, we have drawn from sources in the ASTM publications, ISO Standards and various papers written by some of the most renowned figures in the particle technology world. Additionally, Advantech personnel have contributed sieving technology developments after having logged numerous years of "hands-on" experience with many experts in the field. The result is a melding of standards, research and opinion to provide a solid foundation for your own particle size analysis program.

If additional help is desired in establishing your sieve analysis procedure, or if you desire a list of suppliers of the equipment highlighted in this manual, please contact Advantech Manufacturing, Inc. 2450 S Commerce Dr., New Berlin, WI 53151

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## WHAT IS SIEVING?

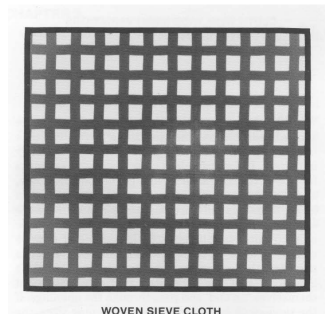
A simplistic definition of sieving is the separation of fine material from coarse material by means of a meshed or perforated vessel. Professor Terence Allen characterizes sieving as "The aperture of a sieve may be regarded as a series of gauges which reject or pass particles as they are presented to the aperture." (1) This theory was actually in practice during the early Egyptian era as grains were sized with 'sieves' of woven reeds and grasses.

The level of sophistication increased with the rise of the industrial revolution and the need for more sophisticated methods for classifying material by their particle size. As requirements for sized material rose, technology in producing uniform sieving media increased. Woven wire cloth was introduced as an alternative, providing greater accuracy and durability. At present, this woven cloth is available in a range of sizes from 125 mm (5") openings to 20 micrometer openings.

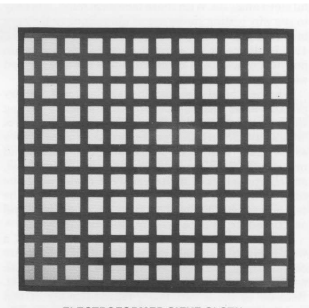
All mesh sizes are covered by both national and international standards.

The need for particle size analysis in the finer size ranges (i.e. 38 micrometers and less) prompted the development of the electrodeposited sieve. These sieves, sometimes called electroformed or micromesh, are currently being produced with openings as fine as 3 micrometers. The mesh openings are extremely uniform in both size and shape and maintain exacting tolerances.

While the technology related to sieve analysis has come a long way since the reed sieves of ancient Egypt, few new developments have come along since the 1940's. Professor Kurt Leschonski wrote "Sieve analysis is one of the few methods of particle size analysis which has escaped modernization." <sup>(2)</sup>While the modernization has not come in the actual hardware of sieving, refinements in the application and utilization of existing equipment has proceeded.



Variations in opening size and shape are common.



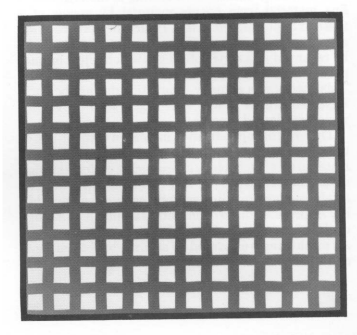
**ELECTROFORMED SIEVE CLOTH**Electrodeposited material showing uniformity in opening size and shape.

## USES, LIMITATIONS AND ADVANTAGES

Harold Heywood wrote "I often refer to sieving as the 'Cinderella' of particle size analysis methods; it does most of the hard work and gets little consideration." (3)

There are numerous reasons for the selection of high quality testing sieves as a first choice in particle size analysis work. Leschonski said "... because simplicity everyone immediately understands the purpose of a stack of operation and its -and sieves inexpensive- ness." (4) Standard sieve analysis is probably the fastest and most widely used quality control procedure in any powder process control industry. Used frequently as a mediating device between the production and sales divisions of a process corporation or between the sales force and the customer, test sieve analysis work enjoys the universal recognition of being the best 'quick and dirty' test procedure for rapid particle size distribution data. The outcome of the analysis is easily calculated and interpreted for comparison between laboratories. Start-up cost to institute a basic sieving quality control program is minimal, and operators at most levels of training are capable of performing a successful sieve analysis. With these factors in mind, it is easy to see why testing sieves are as ubiquitous as they are in industry. Materials from crushed ore chunks of over 114.3 mm (4 1/2") in diameter to slurred alumina and porcelain powders of less than 20 micrometers are all analyzed with test sieves on a regular basis.

Whether hand or machine sieving, wet or dry preparations, analysis or production work, testing sieves have found a niche in the quality control laboratory. Given this overall acceptance of test sieves as a viable analytical device and the widespread presence of the sieve in laboratories of all industries, any shortcomings of such an analytical device would be magnified. For all of the advantages available to the test sieve user, limitations must be recognized



**CLOTH WITH WIDE MESH VARIATIONS** 

Alternating areas of narrow and wide mesh openings can significantly change sieve analysis results.

and accounted for in the presentation and analysis data.

individuals. Test sieves are Beina fabricated of a woven mesh material, variations in the weave are common. The chances of locating two sieves with an identical distribution of opening sizes are extremely remote. Due to these variations, the reproducibility of test results between sieves can be adversely affected. The stringent standards imposed by ASTM, ISO or other regulating bodies have established tolerance factors which allow for the permissible variations in the weave while striving to maintain a level of uniformity in the performance of the 'test grade' sieve cloth. (See Table 1)

With this variation of opening sizes present, some smaller than the nominal and some larger, the time interval of the sieve analysis becomes extremely important. If, for example, a sieve has several openings far above the nominal opening size for the particular mesh size,

and the test is run for 30 minutes, the probability of larger-than-nominal particles finding those oversized openings is much greater than if the test was run for only 15 minutes. Similarly, if the sample of powder contains a large percentage of elongated or needle like particles, a longer test interval would provide a greater likelihood that the elongated particles will orient themselves 'on end' and pass through the openings. If the sieving cloth has a wide range of opening sizes, the sieving of this type of material has a compounded error.

Another factor which must be considered is the reaction of the material to ambient conditions. The most accurate test sieve available would be of minimal use if the relative humidity in the test lab was 99%. Extremely dry conditions can cause fine powders to adhere to the sieve

components and each other with strong electrostatic charges. Additional types of sieving problems are discussed in the glossary section.

To minimize error caused by wire cloth variation, steps must be taken at every stage of fabrication that will assure the uniformity of the woven mesh as well as compliance with the applicable standards. Both the weaver and the test manufacturer must maintain constant monitoring program measuring the actual opening sizes of the wire cloth as well as the uniformity of those openings. The loss to the manufacturers in rejected out of specification sieve cloth is a gain to the end-user in uniformity and compliance.



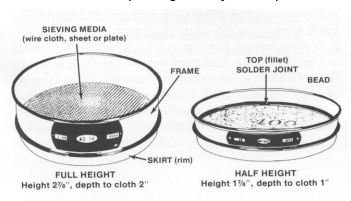
**COMPARATOR**Profile projector specially designed and built for wire cloth and sieve inspection.

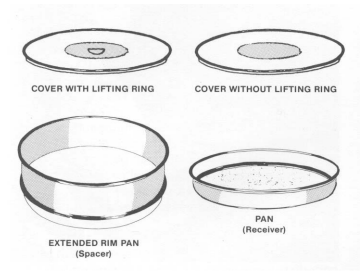
## **GLOSSARY OF SIEVING TERMINOLOGY**

Sieving terminology is frequently used and abused in writing specifications for materials. Listed below are some of the most frequently used terms and a general discussion of their meaning:

**Agglomerate:** natural tendency of materials to clump or ball together. This condition is very common in materials with high moisture, fat or oil content or those with fibrous or extremely irregular topography.

**Blinding:** plugging of the screen openings with particles either exactly the same size as the sieve opening or by fine particles



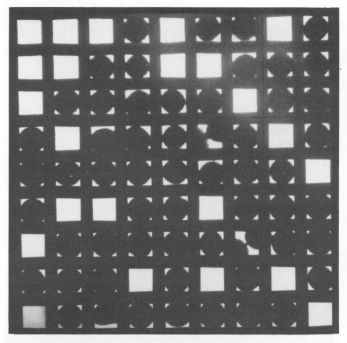


which build up on the wire mesh and eventually close off the openings. Frequently referred to as pegging. (Photo Page 4)

**Cover:** stamped or spun lid that tightly covers the top of a sieve to prevent the loss of the material sample during sifting or mechanical agitation.

**Electrostatic charges:** accumulation of electrical charges on the particles and sieve components causing clinging, agglomeration or blinding. This condition is frequently seen in hydrocarbon-based materials, plastics, reactive metals, paint pigments and powders with a large fraction finer than 20 micrometers.

**Extended rim pan:** a sieving pan with a skirt designed to nest within a sieve stack, allowing multiple tests to be performed simultaneously. Frequently called a nesting pan or spacer.



