
**Radiological protection — Monitoring
and internal dosimetry for staff
members exposed to medical
radionuclides as unsealed sources**

*Radioprotection — Surveillance et dosimétrie interne des travailleurs
exposés lors des utilisations médicales des radioéléments en sources
non scellées*



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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).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

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For an explanation on the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the WTO principles in the Technical Barriers to Trade (TBT) see the following URL: [Foreword - Supplementary information](#)

The committee responsible for this document is ISO/TC 85, *Nuclear energy, nuclear technologies, and radiological protection*, Subcommittee SC 2, *Radiological protection*.

Introduction

In the course of employment, individuals might work with radioactive materials that, under certain circumstances, could be taken into the body. Protecting workers against risks of incorporated radionuclides requires the monitoring of potential intakes and/or the quantification of actual intakes and exposures. The doses resulting from internal radiation exposure arising from contamination by radioactive substances cannot be measured directly. The selection of measures and programmes for this purpose requires decisions concerning methods, techniques, frequencies, etc. for activity measurements and dose assessment. The criteria permitting the evaluation of the necessity of such a monitoring programme or for the selection of methods and frequencies of monitoring usually depend upon the legislation, the purpose of the radiation protection programme, the probabilities of potential radionuclide intakes, and the characteristics of the materials handled.

For these reasons, ISO standards establishing requirements for monitoring programmes (ISO 20553), laboratory requirements (ISO 28218), and dose assessment (ISO 27048) have been developed. These can be applied in a straightforward manner to many workplaces where internal contamination may occur. In order to apply these standards to staff involved in diagnostic or therapeutic uses of radionuclides in medicine, the short effective half-life of radionuclides commonly used for these purposes and the distance between nuclear medicine department and *in vivo* counting facilities or radio-analytical laboratories shall be taken into account. Consequently, guidance on the application of the three International Standards cited above to nuclear medicine staff was requested by a number of countries.

This International Standard establishes criteria to determine whether intake monitoring is required for staff exposed to medical radionuclides as unsealed sources. It also establishes requirements on the design of such monitoring programmes, associated dose assessments, and laboratory requirements. Recommendations of international expert bodies and international experience with the practical application of these recommendations in radiological protection programmes have been considered in the development of this International Standard. Its application facilitates the exchange of information between authorities, supervisory institutions, and employers. This International Standard is not a substitute for legal requirements.

Radiological protection — Monitoring and internal dosimetry for staff members exposed to medical radionuclides as unsealed sources

1 Scope

This International Standard specifies the minimum requirements for the design of professional programmes to monitor workers exposed to the risk of internal contamination via inhalation by the use of radionuclides as unsealed sources in nuclear medicine imaging and therapy departments. It establishes principles for the development of compatible goals and requirements for monitoring programmes and, when adequate, dose assessment. It presents procedures and assumptions for the risk analysis, for the monitoring programmes, and for the standardized interpretation of monitoring data.

This International Standard addresses the following items:

- a) purposes of monitoring and monitoring programmes;
- b) description of the different categories of monitoring programmes;
- c) quantitative criteria for conducting monitoring programmes;
- d) suitable methods for monitoring and criteria for their selection;
- e) information that has to be collected for the design of a monitoring programme;
- f) general requirements for monitoring programmes (e.g. detection limits, tolerated uncertainties);
- g) frequencies of measurements;
- h) procedures for dose assessment based on reference levels for routine and special monitoring programmes;
- i) assumptions for the selection of dose-critical parameter values;
- j) criteria for determining the significance of individual monitoring results;
- k) interpretation of workplace monitoring results;
- l) uncertainties arising from dose assessments and interpretation of bioassays data;
- m) reporting/documentation;
- n) quality assurance.

This International Standard does not address the following:

- monitoring and internal dosimetry for the workers exposed to laboratory use of radionuclides such as radioimmunoassay techniques;
- monitoring and internal dosimetry for the workers involved in the operation, maintenance, and servicing of PET cyclotrons;
- detailed descriptions of measuring methods and techniques;
- dosimetry for litigation cases;
- modelling for the improvement of internal dosimetry;

- the potential influence of medical treatment of the internal contamination;
- the investigation of the causes or implications of an exposure;
- dosimetry for ingestion exposures and for contaminated wounds.

2 Normative references

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 20553, *Radiation protection — Monitoring of workers occupationally exposed to a risk of internal contamination with radioactive material*

ISO 27048:2011, *Radiation protection — Dose assessment for the monitoring of workers for internal radiation exposure*

ISO 28218, *Radiation protection — Performance criteria for radiobioassay*

ISO/IEC Guide 99, *International vocabulary of metrology — Basic and general concepts and associated terms (VIM)*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO/IEC Guide 99, ISO 20553, ISO 28218 and ISO 27048 and the following apply.

3.1 absorption

movement of material to blood regardless of mechanism, generally applied to dissociation of particles and uptake into blood of soluble substances and material dissociated from particles

3.2 absorption type F

as defined by ICRP, deposited materials that have high (fast) rates of *absorption* (3.1) into body fluids from the respiratory tract

3.3 absorption type M

as defined by ICRP, deposited materials that have intermediate (moderate) rates of *absorption* (3.1) into body fluids from the respiratory tract

3.4 activity

number of spontaneous nuclear transformations per unit time

Note 1 to entry: The activity is stated in becquerel (Bq), i.e. the number of transformations per second.

3.5 activity median aerodynamic diameter AMAD

value of aerodynamic diameter such that 50 % of the airborne *activity* (3.4) in a specified aerosol is associated with particles smaller than the AMAD, and 50 % of the activity is associated with particles larger than the AMAD

Note 1 to entry: The aerodynamic diameter of an airborne particle is the diameter that a sphere of unit density would need to have in order to have the same terminal velocity when settling in air as the particle of interest.