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Standard Guide for Development of Laser Diffraction Particle Size Analysis Methods for Powder Materials¹

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1. Scope

1.1 This guide sets out the general approach to the particle size distribution measurement of powders, suspensions, or slurries using an appropriate wet or dry methodology by the laser diffraction technique. It is recommended for use in measurements of broad particle size distributions.

1.2 The guide provides guidelines to the parameters that should be specified and a generalized guideline to reasonable and acceptable tolerances for points in the volume-based distribution curve such as x_{10} ($D_{v,10}$), x_{50} ($D_{v,50}$), x_{90} ($D_{v,90}$), and $D[4, 3]$ (volume moment mean). It is noted that ISO prefers the term x for particle size as opposed to other usage of d or D (implying diameter).

1.3 This guide provides guidance on the verification of instrument performance in conjunction with the internal quality control (QC) audit functions of the instrument owner. Results should be reported in the format indicated by Practice E1617 and ISO 13320.

1.4 *Units*—The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

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

1.6 *This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.*

¹ This guide is under the jurisdiction of ASTM Committee E29 on Particle and Spray Characterization and is the direct responsibility of Subcommittee E29.02 on Non-Sieving Methods.

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2. Referenced Documents

2.1 ASTM Standards:²

B821 Guide for Liquid Dispersion of Metal Powders and Related Compounds for Particle Size Analysis

E1617 Practice for Reporting Particle Size Characterization Data

E2490 Guide for Measurement of Particle Size Distribution of Nanomaterials in Suspension by Photon Correlation Spectroscopy (PCS)

2.2 ISO Standards:³

ISO 13320 Particle size analysis — Laser diffraction methods

ISO 14887 Sample preparation — Dispersing procedures for powders in liquids

ISO 14488/AMD 1:2019 Particulate materials — Sampling and sample splitting for the determination of particulate properties

3. Terminology

3.1 Definitions:

3.1.1 Some of the definitions in 3.2 will differ slightly from those used within other (non-particle sizing) standards. For further details, see the Terminology section of Guide E2490.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *differential pressure (ΔP)*, n —the pressure difference across a venturi that causes acceleration (and thus shear forces) leading to dispersion (and possibly attrition) in a dry laser diffraction measurement.

3.2.2 *laser diffraction*, n —light scattering technique primarily used for particle size distribution analysis, applicable to the approximate range 0.1 – 3000 μm , where multiple angles are

² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

³ Available from International Organization for Standardization (ISO), ISO Central Secretariat, Chemin de Blandonnet 8, CP 401, 1214 Vernier, Geneva, Switzerland, <https://www.iso.org>.

used to analyze the (laser light) scattering from an ensemble of particles in liquid or dry suspension. See ISO 13320.

3.2.3 *method development, n—in laser diffraction*, this refers to the actions involved in understanding how energy affects the apparent particle size distribution of the material in line with the measurement objectives (that is, bulk or dispersed size required).

3.2.4 *repeatability, n—in particle sizing techniques*, this usually refers to the precision of repeated consecutive measurements on the same group of particles and is normally expressed as a relative standard deviation (RSD) or coefficient of variation (C.V.).

3.2.4.1 *Discussion*—The repeatability value reflects the stability (instrumental, but mainly the sample) of the system over time. Changes in the sample could include dispersion and settling.

3.2.5 *reproducibility, n—in particle sizing*, this usually refers to second and further aliquots of the same bulk sample (and therefore is subject to the heterogeneity of the starting material and the sampling method employed).

3.2.5.1 *Discussion*—In a slurry system, it is often the largest error when repeated samples are taken. Other definitions of reproducibility also address the variability among single test results gathered from different laboratories when inter-laboratory testing is undertaken. It is to be noted that the same group of particles can never be measured in such a system of tests and therefore reproducibility values are typically considerably in excess of repeatability values.

3.2.6 *robustness, n*—a measure of the change of the required parameter with deliberate and systematic variations in any or all of the key parameters that influence it.

3.2.6.1 *Discussion*—For example, dispersion time (ultrasound time and power setting) almost invariably will affect the reported results. Variation in pH is likely to affect the degree of agglomeration as will the addition (deliberate or accidental) of any ionic species and so forth. Changes to the input optical constants may also have an effect on the end result.

3.2.7 *standard operating procedure (SOP), n*—the formulation of a fixed measurement protocol once method development has taken place.

4. Summary of Guide

4.1 This guide provides a general overview of the methodologies involved with the development of wet and dry dispersion techniques for measuring the particle size distribution of powdered systems. Specific details are provided to the approach taken when a powder is either measured directly with dry dispersion or wetted in a liquid for wet dispersion.

5. Significance and Use

5.1 The technique of laser diffraction for particle size distribution analysis is extensively used in industry and academia both for on-line control and laboratory needs. Guidance is obviously useful in this regard.

5.2 This guide can be used to develop methods of particle size analysis where well-established analysis procedures do not

already exist. See Guide B821 for similar guidance and useful procedures for wet dispersion of metal powders and related compounds.

6. Reagents

6.1 In general, no reagents specific to the laser diffraction technique are necessary. For dry measurement the dispersant ‘fluid’ is usually dry, particulate-matter-free air. However, in the wet methodology, dispersing, and stabilizing agents can be used for a specific test sample in order to preserve stability during the measurement. A suitable diluent is used to achieve a particle concentration appropriate for a laser diffraction measurement (typically 0.001 – 0.1 volume %). Particle size may undergo change on dilution, as the ionic environment within which the particles are dispersed, changes in nature or concentration. The apparent particle size may change in line with increased concentration (obscuration), especially with particles smaller than around 2 μm , due to multiple scattering.

7. Procedures

7.1 *Overview and Introduction*—The four basic questions to answer prior to performing a particle size distribution measurement are:

7.1.1 What is the acceptable quality level (AQL) of the organization? This sets the desired or required precision for a specification and has an impact on the minimum mass to meet this requirement.

7.1.2 What is the top end point of the distribution that requires specifying or control? This has impact on the minimum mass required.

7.1.3 Is a bulk size (“as is”/with agglomerates) required or is a dispersed (primary) size desired? In other words, “What is the purpose of taking the measurement?” The answers to these questions direct the routes to measurement and the level of energy input required to keep the material in bulk phase or to disperse it to primary particle size.

7.1.4 What is the polydispersity (width/spread) of the particle size distribution and the density of the material? Again, the answer to this question has direct impact on the minimum mass required to meet any proscribed specification or material variation.

7.1.5 To estimate the minimum standard error (fundamental sampling error, FSE) in an experiment due solely to the heterogeneity of the particle size distribution, the following fifth question requires answering:

7.1.5.1 What is the mass of sample that is utilized in the particle size experiment?

7.1.5.2 Note that any formulated specification needs to be ‘just good enough’ and ‘fit for purpose’. The link between the particle size distribution and product performance (product performance indicators) or critical quality attributes to be well understood before any specification is developed.

7.2 The purpose of method development is to understand how the input of energy affects the apparent particle size distribution. The effect of changes in pH and chemistry can also be investigated at this stage. When the effect of these parameters is understood, a standard operating procedure (SOP) can be formulated in line with the objectives of the