**BLINDED SIEVE** 

Spherical and near-size particles can blind or peg in the sieve openings.

Flow additive: powdered substance added to the sample to reduce agglomeration, neutralize static charges and improve the flow characteristics of the sample. Common additives are fine silica, activated charcoal, talc, and other commercially produced natural or synthetic substances. Generally, the additive is pre- screened to a known average particle size, blended with the sample (approximately 1% additive by weight) and then screened with the additives value removed from the reported data.

**Frame:** a rigid sidewall used to form the body of the testing sieve. Common depths are 50.8 mm (2" full height) for 8" sieves and 25.4 mm (1" half height). Special application sieves of other depths are also in use.

**Mesh:** screening medium with openings of uniform size and shape made of woven, punched or electrodeposited material.

**Pan:** stamped or spun receiver of materials passing through the finest sieve.

**Skirt:** section of test sieve below the sieve mesh that allows for mating or nesting of the sieves in a test stack.

**Support mesh:** coarse sieve cloth mounted under fine sieve cloth in a test sieve to provide extra strength. This is widely used in wet sieving operations to protect the fragile fine sieve cloth. Frequently called backing cloth or rolled backing cloth.

**Test Sieve:** screening medium (mesh) with openings of uniform size and shape mounted on a rigid frame, usually for laboratory testing or small scale production applications. The frames can be made of various materials, the most common of which are brass and stainless steel in a cylindrical configuration, having a diameter of 3", 5", 6", 8", 10", 12" or larger.

Wet sieving: the separation of fines from the coarse portion of a sample while suspended in an aqueous solution introduced to a testing sieve. The liquid medium is used to negate static charges, break down agglomerates and lubricate near-size particles. After the fines have been washed through the sieve, the residue is oven-dried and re-weighed.

## **CHAPTER 4**

## SIEVE SPECIFICATIONS

-Domestic and International

The U.S. Standard Sieve Series is a metric system based series first suggested by the American Society for Testing and Materials in 1913. The opening sizes in this sieve series are in the ratio of the fourth root of two. This numerical relationship was first suggested by Professor P .R. Rittinger, a German researcher, in 1867.

In the fourth root of two series, every opening size is 1.189 times the opening size of the next smaller sieve. This

relationship continues into sieve opening area measurement. The U. S. Sieve Series provides that the area of each sieve opening size is 1 1/2 times the area of the preceding sieve size.

By using every other sieve in this number series, the relationship becomes based on the square root of two (1.414), with the area of the opening being twice that of the preceding sieve size. Thus, by skipping two sizes, you create an area ratio

of 3 to 1, or by skipping three sizes, you create a ratio of 4 to 1.

When selecting sieves from this series, any number of sieves can be used for an analysis. Care must be taken in selecting each sieve between two points, every other sieve, every fourth sieve, etc., to keep within the mathematical progression of the series.

After World War II, the International Standards Organization (ISO) was formed in an attempt to establish world standards. Though the U.S. Sieve Series had proven to be effective and was in use throughout the world, members of the ISO would not accept the U.S. Sieve Series as a world standard. The ISO chose to adopt the Preferred Number Series based on the roots of ten. The Preferred Number Series

was suggested by Charles Renard of France in 1879. His system is based on the tenth, twentieth and fortieth roots of ten (designated R-10, R-20 and R-40). See Table 2.

A compromise was reached between the ISO and the proponents of the U.S. Sieve Series when it was discovered that every third value in the R-40/3 table is in a step ratio of 1.1885, sufficiently close to the fourth root of two (1.1892) used in the U.S. Sieve Series. In 1970, slight adjustments were made in the U.S. Sieve Series to align the series perfectly with the ISO specifications.

Copies of these tables of specifications can be found in Table 3.

## **CHAPTER 5**

## SIEVE CALIBRATION PROCEDURES

Quantifying and accounting variations in test sieve results become two of the most important topics in particle technology today. Once again, the ubiquitous nature of stacks of test sieves in around the powder labs world contributed to the scope of the dilemma in sieve standardization and calibration. Kaye "The states inaccuracies and uncertainties of characterization by sieve fractionation arise from the experimental problems of determining the sieve residues and from the non-ideal nature of the sieving surfaces." Further, "The presence of a range of aperture sizes in any real sieving surface is a source of error in sieve based characterization studies since the theoretical or nominal size of the sieve is taken to be the boundary limit for the sieve residue." (5)

Not only is the test sieve user plagued with variations in the weave of the cloth, but also confronted with the effects of particle shape on sieving results. 50 years ago, A.M. Gaudin wrote, "Powders with identical size distributions, densities and chemical composition may behave quite differently as a result of variations in particle shape between samples. example, powders consisting solely of spherical particles are likely to have good flow properties, while powders containing needlelike particles will not." Further, "In addition, it is impossible to isolate the concepts of particle size and shape, since the method of size measurement will influence the particle size determined." (6)

Numerous approaches have been tried to compensate for the effects of variations in wire cloth and particle shape. The methods have fallen into 3 basic categories: 1) inspection of the mesh to determine opening size, 2) material testing of the sieves to determine if sieves fall within performance specifications, and 3) a

combination of methods 1 and 2, assuring compliance with both opening size and performance specifications.

Probably the most elementary of the inspection methods is the use of the etched glass slide. This procedure relies on what is referred to as the 'Moire Effect', which compares the number of wires per inch in the wire cloth sample to the number of lines per inch etched on the glass slide. By microscopically measuring the diameters, a rough estimate of the opening size can be approximated. One major short- coming of this procedure is the assumption that all wire diameters within the sample are the same. A slight variation in wire diameter can translate to a significant change in opening size.

An alternative to this measurement approach is the use of a high-powered optical comparator or profile projector. In method. light powerful illuminate the mesh from both above and below and project the image onto a glass screen. Calibrated micrometer stages move the mesh sample in relation to a reference point allowing measurements with an accuracy of 1 micrometer to be made on both the opening and wire diameter. The results are displayed on a numerical readout. The broad field of view of the comparator allows for the scanning of a large number of sieve openings, facilitating a more comprehensive picture of the nature of the sieve cloth.

In the material testing of sieves, powder samples are run on subject sieves and the residue calculated. These values are then compared with other sieves in selecting what are often referred to as 'matched' sieves. There are a number of shortcomings in this procedure also. The first and foremost problem encountered is that of compliance. Conceivably, it is possible to find hundreds of sieves that will provide the same performance data when tested with a reference material and still not meet ASTM standards.

While the sieves perform comparably, they do not meet the basic criteria of ASTM

specifications, which should disqualify them from use as a U.S. Standard Testing sieve. Another problem encountered with material matching is the use of reference samples that are different in shape, size or density than the users' products. For example, a manufacturer of spherical steel shot would yield significantly different results on a sieve that had been matched with an angular ground silica material. In this case, both shape and density are considerably different. The key to proper matching is using the end-users own product or a material that approximates the product most closely.

The final approach is a combination of the first two methods. First, the sieve is inspected optically for compliance with all applicable standards. Openings and wire diameters are measured, not averaged. After the sieve opening distribution has been characterized and evaluated, actual material testing can begin. During the material testing, samples of the user's product are used for the standardization procedure. ΑII tests are run for repeatability and the variation between test results calculated. This procedure yields a testing sieve with known values in the two most essential parameters compliance with specifications and performance duplicate test conditions.

An alternative that has been used with some success is the use of correction factors between sieves. Once a 'master set' of sieves has been established, a reference sample is tested on the stack. The values are calculated and retained. As new sieves are acquired, the original reference sample is tested on the new set and the values calculated. Any variations between the sieve stacks can be compensated for with correction factors or multipliers. example, a sieve in stack 3 may retain more or less than the comparable sieve in the master set. A multiplier of magnitude greater than or less than 1 is necessary to calculate the comparable retention value on that sieve when compared to the master set. In this way, every sieve in use can be compared to the master set to standardize sieving results. Whatever method you use, it is essential that your starting point is based on ASTM specifications. This

compliance is necessary to assure uniformity between and within industries.

## **CHAPTER 6**

## PERFORMING THE SIEVE ANALYSIS

In obtaining meaningful sieve analysis data, six major steps are recommended. 1) Obtain a representative sample of the material to be evaluated. 2) Prepare the sample for evaluation; this may involve washing and/or drying the sample. Reduce the sample to a size suitable for the sieve analysis procedure. 4) Perform the actual sieve analysis procedure. Compute the data and convert the data into a usable format. 6) Organize the data assemble information and the presentation.

Granular and powder materials are prone to segregation during movement and storage of the products. This segregation can be due to the disparity of the particle sizes and the varied densities for blended When forming a stockpile of products. material, the larger, coarser particles are heavier and tend to roll to the lowest portion and outer perimeter of the cone. The finer particles are lighter and more angular and remain concentrated at the top and through the vertical center of the cone. Obtaining samples from only the outer perimeter or from the top of the cone would not provide a sample which would be representative of the entire batch.

