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Nanotechnologies – Compilation and description of sample preparation and dosing methods for engineered and manufactured nanomaterials

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Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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The committee responsible for this document is ISO/TC 229, *Nanotechnologies*.

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Introduction

This document provides guidance regarding the preparation of nanomaterials for toxicological, including eco-toxicological, testing. The goal of this document is to assist health and environmental scientists and scientists and experts from other disciplines to understand, plan, choose and address issues relevant to nanomaterials before and during conducting toxicological tests. These issues include the effects of the properties of the material on preparation methods and of the media into which the samples of nanomaterials will be added. Failure to consider these effects might lead to erroneous conclusions regarding the relationship between the nature of the nanomaterial and observed toxicological responses. In particular, the composition and other characteristics of test media can affect the dose to which an organism that is the subject of a test will be exposed. Information on preparation of the test material is necessary prior to any biological or ecological evaluation. Information such as this is consistent with other ISO documents. For example, ISO 10993-18^[1] specifically addresses the evaluation of the chemical characterization of materials used in medical devices, ISO 14971^[2] specifies that a toxicological risk analysis should take into account the chemical nature of the materials, ISO/TR 13014^[3] addresses issues pertaining to the materials themselves and ISO/TS 19337^[55] points out the need to clarify whether observed toxic effects come from tested nano-objects themselves or from other uncontrolled sources. Some examples are provided of methods that establish test conditions that are relatable to environmentally relevant conditions.

This document uses a number of technical terms which have been defined earlier in other documents. Some of these terms have been defined in multiple documents, in different areas of science and technology, providing potentially or seemingly conflicting definitions. This document does not provide new, authoritative definitions for the terms used herein. Instead, this clause provides short descriptions for the terms used. Where possible, reference is made to existing documents.

Nanotechnologies — Compilation and description of sample preparation and dosing methods for engineered and manufactured nanomaterials

1 Scope

This document provides guidance regarding the preparation of nanomaterials for eco- and bio-toxicological testing. It provides guidance regarding factors pertaining to sample preparation and dose determination that might be useful in toxicological, including ecotoxicological, testing of engineered and manufactured nanoscale materials.

The descriptions of sample preparation method factors for both *in vitro* and *in vivo* toxicological testing of engineered and manufactured nanoscale materials include considerations about physico-chemical properties, media, methods for transformation and accumulation studies, health effects and dosimetry. The document is not intended to be a literature review nor a thorough assessment of the quality of the methods or data generated. The document is intended to complement other international efforts.

The focus of this document is on factors that might lead to results that are not relevant to safety evaluations. When featured, referenced methods are considered for their general interest and potential applicability. It is likely that most of the described methods are not generally applicable to all nanomaterials but they do demonstrate important factors and limitations that are common for a variety of nanomaterials.

2 Normative references

There are no normative references in this document.

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- IEC Electropedia: available at <http://www.electropedia.org/>
- ISO Online browsing platform: available at <http://www.iso.org/obp>

3.1 particle

minute piece of matter with defined physical boundaries

Note 1 to entry: A physical boundary can also be described as an interface.

Note 2 to entry: A particle can move as a unit.

Note 3 to entry: This general particle definition applies to nano-objects.

[SOURCE: ISO/TS 80004-2:2015, 3.1]

3.2 structure

arrangement defined by four different aspects (crystallinity, crystal structure, molecular structure and microstructure)

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3.2.1

crystallinity

presence or absence of crystalline structure in the arrangement of the atoms of which a material consists

3.2.2

crystal structure

lattice structure in which atoms of an individual crystal are arranged, using lattice parameters and lattice type, such as face-centred cubic, hexagonal close-packed, body-centred, cubic, etc.

3.2.3

molecular structure

arrangement of atoms of an individual molecule

3.2.4

microstructure

arrangement of individual crystals or amorphous phases in a polycrystalline or multiphase material

3.3

measurand

quantity intended to be measured or a quantity that is being determined by measurement

Note 1 to entry: The specification of a measurand requires knowledge of the kind of quantity, description of the state of the phenomenon, body, or substance carrying the quantity, including any relevant component, and the chemical entities involved. The measurement, including the measuring system and the conditions under which the measurement is carried out, might change the phenomenon, body, or substance so that the quantity being measured may differ from the measurand as defined.

[SOURCE: ISO/IEC Guide 99, 2007, 2.3 — modified]

3.4

nanomaterial

NM

material with any external dimension in the *nanoscale* or having internal structure or surface structure in the nanoscale

Note 1 to entry: This generic term is inclusive of *nano-object* and *nanostructured material*.

Note 2 to entry: See also engineered nanomaterial, manufactured nanomaterial, incidental nanomaterial.

[SOURCE: ISO/TS 80004-1:2015, 2.4]

3.5

nano-object

discrete piece of material with one, two or three external dimensions in the *nanoscale*

Note 1 to entry: The second and third external dimensions are orthogonal to the first dimension and to each other.

[SOURCE: ISO/TS 80004-2:2015, 2.2]

3.6

nanoparticle

NP

nano-object with all external dimensions in the *nanoscale* where the lengths of the longest and the shortest axes of the nano-object do not differ significantly

Note 1 to entry: If the dimensions differ significantly (typically by more than three times), terms, such as *nanofibre* or *nanoplate*, may be preferred to the term nanoparticle.

[SOURCE: ISO/TS 80004-2:2015, 4.4]

3.7

nanoscale

length range approximately from 1 nm to 100 nm

Note 1 to entry: Properties that are not extrapolations from a larger size are predominantly exhibited in this length range.

[SOURCE: ISO/TS 80004-2:2015, 2.1]

4 Abbreviated terms

BET Brunauer–Emmett–Teller isotherm

CNT carbon nanotube

DLS dynamic light scattering

ICP-MS inductively coupled plasma mass spectrometry

NOAA nano-objects, and their aggregates and agglomerates greater than 100 nm

NOM natural organic material

TEM transmission electron microscopy

5 Background

5.1 Discussion of the importance of sample preparation and dosing

Nanomaterials are diverse, being based on endless combinations of composition, particle size and distribution, surface chemistry and many other key properties. With this diversity, nanomaterials cannot be treated as a single class of substances. Just as in other areas of toxicology, the assessment of biological effects should consider how samples and doses are prepared and dosimetry is assured so that the observed effects are meaningful and test results can be used in a realistic way such as in safety assessments.

Screening tests are used for rapid evaluations and are typically conducted using cell culture or other *in vitro* techniques due to fast response time, cost, infrastructure and time constraints, factors that limit most whole animal studies. The purpose of a screening test is to provide an indicator of potential adverse outcomes and effects on human health or the environment. Although there are many definitions available for the term screening test, for the purposes of this document, a screening test can be generally defined as a relatively simple, inexpensive test that can be administered easily and provides rapid results. The screening tests should reflect the compromise between simplicity, rapidity and low-cost while still providing results that have meaning to safety-relevant situations, the way that samples are prepared and the doses administered will ideally be relatable to realistic situations. Therefore, the points considered in this document also apply to screening tests and should also be taken into account in tiered testing to ensure consistent conditions with each step of the tier.