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SINGAPORE STANDARD Medical laboratories — Requirements for quality and competence



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Contents

Page

Nation	al Foreward	6			
Forewa	ard	7			
Introduction					
1	Scope				
2	Normative references				
3	Terms and definitions				
4	nagement requirements				
•	A 1 Organization and management responsibility	15			
	4.2 Quality management system	18			
	4.2 Quanty management system	20			
	AA Service agreements	21			
	4.5 Evamination by referral laboratories	22			
	1.6 External convices and cumplies	22			
	4.7 Advisory services	23 72			
	4.7 Advisory services	23 22			
	4.0 Identification and control of nonconformition	23			
	4.7 Identification and control of noncomor mittes	24			
	4.10 Corrective action	24			
	4.11 Fleventive action	25			
	4.12 Control of records	23			
	4.15 Control of records	23			
	4.14 Evaluation and addits	27			
_	4.15 Management review	29 20			
5	Technical requirements	30			
	5.1 Personnel	30			
	5.2 Accommodation and environmental conditions	33			
	5.3 Laboratory equipment, reagents, and consumables	35			
	5.4 Pre-examination processes	39			
	5.5 Examination processes	1 3			
	5.6 Ensuring quality of examination results	16			
	5.7 Post-examination processes	49 49			
	5.8 Reporting of results	49			
	5.9 Release of results	51			
	5.10 Laboratory information management	52			
Annex	Annex A (informative) Correlation with ISO 9001:2008 and ISO/IEC 17025:2005				
Annex	Annex B (informative) Comparison of ISO 15189:2007 to ISO 15189:2012				
Bibliography					

National Foreword

This Singapore Standard was prepared by the National Mirror Technical Committee for ISO/TC 212 under the direction of the Biomedical and Health Standards Committee. This Singapore Standard is an identical adoption of International Standard ISO 15189 : 2012 "Medical laboratories — Requirements for quality and competence" published by the International Organization for Standardization.

Attention is drawn to the following:

- 1. Where appropriate, the words 'International Standard' shall be read as 'Singapore Standard'.
- 2. The references to International Standards shall be replaced by the following Singapore Standards:

International Standard	Corresponding Singapore Standard	
ISO 9000	SS ISO 9000	
ISO 9001	SS ISO 9001	
ISO 15189	SS ISO 15189	
ISO 17000	SS ISO 17000	
ISO 17025	SS ISO 17025	

Attention is drawn to the possibility that some of the elements of this Singapore Standard may be the subject of patent rights. Enterprise Singapore shall not be held responsible for identifying any or all of such patent rights.

NOTE

- 1. Singapore Standards (SSs) and Technical References (TRs) are reviewed periodically to keep abreast of technical changes, technological developments and industry practices. The changes are documented through the issue of either amendments or revisions.
- 2. An SS or TR is voluntary in nature except when it is made mandatory by a regulatory authority. It can also be cited in contracts making its application a business necessity. Users are advised to assess and determine whether the SS or TR is suitable for their intended use or purpose. If required, they should refer to the relevant professionals or experts for advice on the use of the document. Enterprise Singapore shall not be liable for any damages whether directly or indirectly suffered by anyone or any organisation as a result of the use of any SS or TR.
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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.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

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ISO 15189 was prepared by Technical Committee ISO/TC 212, *Clinical laboratory testing and in vitro diagnostic test systems*.

This third edition cancels and replaces the second edition (ISO 15189:2007), which has been technically revised.

A correlation between the second and third editions of this International Standard is provided as Annex B. The third edition continues the alignment established in ISO/IEC 17025:2005.

This corrected version of ISO 15189:2012 includes various editorial corrections.

Introduction

This International Standard, based upon ISO/IEC 17025 and ISO 9001, specifies requirements for competence and quality that are particular to medical laboratories¹. It is acknowledged that a country could have its own specific regulations or requirements applicable to some or all its professional personnel and their activities and responsibilities in this domain.

Medical laboratory services are essential to patient care and therefore have to be available to meet the needs of all patients and the clinical personnel responsible for the care of those patients. Such services include arrangements for examination requests, patient preparation, patient identification, collection of samples, transportation, storage, processing and examination of clinical samples, together with subsequent interpretation, reporting and advice, in addition to the considerations of safety and ethics in medical laboratory work.

Whenever allowed by national, regional or local regulations and requirements, it is desirable that medical laboratory services include the examination of patients in consultation cases, and that those services actively participate in the prevention of disease in addition to diagnosis and patient management. Each laboratory should also provide suitable educational and scientific opportunities for professional staff working with it.

While this International Standard is intended for use throughout the currently recognized disciplines of medical laboratory services, those working in other services and disciplines such as clinical physiology, medical imaging and medical physics could also find it useful and appropriate. In addition, bodies engaged in the recognition of the competence of medical laboratories will be able to use this International Standard as the basis for their activities. If a laboratory seeks accreditation, it should select an accrediting body which operates in accordance with ISO/IEC 17011 and which takes into account the particular requirements of medical laboratories.

This International Standard is not intended to be used for the purposes of certification, however a medical laboratory's fulfilment of the requirements of this International Standard means the laboratory meets both the technical competence requirements and the management system requirements that are necessary for it to consistently deliver technically valid results. The management system requirements in Clause 4 are written in a language relevant to a medical laboratory's operations and meet the principles of ISO 9001:2008, *Quality management systems* — *Requirements*, and are aligned with its pertinent requirements (Joint IAF-ILAC-ISO Communiqué issued in 2009).

The correlation between the clauses and subclauses of this third edition of ISO 15189 and those of ISO 9001:2008 and of ISO/IEC 17025:2005 is detailed in Annex A of this International Standard.

Environmental issues associated with medical laboratory activity are generally addressed throughout this International Standard, with specific references in 5.2.2, 5.2.6, 5.3, 5.4, 5.5.1.4 and 5.7.

¹ In other languages, these laboratories can be designated by the equivalent of the English term "clinical laboratories."

Medical laboratories — Requirements for quality and competence

1 Scope

This International Standard specifies requirements for quality and competence in medical laboratories.

This International Standard can be used by medical laboratories in developing their quality management systems and assessing their own competence. It can also be used for confirming or recognizing the competence of medical laboratories by laboratory customers, regulating authorities and accreditation bodies.

NOTE International, national or regional regulations or requirements may also apply to specific topics covered in this International Standard.

2 Normative references

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

ISO/IEC 17000, Conformity assessment — Vocabulary and general principles

ISO/IEC 17025:2005, General requirements for the competence of testing and calibration laboratories

ISO/IEC Guide 2, Standardization and related activities — General vocabulary

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