Sample extraction and preparation is the most commonly overlooked variable in sieve standardization programs. Testing bias can be added at many places along the progression from the raw materials received from a supplier, samples taken at each stage of production, sample reduction procedures and samples when the product is ready for shipment to the customer. The way the samples are extracted from the original bulk volume varies with the way the materials are received, produced or stored. The ideal sampling method is one which provides the most representative sample with the least amount of material required.

The following paragraphs were first published in the ASTM technical publication STP 447 A. The collaborative efforts of the authors have produced a section on sampling technique which will aid in obtaining representative test samples from larger test sources...<sup>(7)</sup>

### Sampling from a chute or belt

Accuracy in sampling is obtained where material is flowing from a chute or belt conveyor. The ideal place to collect the sample is where the material drops from the chute or belt. If the material stream is small enough, use a pail or other suitable receptacle which can be swung completely across the flowing stream in a brief interval of time and with uniform movement. The sampling receptacle should not be allowed to overflow, because the overflow would tend to reject a higher proportion of the particles that exist larger in representative Mechanical sample. sampling devices are available for selecting samples automatically from a stream at uniform time intervals.

## Sampling from carload shipments of coarse bulk material

For coarse materials, such as crushed stone and gravel, shipped in railroad cars, a recommended method is to dig three or more trenches at least 30.48 cm (1 foot ) deep and approximately 30.48 cm foot)) wide at the bottom. Equal portions are taken at seven equally spaced points along the bottom of the trench by pushing a shovel downward into the material and not by scraping horizontally. Samples from trucks, barges, or boats should be taken in the same manner as from railroad cars, except that the number of trenches should adjusted to the size of the transportation unit and tonnage involved.

## Sampling from carload shipments of fine bulk materials

One established method for sampling a carload of bulk granular material is to take eight equal samples, (approximately 700 to 1000 grams each) from the bottom of a 30.48 cm (1 foot)) conical excavation. Samples should be suitably spaced to represent the length and width of the car and then combined into a single gross sample.

## Sampling bulk shipments of fine material with a sampling tube

An alternate and simpler method of sampling a carload, or other bulk quantity of fine or granular material is by use of a sampling tube which, for this purpose, should be 38.1 mm (1 1/2 inches ) by approximately 1.829 m (6 feet ). Five or six insertions of the tube will produce approximately, a 2 pound (907g) sample.

## Sampling from a carload of bagged material

One method of sampling a carload of material shipped in bags is to select, at random, a number of bags equal to the cube root of the total number of bags in the car and to take suitable portions (800 to 1000 grams for minus 6 mm material) from each of the selected bags for a combined gross sample.

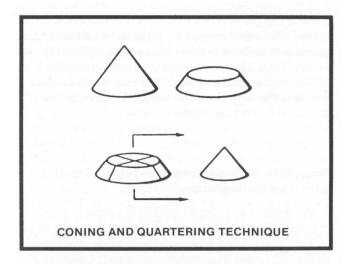
#### Sampling from a pile

In sampling from a pile, particularly material like crushed stone or containing large particles, it is extremely difficult to secure samples that are truly representative. At the apex of a conical pile, the proportion of fines will be greater, while at the base; the percentage of coarse particles will be greater. Therefore, neither location will be representative of the whole. In a shoveling process, every fifth or tenth shovel, etc., should be taken depending on the amount of the sample desired. The sample should consist of small quantities taken at random from as many parts of the pile as are accessible and taken in a

manner that the composite will have the same grading as the larger amount.

## Reduction of gross sample to test size for sieve analysis

After the gross sample has been properly obtained, the next step is to reduce it to a suitable size for sieve analysis without impairing in any way the particle size distribution characteristics of the original sample. This phase of the operation should follow the applicable procedures described in the succeeding sections and should be performed with as much care as was used in the collection of the gross sample and in performing the sieve test.



#### Coning and quartering

Pile the gross sample in a cone, place each shovel full at the apex of the cone, and allow it to run down equally in all directions. This will mix the sample. Then spread the sample in a circle and walk around the pile, gradually widening the circle with a shovel until the material is spread to a uniform thickness.

Mark the flat pile into quarters, and reject two opposite quarters. Mix again into a conical pile, taking alternate shovel-fulls from the two quarters saved. Continue the process of piling, flattening, and rejecting two quarters until the sample is reduced to the required size.

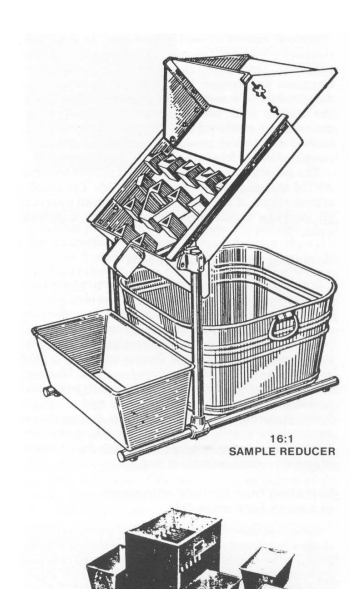
#### Sample splitters and reducers

Gross samples, if not too large, may be reduced to test sample size by one or more passes through a sample splitter or Jones type riffle, which will divide a sample in half while maintaining the particle size distribution of the original sample. By repeated passes, the sample can be split into quarters, eighths, and so on until the size of the sample desired is obtained. For larger gross samples, sample reducers are available which will select a representative 1/16 part with a single pass. After just two unit. passes through such а representative one pound sample can be obtained from an original 256 pounds. Three passes will give a one pound sample from two tons of material. Always make sure that the passages in the splitter or reducer are at least three times the size of the largest particle in the sample. Do not attempt to arrive at exactly the amount of material specified for the test. If a 50 gram sample is desired, arrive as near to this amount as practicable, because it will make no difference in the test percentage results whether the sample is slightly larger or smaller. In attempting to arrive at an exact weight, the tendency is to discriminate by the removal of sizes that are not representative of the whole, thus destroying the representative quality of the sample.

#### Size of Sample in the Test

There is a natural tendency, although incorrect, to use an excessively large sample in the test. In most cases, a smaller sample will provide a more accurate analysis. Beware, however, that the more you split, the greater the chance of error. Testing sieves are a go or no go gauge; if the sample is too large it will not permit each of the particles an opportunity to present themselves to the screen surface. Often the limiting factor for reducing the sample size is the accuracy of the weighing device used to determine the amount of material retained on the sieve.

Generally a 25 to 100 gram sample is recommended. However, if it is necessary to establish the correct sample size, utilize the following procedure: Using a sample splitter, reduce samples to weights (i.e. 25, 50, 100, 200 grams). Analyze these



various sample sizes on a selected nest of sieves for a period of five minutes preferably using a mechanical sieve shaker. If the test with the 100 gram sample shows approximately the same percentage passing the finest sieve as the 50 gram sample, whereas the 200 gram sample shows a lower percentage, this would indicate that the 200 gram sample is too large and the 100 gram samples would be satisfactory. Then run the 100 gram sample on the same set of sieves for the same time period to see if repetitive results are obtainable.

SAMPLE SPLITTER

A useful table of recommended sample sizes for tests with 200 mm or 8" diameter sieves is presented in Table 4. Note that the table gives sample sizes listed by volume. Recommended sample weights in grams can be determined by multiplying the values in Column 3 and 4 by the bulk density (grams per cubic centimeter) of the material to be tested rounded out within a reasonable tolerance. If the actual bulk density of a certain material is not known, the typical density factor for the most nearly similar material listed in Table 5 may by used.

To perform the actual sieve analysis, sieves should be chosen in a sequence as described earlier. Use every sieve, every other sieve, or every third sieve, etc. between the desired size parameters. The use of sieves in this sequential order will allow for better data presentation and a more meaningful analysis of the test results. Care should also be taken in selecting the proper sieves to avoid overloading any sieve with an especially large material peak. For example, a specification may require 96% of the sample be retained above a #50 mesh The proper way to perform an analysis of this nature is to use 'relief screen', that is, sieves in the 30, 35, 40 and 45 mesh ranges to remove some of the burden from the critical cut point of 50 mesh. If the relief sieves are not used, the particles of exactly 50 mesh size or slightly larger may become wedged in or forced through the sieve openings by the mass of material resting above them. Large concentrations of material on one sieve reduce the opportunity for near sized material to pass through the sieve resulting in a larger portion of the material retained on the test sieve. The sieve cut point would be inaccurate and the sample would not meet the specifications for the test.

The selected sieves should assembled with the coarsest sieve at the top of the stack and the balance of the stack in increasing magnitude of fineness (increasing sieve numbers with smaller openings). The stack should include a cover on the top sieve and a pan below the finest sieve. The sieve stack can be shaken then rapped by hand or mounted in sieve shaker with a motorized electrostatic drive mechanism.



While many applications still use the handshaken method for sieving, motor driven shakers have proven to be much more consistent, minimizing variations related to operator procedures. In powder analysis below the 100 mesh range, the sieve shaker should be equipped with a device to impart a shock wave to the sieve stack at regular intervals. This hammer or rapping device is necessary to reorient the particles on the sieve and impart some shear forces to near-sized particles blocking the sieve openings.

#### **Recommended Time Intervals**

The duration of the sieving interval is usually regulated by industry standards, or in-house control specifications. Commonly, 10, 15 or 20 minute tests are used as arbitrary sieving intervals. determine the best interval for a new material, or to double check the accuracy of existing specifications, the following procedure can be used. Select the desired sieves for the analysis. 1) Weigh up a sample of the material to be tested and introduce it to the complete sieve stack. 2) Shake the sieve stack for a period of 5 minutes. 3) Weigh the residue in the pan and calculate the percentage in relation to the starting weight. 4) Reassemble the stack and shake for one additional minute. 5) Repeat the weigh-up procedure and percentage. calculate the lf the percentage of fines increased more than 1% between 5 minutes and 6 minutes, reassemble the stack and shake for an additional minute. The data can be plotted as percentage throughput versus time for each data point you calculate. When the change in the percentage of fines passing in the 1 minute period drops below 1%, the test can be considered complete. Record the total testing time for subsequent analyses.

Another type of sieve analysis is the wet sieve test. In this method, the sample is weighed and then washed through the finest sieve in the stack with water, a wetting agent (water based), or some other compatible solvent. After thoroughly washing the fines from the raw sample, the residue is dried either over a hot plate or in an oven. The temperature of the sieve should be maintained below (300°F)<sup>1</sup> to avoid loosening of the sieve cloth or failure of the solder joint. After drying, the residue is then sieved normally on the balance of the sieve stack. The loss in weight not accounted for on the coarse screens is assumed to be fines or soluble material.

Wet sieve analysis is especially helpful when working with naturally agglomerated materials, ultra-fine powders with severe static changes and in samples where fine particles tend to cling to the coarse fractions in the blend. The disadvantages associated with wet sieving are primarily the time period required to perform the analysis due to the additional washing and drying time and the possible damage to the sieve mesh by overloading. A common practice with wet sieving operations is brushing or forcing the sample through the mesh while the liquid medium is directed on the sieve. This pressure can distort the sieve openings or tear the mesh at the solder joint through stress. Therefore, this procedure is not recommended. Once the sieving interval is complete, whether dry or wet sieving is used, the residue on each sieve is removed by pouring the residue into a suitable weighing vessel. To remove material wedged in the sieve's openings, the sieve is inverted over a sheet of paper or suitable collector and the underside of the wire cloth brushed **gently** with a nylon paint brush with bristles cut to a 25.4 mm (1") length. The side of the sieve frame may be tapped gently with the handle of the brush to dislodge the particles between brush strokes. At no time should a needle or other sharp object be used to remove the particles lodged in the wire cloth.

Special care should be taken when brushing sieves finer than 80 mesh. Brushing can cause distortions and irregularities in the sieve openings. The procedure is repeated for each sieve in the stack and contents of the pan.

The individual weights retained on the sieves should be added and compared to the starting sample weight. Wide variations or sample losses should be determined immediately. If the finished sample weight varies more than 2% from the initial weight, the analysis and sample should be discarded and the test performed another sample. If the sample weights are acceptable, complete the calculations and report the individual weights retained on each sieve.

Presentation and analysis resulting data is frequently made easier by plotting on one of a number of graph formats. The most common graphic presentation is the plotting of cumulative percentage of material retained on a sieve (plotted on a logarithmic scale) versus percentage (plotted on a linear scale). The resulting curve allows a quick approximation of the sieve size at the fiftypercentile point of accumulation. The curve shows the smoothness distribution by revealing the presence of bimodal blends in the sample. Other plotting techniques include log-log and direct plotting of micron size versus percentage retained.



<sup>&</sup>lt;sup>1</sup> Advantech metal framed sieves should not exceed 261° F (127° C). Solder will begin to soften at this point.

Care should be exercised in the analyzing the data in relation to the length of time the test was run. If the sample contains a large amount of elongated or near- size particles, the test results can be misleading. The longer the sieving interval, the greater the opportunity for these problem particles to pass through the sieve's openings. Ideally each fraction

should be inspected microscopically after sieving to determine the integrity of the sieve cut point.

Table 6 lists many of the ASTM published standards on sieve analysis procedures for specific materials or industries.

## **CHAPTER7**

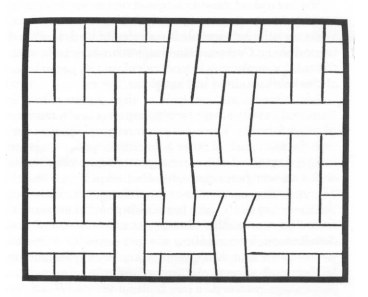
## SIEVE CARE AND CLEANING

Test sieves, like any other piece of analytical laboratory equipment, require regular care to maintain their performance standards. Sieves should be kept clean and dry at all times, and stored either in the cardboard carton provided or in a suitable cabinet. The wire cloth must be taut and free from variations in opening size. For this reason, cleaning procedures must be clearly delineated as part of a comprehensive sieving program.

Test sieves should be cleaned ultrasonically on a regular basis.\* For some installations, this may be done at the end of a shift or at the end of a week, but must be done regularly to assure accurate sieving results. The sieves should be immersed in an ultrasonic cleaner filled with a solution of a mild detergent and water. Prior to reuse, ensure that the test sieves are dried thoroughly. Ultrasonic cleaning prevents the buildup of particles trapped in the sieve openings and prolongs the useful life of the sieve. Between test clean-up, brushing of the mesh, sizes 100 and coarser, is recommended. For best results, use a nylon bristle paint brush with the bristles cut to а length approximately 25.4 mm (1"). The sieve openings should be brushed from the underside only with a gentle circular

motion. Vigorous brushing will distort the sieve openings and reduce the effective life of the sieve. Particles lodged in the sieve openings should never be removed with a sharp object. These particles should be removed in an ultrasonic cleaner only. Brushing should be avoided on sieves finer than 100 mesh, as the fine wires are more likely to bend, distort or even break. Brushing can often loosen the wire cloth; the finer mesh sizes are most susceptible to this damage.

Similarly, cleaning sieves with a compressed air jet is common, but this can



WIRE CLOTH DAMAGED BY IMPROPER BRUSHING
Note the irregularities in both opening size and shape.

 $<sup>^2</sup>$  \*Do  $\underline{\rm NOT}$  ultrasonically clean precision electroformed test sieves. Refer to the Handling and Use Instructions on the sieve jewel case.

damage the sieve openings on the finer mesh sieves. The concentrated jet of air can cause severe 'local' damage to the wire cloth, and significantly reduce the accuracy of the sieve mesh.

With proper care, sieves will perform accurately for many years. Typical wear does not usually change the opening sizes, but can abrade the 'knuckles' or crimps of the wire. A sieve with noticeable sagging of the cloth should be replaced. Fine mesh sieves that are torn should not be resoldered, as the localized heat of the soldering iron can distort the openings. Epoxies have been used for repairs, but

tend to block a large percentage of the openings reducing the opportunity for particles to pass through the openings in the allotted agitation time. Epoxies may become too brittle for the flexing of the wire cloth and can fracture with use.

Good general laboratory procedures should be observed with testing sieves as with any other piece of test equipment. Testing should be performed with clean, uncontaminated sieves, especially when using a sieve for the first time. With proper care and cleaning coupled with a good calibration procedure, any test sieve should provide many years of consistent service.

## **EPILOG**

We hope that the characterization of testing sieves and their uses presented in this manual will serve as an enhancement to your current particle size analysis program. By maximizing the analytical advantage potential of testing sieves while minimizing and compensating for shortcomings and inaccuracies, the testing sieve can be a viable and precise testing tool. Care, maintenance and proper test procedures are as critical with a testing sieve as they are with other, more sophisticated particle size analyzers.

Compliance with applicable industry, national and international specifications is essential. The intent of these regulating bodies is the formulation of general standards to assure uniformity in testing standards observed by both the buyer and producer. The accepted specification should be the foundation for the in-house testing procedure.

Testing accuracy is highly dependent on the technique of the operators. Interpretation of data should be neither overstated nor understated in terms of importance. The effects of variables must be understood, accepted and factored into final data analysis to avoid these shortcomings.

NOTE: To aid in making this manual as understandable and comprehensive as possible, minor changes in spelling and grammar have been made to some of the quoted references. These changes have not altered the statements made but have aided in clarifying the thoughts of the authors.

## **BIBLIOGRAPHY**

- Allen, Terence, Particle Size Measurement, Chapman and Hall, New York 1981.
- Leschonski, Kurt "Sieve Analysis, The Cinderella of Particle Size Analysis Methods?", Powder Technology, Elsevier Sequoiz S.A. , Lausanne, 24 (1979)
- Heywood, Harold, Proc Particle Size Analysis Conference, Bradford,
   1970.
- 4. Leschonski, Kurt, Ibid.
- 5. Kaye, Brian, Direct Characterization of Fine Particles, John Wiley and Sons, New York, 1981.
- Gaudin, A.M. Principles of Meneral Dressing, McGraw-Hill, New York 1939.
- 7. Manual on Test Sieving Methods-STP 447 A, American Society of Testing and Materials, Philadelphia, 1969.

## STANDARD SPECIFICATION FOR WOVEN WIRE TEST SIEVE CLOTH AND TEST SIEVES ASTM E11 - 15

Nominal Dimensions and Permissible Variations for Sieve Cloth and Compliance, Inspection and Calibration Test Sieves (1) (6)(13)(14)(15)Permissible Range + X Resulting Sieve Designation ±Υ of Choice Nominal Sieve Maxim um Maximum Typical Variation for Opening (in.) Variation for Individual Wire Diameter Standard Alternative Average Opening Max Opening Opening millimeter millimeter millimeter millimeter millimeter inches 125 5 in. 5 3.66 4.51 129.51 8 6.8 9.2 4.24 in. 5.4 106 4.24 3.12 3.99 109.99 6.3 7.2 100 4 in. 2.94 3.82 103.82 6.3 5.4 7.2 90 3 1/2 in. 3.5 2.65 3.53 93.53 6.3 5.4 7.2 75 3.09 78.09 5.4 7.2 3 in. 3 2.22 6.3 63 2 1/2 in. 2.5 1.87 2.71 65.71 5.6 4.8 6.4 53 2.12 in. 2.12 1.58 2.39 55.39 5 4.3 5.8 2.29 50 2 in. 2 1.49 52.29 5 4.3 5.8 1 3/4 in 47.12 3.8 45 1.75 1.35 2.12 4.5 5.2 37.5 1 1/2 in. 1.5 1.13 1.85 39.35 4.5 3.8 5.2 31.5 1 1/4 in. 1.25 0.95 1.63 33.13 4 3.4 4.6 3.55 26.5 1.06 in. 1.06 0.802 1.44 27.94 3 4.1 25 1.00 in. 0.758 1.38 26.38 3.55 3 4.1 1 22.4 0.875 1.27 7/8 in. 0.681 23.67 3.55 3 4.1 19 3/4 in. 0.750 0.579 1.13 20.13 3.15 2.7 3.5 0.490 0.99 2.7 3.6 16 5/8 in. 0.625 16.99 3.15 13.2 0.530 in. 0.530 0.406 0.86 14.06 2.8 2.4 3.2 12.5 1/2 in. 0.500 0.385 0.83 13.33 2.5 2.1 2.9 11.2 7/16 in. 0.438 0.346 0.77 11.97 2.5 2.1 2.9 9.5 3/8 in. 0.375 0.295 0.68 10.18 2.24 1.9 2.6 8 5/16 in. 0.312 0.249 0.60 8.60 2 1.7 2.3 6.7 0.265 in. 0.265 0.210 0.53 7.23 1.8 1.5 2.1 0.250 0.51 1.5 2.1 6.3 1/4 in. 0.197 6.81 1.8 5.6 No. 3 1/2 0.223 0.176 0.47 6.07 1.6 1.3 1.9 4.75 0.41 1.9 No. 4 0.187 0.150 5.16 1.6 1.3 4 No. 5 0.157 0.127 0.37 4.37 1.4 1.2 1.7 3.35 0.32 1.25 1.06 No. 6 0.132 0.107 3.67 1.5 2.8 No. 7 0.110 0.090 0.29 3.09 1.12 0.95 1.3 2.36 No. 8 0.0937 0.076 0.25 2.61 0.85 1.15 1 No. 10 2 0.0787 0.065 0.23 2.23 0.9 0.77 1.04 1.7 No. 12 0.0661 0.056 0.20 8.0 0.68 0.92 1.90 1.4 No. 14 0.0555 0.046 0.18 1.58 0.71 0.6 0.82 1.18 No. 16 0.0469 0.040 0.16 1.34 0.63 0.54 0.72 0.0394 No. 18 0.034 0.14 1.14 0.56 0.48 0.64 1 micrometer inches micrometer micrometer micrometer millimeter No. 20 0.0331 127 0.43 0.58 850 29.1 977 0.5 710 0.38 No. 25 0.0278 24.7 112 822 0.45 0.52 600 No. 30 0.0234 21.2 101 701 0.4 0.34 0.46 0.0197 0.315 0.27 500 No. 35 18.0 89 589 0.36 425 No. 40 0.0165 15.5 81 506 0.28 0.24 0.32 No. 45 355 0.224 0.19 0.0139 13.3 72 427 0.26 300 No. 50 0.0117 11.5 65 365 0.2 0.17 0.23 250 No. 60 0.0098 9.9 58 308 0.16 0.13 0.19 212 0.12 8.7 No. 70 0.0083 52 264 0.14 0.17 180 No. 80 0.0070 7.6 47 227 0.125 0.106 0.15 150 No. 100 0.0059 6.6 43 193 0.1 0.085 0.115 125 No. 120 0.0049 5.8 38 163 0.09 0.077 0.104 106 No. 140 0.0041 5.2 35 141 0.071 0.06 0.082 90 No. 170 0.0035 4.6 32 122 0.063 0.054 0.072 75 No. 200 0.0029 4.1 29 104 0.05 0.043 0.058 63 No. 230 0.0025 3.7 26 89 0.045 0.038 0.052 No. 270 0.0021 0.036 0.031 0.041 53 3.4 24 77 45 No. 325 0.0017 3.1 22 67 0.032 0.027 0.037 38 No. 400 0.0015 2.9 20 58 0.03 0.024 0.035 32 2.7 18 50 No. 450 0.0012 0.028 0.023 0.033 25 0.0010 2.5 16 41 0.025 0.021 0.029 No. 500 20 No. 635 0.0008 2.3 15 35 0.02 0.017 0.023 Column 3 - These numbers are only approximate but are in use for reference; the sieve shall be identified by the standard designation in millimeters or micrometers.

## INTERNATIONAL STANDARDS ORGANIZATION (ISO PREFERRED NUMBER SERIES

Values in millimeters unless specified as micron  $(\mu)$ .

R 20/3	R 20	* R 40/3	Equivalent in		
			inches		
125	125	125	4.921		
	112		4.409		
		106	4.173		
	100		3.937		
90	90	90	3.543		
	80		3.150		
		75	2.953		
	71		2.795		
63	63	63	2.480		
	56		2.205		
		53	2.087		
	50		1.969		
45	45	45	1.772		
	40		1.575		
		37.5	1.476		
	35.5		1.398		
31.5	31.5	31.5	1.240		
	28		1.102		
		26.5	1.043		
	25		0.984		
22.4	22.4	22.4	0.882		
	20		0.787		
		19	0.748		
	18		0.709		
16	16	16	0.630		
	14		0.551		
		13.2	0.520		
	12.5		0.492		
11.2	11.2	11.2	0.441		
	10		0.394		
		9.5	0.374		
	9		0.354		
8	8	8	0.315		
	7.1		0.280		
		6.7	0.264		
	6.3		0.248		
5.6	5.6	5.6	0.220		
	5		0.197		
		4.75	0.187		
	4.5		0.177		
4	4	4	0.157		
	3.55		0.140		
		3.35	0.132		
	3.15		0.124		
2.8	2.8	2.8	0.110		
	2.5		0.098		
		2.36	0.093		
	2.24		0.088		
2	2	2	0.079		
	1.8		0.071		

R 20/3	R 20	* R 40/3	Equivalent in		
K 20/0	11 20		inches		
		1.7	0.0669		
	1.6		0.0630		
1.4	1.4	1.4	0.0551		
	1.25		0.0492		
		1.18	0.0465		
	1.12		0.0441		
1	1	1	0.0394		
	900µ		0.0354		
		850µ	0.0335		
	800µ		0.0315		
710µ	710µ	710µ	0.0280		
	630µ		0.0248		
		600µ	0.0236		
	560µ		0.0220		
500µ	500µ	500µ	0.0197		
	450µ	- 1 months = 1 mm.	0.0177		
		425µ	0.0167		
	400µ	•	0.0157		
355µ	355µ	355µ	0.0140		
	315µ	771341301 <b>3</b> 00	0.0124		
		300µ	0.0118		
	280µ		0.0110		
250µ	250µ	250µ	0.0098		
	224µ		0.0088		
		212µ	0.0083		
	200µ		0.0079		
180µ	180µ	180µ	0.0071		
	160µ		0.0063		
	тоор	150µ	0.0059		
1	140µ	TOOR	0.0055		
125µ	125µ	125µ	0.0049		
1204	112µ	IZOM	0.0044		
	пи	106µ	0.0042		
1	100µ	ΙΟΟμ	0.0039		
90µ	90µ	90µ	0.0035		
- 50μ	80µ	ООД	0.0033		
$\vdash$	υυμ	75µ	0.0030		
	71µ	γSμ	0.0038		
63µ	63µ	63µ	0.0025		
σομ	56µ	σσμ	0.0023		
	σομ	53µ	0.0022		
<del>                                     </del>	50µ	υυμ	0.0021		
45	45μ	45	0.0020		
45µ	40µ	45µ	0.0018		
$\vdash$	·πυμ	38µ	0.0016		
R'10	36µ	оор	0.0013		
2011/2006	ουμ		SAME TO SOUTH STORM		
32µ			0.0013 0.0010		
25μ 20μ			0.0010		
20μ			0.0000		

<sup>\*</sup> Same as ASTM E 11 USA Standard Sieve Series

R'10 = Tenth root of ten ratio

R 20 = Twentieth root of ten

R 20/3 = Every third number of R 20 Series

R 40/3 = Every third number of fortieth root of ten series

## COMPARISON TABLE INTERNATIONAL TEST SIEVE SERIES

INTERNATIONAL ISO 3310-1:2000	AMER ASTM I	200000000000000000000000000000000000000		TISH 0:2000	CANADA CGSB-8.2-M88		NCE IFX11-501	GERI DIN (ISO) 3		JAPAN JIS
Aperture mm	Opening mm	Equiv. inch/No.	Aperture mm	Equiv. BS Mesh	Aperture mm	Aperture mm	Tamis No.	Aperture mm	Approx. DIN No.	Aperture mm
125.00	125.00	5"	125.00							
112.00			112.00							
106.00	106.00	4.24"	106.00							
100.00	100.00	4"	100.00		100.00	100.00		100.00		
90.00	90.00	3 1/2"	90.00	33	90.00	90.00	0	90.00		5
80.00			80.00							
75.00	75.00	3"	75.00							
71.00			71.00		71.00	71.00		71.00		71.00
63.00	63.00	2 1/2"	63.00		63.00	63.00		63.00		90 WEST 5 10
56.00			56.00							
53.00	53.00	2.12"	53.00		53.00	53.00		53.00		
50.00	50.00	2"	50.00		50.00	50.00		50.00		50.00
45.00	45.00	1 3/4"	45.00		45.00	45.00		45.00		00.00
40.00	40.00	, 0, 1	40.00		10.00	40.00		40.00		
37.50	37.50	1 1/2"	37.50							
35.50	07.00	1 1/2	35.50	i i		<b>t</b>				
31.50	31.50	1 1/4"	31.50		31.50	31.50		31.50		
28.00	01.00	LOUT	28.00		28.00	28.00		28.00		
26.50	26.50	1.06"	26.50		20.00	20.00		20.00		26.50
25.00	25.00	1.00"	25.00		25.00	25.00		25.00		20.00
22.40	22.40	7/8"	22.40	<del>                                     </del>	22.40	22.40		22.40		22.40
20.00	22.40	110	20.00		20.00	20.00		20.00		22.40
19.00	19.00	3/4"	19.00		20.00	20.00		20.00		19.00
18.00	19.00	3/4	18.00			18.00		10.00		15.00
and the second second	16.00	5/8"			16.00			18.00 16.00		16.00
16.00	16.00	5/6	16.00	3	16.00	16.00	2 0			16.00
14.00	46.00	E0.011	14.00		14.00	14.00		14.00		
13.20	13.20	.530"	13.20		72 220	10.50		40.50		10.55
12.50	12.50	1/2"	12.50		12.50	12.50		12.50		12.50
11.20	11.20	7/16"	11.20		11.20	11.20		11.20		11.20
10.00			10.00	7	10.00	10.00		10.00		
9.50	9.50	3/8"	9.50							9.50
9.00	74270 males (442	1207000000000	9.00		Marine de la constante de la c	9.00		9.00		027094040404
8.00	8.00	5/16"	8.00		8.00	8.00		8.00		8.00
7.10			7.10		7.10	7.10		7.10		
6.70	6.70	.265"	6.70							6.70
6.30	6.30	1/4"	6.30	*	6.30	6.30		6.30		
5.60	5.60	No.3 1/2	5.60	3	5.60	5.60		5.60		
5.00			5.00			5.00	38	5.00		5.00
4.75	4.75	No.4	4.75	3 1/2						
4.50			4.50			4.50		4.50		
4.00	4.00	No.5	4.00	4	4.00	4.00	37	4.00	2E	
3.55			3.55	5		3.55		3.55		
3.35	3.35	No.6	3.35							
3.15			3.15		3.15	3.15	36	3.15		
2.80	2.80	No.7	2.80	6	2.80	2.80		2.80		
2.50			2.50		2.50	2.50	35.00	2.50		
2.36	2.36	No.8	2.36	7		DUNCKTORE	10-8753803	FOLOTO \$ 3450		
2.24	J encountries	110000000000000	2.24	10		2.24		2.24		
2.00	2.00	No.10	2.00	8	2.00	2.00	34	2.00	3E	
1.80	42,000,000	994/4/1874/1757	1.80	39	YV650534450	1.80	C550V	1.80	43700	
1.70	1.70	No.12	1.70	10		V250004E)		Ø15¢Ma∓.		
1.60			1.60		1.60	1.60	33	1.60		
1.40	1.40	No.14	1.40	12	1.40	1.40	5.7	1.40		1.40
1.25	110	. 10. 17	1.25	"-	1.40	1.25	32	1.25		1.40
1.18	1.18	No.16	1.25	14		1.20	32	1.20	5	
1.12	1.10	140.10		14	1.12	1.12		1.12		
	1.00	No 10	1.12	10			24		6	
1.00	1.00	No.18	1.00	16	1.00	1.00	31	1.00	6	
900µ	050	NI - 00	900µ	,,		900µ		900µ		050
850µ 800µ	850µ	No.20	850µ	18	000	000	22	000		850µ
		ı	800µ	ı I	800µ	800µ	30	800µ		L

## COMPARISON TABLE INTERNATIONAL TEST SIEVE SERIES

INTERNATIONAL ISO 3310-1:2000		RICAN E 11-01		TISH 0:2000	CANADA CGSB-8.2-M88		NCE NFX11-501	12-20-11-00-00-0	MANY 310-1:2000	JAPAN JIS
Aperture mm	Opening mm	Equiv. inch/No.	Aperture mm	Equiv. BS Mesh	Aperture mm	Aperture mm	Tamis No.	Aperture mm	Approx. DIN No.	Aperture mm
630µ 600µ 560µ	600µ	No.30	630µ 600µ 560u	25		630µ 560µ	29	630µ 600µ 560µ	10	600µ
500µ	500µ	No.35	500μ 500μ	30	500µ	500μ 500μ	28	500μ 500μ	12	500µ
450μ 425μ 400μ	425µ	No.40	450μ 425μ 400μ	36	400ր	450μ 400μ	27	450μ 430μ 400μ	14 16	425µ
466µ 355µ 315µ	355µ	No.45	355µ 315µ	44	355µ 315µ	355µ 315µ	26	466µ 355µ 315µ	0	355µ
300µ 280µ	300µ	No.50	300µ 280µ	52	252	280µ	0.5	300µ 280µ	20	300µ
250µ 224µ 212µ	250µ 212µ	No.60 No.70	250µ 224µ 212µ	60 72	250µ	250µ 224µ	25	250μ 224μ	24	250µ 212µ
200μ 180μ	180µ	No.80	200µ 180µ	85	200µ 180µ	200μ 180μ	24	200µ 180µ	30	180µ
160μ 150μ 140μ	150µ	No.100	160µ 150µ 140µ	100	140µ	160µ 140µ	23	160µ 150µ 140µ	40	150µ
125µ 112µ	125µ	No.120	125µ 112µ	120	125µ	125µ 112µ	22	125µ 112µ	50	125µ
106µ 100µ 90µ	106µ 90µ	No.140 No.170	106µ 100µ 90µ	150 170	100µ 90µ	100µ 90µ	21	100µ 90µ	60 70	106µ 90µ
80μ 75μ 71μ	75µ	No.200	80µ 75µ 71µ	200	71µ	80µ	20	80μ 75μ 71μ	80	75µ
71µ 63µ 56µ	63µ	No.230	63µ 56µ	240	7 гр 63µ 56µ	71µ 63µ 56µ	19	7 гр 63µ 56µ	110	63µ
53µ 50µ 45µ	53µ 45µ	No.270 No.325	53µ 50µ 45µ	300 350	45µ	50µ 45µ	18	50μ 45μ	120	53µ 45µ
40μ 38μ	43μ 38μ	No.400	40µ 38µ	400	70µ	40µ	17	40µ	i,	43μ 38μ
36µ 32µ 25µ 20µ	32µ 25µ 20µ	No.450 No.500 No.635	36µ 32µ 25µ 20µ	440	36µ	36µ 32µ 25µ 20µ		36µ 32µ 25µ 20µ	130 200	32µ

## RECOMMENDED REPRESENTATIVE BULK VOLUMES OF TEST SAMPLES

Used in 8" (203mm) Testing Sieves

Standard Sieve Design	ation	В	sulk Volume of Material
		Recommended Volume	Maximum Permitted
		of Material for Test	Volume on Sieve on
Standard	Alternate	Sample	Completion of Sieving
25.0mm	1.00"	1800cm³	900cm³
22.4mm	7 <i>/</i> 8''	1600cm³	800cm <sup>3</sup>
19.0mm	3/4"	1400cm³	700cm³
16.0mm	5/8"	1000cm³	500cm <sup>3</sup>
12.5mm	1/2"	800cm³	400cm <sup>3</sup>
11.2mm	7/16''	800cm³	400cm <sup>3</sup>
9.50mm	3/8"	600cm³	300cm <sup>3</sup>
8.00mm	5/16"	500cm³	250cm <sup>3</sup>
6.30mm	1/4''	400cm³	200cm <sup>3</sup>
5.60mm	No. 3 1/2	400cm³	200cm <sup>3</sup>
4.00mm	No. 5	350cm³	150cm³
2.80mm	No. 7	240cm³	120cm³
2.00mm	No. 10	200cm <sup>3</sup>	100cm <sup>3</sup>
1.40mm	No. 14	160cm³	80cm <sup>3</sup>
1.00mm	No. 18	140cm³	70cm <sup>3</sup>
710µ	No. 25	120cm³	60cm³
500µ	No. 35	100cm³	50cm <sup>3</sup>
355µ	No. 45	80cm³	40cm <sup>3</sup>
250µ	No. 60	70cm³	35cm³
180µ	No. 80	60cm³	30cm³
125µ	No. 120	50cm³	25cm³
90µ	No. 170	40cm <sup>3</sup>	20cm <sup>3</sup>
63µ	No. 230	35cm³	17cm³
45µ	No. 325	30cm³	15cm³
38µ	No. 400	25cm³	12cm³

The recommended weight of material for a sieve test sample is calculated by multiplying the bulk volume figure in Column 3 by the particular bulk density in grams per cubic centimeter of the material, rounded out within a tolerance of ±25 percent.

BULK DENSITY OF PULVERIZED MATERIALS IN FREELY POURED CONDITION<sup>a</sup>

<u> </u>	lbs./ft.³	e Weight g/cm³	Material	lbs./ft.3	e Weight g/cm³	Material	lbs./ft.3	e Weight g/cm³
Alumina	44	1.23	Fullers earth	30 to 40	0.48 to 1.04	Rubber, chopped	36	0.58
Aluminum, calcined	128	2.05	Garnet	168	2.69	Rubber, ground	20	0.32
Aluminum oxide	122	1.96	Glass beads	76	1.22	Phosphate rock	75 to 85	1.20 to 1.36
Aluminum shot	96	1.54	Glass, crushed	66	1.06	Salt, flake	61	0.98
Ammonium nitrate	48	0.77	Glass cullet	93	1.49	Salt, rock	66	1.06
Ammonium - sulfate	61	0.98	Granite, crushed	95 to 100	1.52 to 1.60	Salt, table	75	1.20
Bauxite ore	75 to 85	1.20 to 1.36	Gravel	90 to 100	1.44 to 1.60	Sand	90 to 100	1.44 to 1.60
Bentonite	50 to 65	0.80 to 1.04	Gypsum, calcined	58	0.93	Sand, silica	90 to 100	1.44 to 1.60
Bicarbonate of soda	57	0.91	Gypsum, crushed	90 to 100	1.44 to 1.60	Sawdust	18	0.29
Borax	50 to 61	0.80 to 0.98	Iron ore	120 to 150	1.92 to 2.40	Seacoal	42	0.67
Boric acid	58	0.93	Kaolin	160	2.56	Shale	100	1.60
Calcite	90	1.44 to 1.68	Kyanite	68	1.09	Shot, metal	230	3.69
Calcium carbide	75	1.20	Lime, ground	60	0.96	Silica, flour	27	0.43
Calcium carbonate	49	0.79	Lime, hydrated	25	0.40	Silica, gel	45	0.72
Calcium chloride	64	1.03	Limestone, crushed	85 to 100	1.36 to 1.60	Soapstone, pulverized	40	0.64
Calcium phosphate	57	0.91	Limestone, agricultural	70	1.12	Soda ash, light	25 to 35	0.40 to 0.56
Carbon black	24	0.33	Magnesite	106	1.70	Soda ash, hea∨y	55 to 65	0.88 t 1.04
Cellulose powder	16	0.26	Magnetite	155	2.49	Soda, bicarbonate	57	0.91
Cement, portland	90 to 100	1.44 to 1.60	Manganese ore	120 to 136	1.92 to 2.18	Sodium nitrate	78	1.25
Cement clinker	75 to 80	1.20 to 1.28	Marble, crushed	90 to 95	1.44 to 1.52	Sodium phosphate	43	0.69
Chrome ore	140	2.25	Metals, powdered			Sodium sulfate	96	1.54
Clay	30 to 75	0.48 to 1.20	Aluminum	80	1.28	Steel grit	228	3.66
Coal, anthracite	55	0.88	Copper	169	2.71	Stone, crushed	85 to 95	1.36 t 1.52
Coal, bituminous	50	0.88	Copper-lead	364	5.84	Sugar, granulated	5	0.80
Coke breeze	25 to 35	0.40	Iron	243	3.90	Sugar, powdered	37	0.59
Coke, petroleum	25 to 40	0.40 to 0.64	Nickel	263	4.22	Sulphur, crushed	50 to 65	0.80 t 1.04
Copper ore	100 to 150	1.60 to 2.40	Stainless steel	240	3.85	Talc, powder	34	0.55
Coquina shell	80	1.28	Tantalum	300	4.80	Talc, granular	44	0.71
Corn starch	40	0.64	Mica	42	0.67	Traprock, crushed	105 to 110	1.68 to 1.76
Diatomaceous earth	31	0.50	Ore, sintered	144	1.83	Triple superphosphate,		
Dicalcium phosphate	64	1.03	Oyster shells, ground	29	0.47	granular	64	1.03
Dolomite, crushed	90 to 100	1.44 to 1.60	Perlite ore	65 to 75	1.04 to 1.20	Tungsten carbide	550	8.82
Feldspar, crushed	65 to 84	1.04 to 1.35	Plaster, calcined	64	1.03	Urea prills	43	0.69
Ferrophosphorous	196	3.14	Polyethylene pellets	36	0.58	Vermiculite ore	80	1.28
Fire clay	80	1.28	Polyethylene powder	18	0.29	Wood chips	13	0.21
Flour, wheat	24	0.38	Poly vinyl chloride	30	0.48	Zinc dust	144	2.31
Flour, maize	37 90 to	0.59 1.44 to	Potash	77	1.23	Zirconium oxide	200	3.22
Fluorspar	120	1.92	Potassium carbonate	79	1.27	Zirconium sand	162	2.60

<sup>&</sup>lt;sup>a</sup> - Where a single figure is given, it represents an actual weight of a typical average sample of the material recorded by a research laboratory; therefore, the figure can be expected to vary from sample to sample of the same material.

## LIST OF ASTM PUBLISHED STANDARDS ON SIEVE ANALYSIS PROCEDURES FOR SPECIFIC MATERIAL OR INDUSTRIES

Material	ASTM Designation	Title of Standard	Sieve No. or Size Range
Aggregates	C117-95	Standard Test Method for Materials Finer Than 75-µm (No.200) Sieve in Mineral	No.200
, iggi ogutoo	0111 00	Aggregates by Washing	110.200
	C136-01	Standard Test Method for Sieve Analysis of Fine and Coarse Aggregates	3½ in No.200
	C142-97	Standard Test Method for Clay Lumps and Friable Particles in Aggregates	1½ in No.20
	C330-00	Standard Specifications for Lightweight Aggregates for Structural Concrete	1 in No.100
	C331-01	Standard Specifications for Lightweight Aggregates for Concrete Masonry Units	¾ in No.100
	D4791-99	Standard Test Method for Flat Particles, Elongated Particles, or Flat and	
		Elongated Particles in Coarse Aggregate	
	D5821-01	Standard Test Method for Determining the Percentage of Fractured Particles in	
		Coarse Aggregate	
Asbestos	D2589-88 (1997)	Standard Test Method for McNett Wet Classification of Duel Asbestos Fiber	No.4 - No.325
	D2947-88 (1997)	Standard Test Method for Screen Analysis of Asbestos Fibers	110.1 110.020
O		5.	N = 400
Carbon black	D1508-99	Standard Test Method for Carbon Black, Pelleted-Fines and Attrition	No.100
	D1511-00	Standard Test Method for Carbon Black-Pellet Size Distribution	No.10 - No.120
	D1514-00	Standard Test Method for Carbon Black-Sieve Residue	No.30 - No.325
Cement	C184-94	Standard Test Method for Fineness of Hydraulic Cement by the 150-µm (No.100)	No.100 & No.200
		and 75-µm (No.200) Sieves	
	C430-96	Standard Test Method for Fineness of Hydraulic Cement by the 45-µm (No.325)	No. 325
		Sieve	
	C796 06		No. 50 No. 200
	C786-96	Standard Test Method for Fineness of Hydraulic Cement and Raw Materials by	No.50 - No.200
		the 300-µm (No.50), 150-µm (No.100), and 75-µm (No.200) Sieves by Wet	
		Methods	
Ceramic	C325-81 (1997)	Standard Test Method for Wet Sieve Analysis of Ceramic Whiteware Clays	No.100 - No.325
	C371-89 (1999)	Standard Test Method for Wire-Cloth Sieve Analysis of Nonplastic Ceramic	No.70 - No.325
	AMERICA DE VICTOR A PROPERTIES A	Powders	
Coal	D197-87 (1994)	Standard Test Method for Sampling and Fineness Test of Pulverized Coal	No.16 - No.200
Jour	D4749-87 (1994)	Standard Test Method for Performing the Sieve Analysis of Coal and Designating	5 in No.400
	D4149-01 (1994)		3 111 140.400
S 00	D0044.00	Coal Size	
Coatings	D3214-96	Standard Test Methods for Coating Powders and Their Coatings Used for	
		Electrical Insulation	
	D3451-01	Standard Guide for Testing Coating Powders and Powder Coatings	
Coke	D293-93 (1999)	Standard Test Method for the Sieve Analysis of Coke	4 in No.200
	D5709-95 (2000)	Standard Test Method for Sieve Analysis of Petroleum Coke	3 in No.200
Enamel	C285-88 (1999)	Standard Test Methods for Sieve Analysis of Wet-Milled and Dry-Milled Porcelain	No.40 - No.325
Litation	0200 00 (1000)	Enamel	110.40 110.020
Class	0420.04		N= 0 N= 200
Glass	C429-01	Method for Sieve Analysis of Raw Materials for Glass Manufacture	No.8 - No.200
	D1214-89 (1994)	Test for Sieve Analysis of Glass Spheres	
Magnesium	D2772-90 (1997)	Standard Test Method for Sieve Analysis of Electrical Grade Magnesium Oxide	
Metal Bearing ores	E276-98	Standard Test Method for Particle Size or Screen Analysis at No.4 (4.75-mm)	No.4 - No.200
		Sieve and Finer for Metal-Bearing Ores and Related Materials	
Metal Powders	B214-99	Test for Sieve Analysis of Metal Powders	No.80 - No.325
Vlineral	D451-91 (1996)	Standard Test Method for Sieve Analysis of Granular Mineral Surfacing for	No.6 - No.100
A Commence of	2 10 1 0 1 (1000)	Asphalt Roofing Products	110.0 110.100
	D452-91 (1997)	Standard Test Method for Sieve Analysis of Surfacing for Asphalt Products	No.12 - No.200
			140.12 - 140.200
	D546-99	Standard Test Method for Sieve Analysis of Mineral Filler for Bituminous Paving	
		Mixtures	
Perlite	C549-81 (1995)	Standard Specification for Perlite Loose Fill Insulation	
Pigments and paint	D185-84 (1999)	Standard Test Methods for Coarse Particles in Pigments, Pastes and Paints	No.325
	D480-88 (1999)	Standard Test Methods for Sampling and Testing of Flaked Aluminum Powders	No.100 - No.325
		and Pastes	
Plastic	D1921-01	Standard Test Method for Particle Size (Sieve Analysis) of Plastic Materials	down to No.400
		선생님은 이번 사용하는 전 경험에는 현실 사용하다면 하다 아니는 이렇게 하다 아니는 이렇게 하다 아니는 이렇게 하다면 보고 있다. 그렇게 하는 사람들이 되었다면 하는 사람들이 아니는	
Porcelain	C285-88 (1999)	Standard Test Methods for Sieve Analysis of Wet-Milled and Dry-Milled Porcelain	No.40 - No.325
		Enamel	
Refractories	C92-95 (1999)	Tests for Sieve Analysis and Water Content of Refractory Materials	3 in No.200
Resins	D2187-94 (1998)	Standard Test Methods for Physical and Chemical Properties of Particulate Ion-	No.8 - No.100
		Exchange Resins	
Rubber additives	D5461-93 (1998)	Standard Test Method for Rubber Additives-Wet Sieve Analysis of Powdered	
	50.0.00 (1000)	Rubber Chemicals	
Soan	D502 90 /4005\		No 12 No 100
Soap Sodo och	D502-89 (1995)	Standard Test Method for Particle Size of Soaps and Other Detergents	No.12 - No.100
Soda ash	E359-00	Standard Test Methods for Analysis of Soda Ash (Sodium Carbonate)	NIS A NISSAN
Soil	D421-85 (1998)	Standard Practice for Dry Preparation of Soil Samples for Particle-Size Analysis	No.4 - No.40
		and Determination of Soil Constants	
	D422-63 (1998)	Standard Test Method for Particle-Size Analysis of Soils	3 in No.200
	D1140-00	Standard Test Methods for Amount of Material in Soils Finer Than the No.200 (75-	No.40 - No.200
		µm) Sieve	
	D2217-85 (1998)	Standard Practice for Wet Preparation of Soil Samples for Particle-Size Analysis	No.10 - No.40
	DZZ11-03 (1880)		140.10 - 140.40
√ermiculite	0540.00 (4000)	and Determination of Soil Constants	97:- N 400
	C516-80 (1996)	Standard Specification for Vermiculite Loose Fill Insulation	3⁄4 in No.100



# Meinzer II Frequently Asked Questions

For specific sieving procedures, please refer to <u>Test Sieving: Principles and Procedures</u> located in the User's Manual.

1. What are the vibrations per minute for the Meinzer II?

A 60Hz machine will produce 3,600 vibrations per minute.

2. What sort of maintenance is required for the Meinzer II?

The Meinzer II just needs to be kept clean. No lubrication or resetting of parts is required. Any alteration or unauthorized maintenance will void the warranty.

3. Does Advantech calibrate/certify test sieves for the Meinzer II?

Yes. Test sieves can be certified using Advantech's <u>Centerline<sup>©</sup> Premium Sieve Certification</u>. For a <u>suggested re-certification schedule</u>, please contact our Customer Service Team at 262.786.1600 or <u>sales@advantechmfg.com</u>.

4. How many sieves can fit in on the Meinzer II?

Please refer to Chart 4A for details on the Meinzer II sieve capacity.

#### Chart 4A

#### Meinzer II Sieve Capacity

	Half Height (min/max)	Full Height (min/max)	Pan	Cover
8" (203.2mm)or 200mm	4/15	2/8	1	1



Figure 5A

5. My Meinzer II is making a lot of noise and the sieve stack is rattling in the machine. What is wrong?

The sieve stack may have been improperly secured. Failing to fully clamp as in **Figure 5A** or adequately tighten the straps as in **Figure 5B** will cause the sieve stack to rattle. For instructions on how to properly build and secure your test sieve stack, please refer to step 8 in <u>Performing a Sieve Analysis using the Meinzer II Testing Sieve Shaker</u> in the front of this manual.



#### 6. What is the warranty on the Meinzer II?

The Meinzer II carries a one year limited warranty against defective material and workmanship.

#### 7. What is an extended rim pan? Do I need this for my test?

An extended rim pan is manufactured with a skirt around the bottom so it can be received by a sieve below it. This will allow the user to run multiple samples in one stack. The extended rim pan can be inserted mid-stack to collect fines of sample one and the bottom pan will collect fines from sample two. See **Figure 7A** for an example.



Figure 7A

#### 8. The fuse has blown in the machine. How do I change it?

The fuse must be replaced with one of identical rating. To replace the fuse, perform the following:

- Disconnect the machine from the power supply.
- Unscrew the central cap of the fuse holder.
- Remove the holder and fuse together.
- Remove the blown fuse and place the new fuse in the metal spring in the central cap.
- Fit the cap and fuse back into the holder and screw in completely. **Do not over tighten.**

#### 9. Does Advantech have a repair facility nearby?

Advantech is pleased to offer telephone repair support for Meinzer II Testing Sieve Shakers. Contact a member of our Tech Support Team at 262.786.1600. Alternatively, machines may be sent in to our location in New Berlin, WI for extensive repair or refurbishing. Contact us for information on how to prepare your machine for receipt and service by our Repair Department.

## 10. My questions have still not been answered.

For further technical support, please contact our Tech Support Team at 262.786.1600 or at sales@advantechmfg.com. We'd be glad to assist.

## **Notes**