

OHUTUSNÕUDED ELEKTRILISTELE MÕÕTMIS-,  
JUHTIMIS- JA LABORATOORIUMISEADMETELE. OSA  
2-101: OHUTUSNÕUDED IN VITRO DIAGNOSTILISTELE  
(IVD) MEDITSIINISEADMETELE

Safety requirements for electrical equipment for  
measurement, control, and laboratory use - Part 2-101:  
Safety requirements for in vitro diagnostic (IVD)  
medical equipment

## EESTI STANDARDI EESSÕNA

## NATIONAL FOREWORD

See Eesti standard EVS-EN IEC 61010-2-101:2022 sisaldab Euroopa standardi EN IEC 61010-2-101:2022 ingliskeelset teksti.	This Estonian standard EVS-EN IEC 61010-2-101:2022 consists of the English text of the European standard EN IEC 61010-2-101:2022.
Standard on jõustunud sellekohase teate avaldamisega EVS Teatajas.	This standard has been endorsed with a notification published in the official bulletin of the Estonian Centre for Standardisation and Accreditation.
Euroopa standardimisorganisatsioonid on teinud Euroopa standardi rahvuslikele liikmetele kättesaadavaks 11.11.2022.	Date of Availability of the European standard is 11.11.2022.
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English Version

Safety requirements for electrical equipment for measurement,  
control, and laboratory use - Part 2-101: Safety requirements for  
in vitro diagnostic (IVD) medical equipment  
(IEC 61010-2-101:2018)

Exigences de sécurité pour appareils électriques de  
mesurage, de régulation et de laboratoire - Partie 2-101:  
Exigences particulières pour le matériel médical de  
diagnostic in vitro (DIV)  
(IEC 61010-2-101:2018)

Sicherheitsbestimmungen für elektrische Mess-, Steuer-,  
Regel- und Laborgeräte - Teil 2-101: Besondere  
Anforderungen an In-vitro-Diagnostik (IVD) Medizingeräte  
(IEC 61010-2-101:2018)

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Comité Européen de Normalisation Electrotechnique  
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CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels

## European foreword

The text of document 66/644/CDV, future edition 3 of IEC 61010-2-101, prepared by IEC/TC 66 "Safety of measuring, control and laboratory equipment" was submitted to the IEC-CENELEC parallel vote and approved by CENELEC as EN IEC 61010-2-101:2022.

The following dates are fixed:

- latest date by which the document has to be implemented at national (dop) 2023-09-26 level by publication of an identical national standard or by endorsement
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The text of the International Standard IEC 61010-2-101:2018 was approved by CENELEC as a European Standard without any modification.

In the official version, for Bibliography, the following notes have to be added for the standards indicated:

IEC 62061	NOTE	Harmonized as EN IEC 62061
IEC 62366-1	NOTE	Harmonized as EN 62366-1
ISO 15223-1	NOTE	Harmonized as EN ISO 15223-1

# INTERNATIONAL STANDARD

## NORME INTERNATIONALE

GROUP SAFETY PUBLICATION  
PUBLICATION GROUÉE DE SÉCURITÉ

**Safety requirements for electrical equipment for measurement, control and laboratory use –  
Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment**

**Exigences de sécurité pour appareils électriques de mesurage, de régulation et de laboratoire –  
Partie 2-101: Exigences particulières pour le matériel médical de diagnostic in vitro (DIV)**



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## INTERNATIONAL ELECTROTECHNICAL COMMISSION

**SAFETY REQUIREMENTS FOR ELECTRICAL EQUIPMENT FOR  
MEASUREMENT, CONTROL AND LABORATORY USE –****Part 2-101: Particular requirements for  
in vitro diagnostic (IVD) medical equipment**

## FOREWORD

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International Standard IEC 61010-2-101 has been prepared by IEC technical committee 66: Safety of measuring, control and laboratory equipment.

It has the status of a group safety publication, as specified in IEC Guide 104.

This document has been prepared in close collaboration with Working Group CENELEC BTTF 88.1.

This third edition cancels and replaces the second edition published in 2015. This edition constitutes a technical revision.

This edition includes the following significant technical changes with respect to the previous edition:

- a) adaptation of changes introduced by Amendment 1 of IEC 61010-1;
- b) added tolerance for stability of AC voltage test equipment to Clause 6.

The text of this International Standard is based on the following documents:

CDV	Report on voting
66/644/CDV	66/669/RVC

Full information on the voting for the approval of this International Standard can be found in the report on voting indicated in the above table.

This document has been drafted in accordance with the ISO/IEC Directives, Part 2.

A list of all parts of the IEC 61010 series, under the general title: *Safety requirements for electrical equipment for measurement, control, and laboratory use*, may be found on the IEC website.

This Part 2-101 is intended to be used in conjunction with IEC 61010-1. It was established on the basis of the third edition (2010) and its Amendment 1 (2016).

This Part 2-101 supplements or modifies the corresponding clauses in IEC 61010-1 so as to convert that publication into the IEC standard: *Particular requirements for in vitro diagnostic (IVD) medical equipment*.

Where a particular subclause of Part 1 is not mentioned in this Part 2, that subclause applies as far as is reasonable. Where this part states “addition”, “modification”, “replacement”, or “deletion” the relevant requirement, test specification or note in Part 1 should be adapted accordingly.

In this standard:

- 1) the following print types are used:
  - requirements: in roman type;
  - NOTES: in smaller roman type;
  - *conformity and test: in italic type*;
  - terms used throughout this standard which have been defined in clause 3: SMALL ROMAN CAPITALS;
- 2) subclauses, figures, tables and notes which are additional to those in part 1 are numbered starting from 101. Additional annexes are lettered starting from AA and additional list items are lettered from aa).

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- reconfirmed,
- withdrawn,
- replaced by a revised edition, or
- amended.

## **SAFETY REQUIREMENTS FOR ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL AND LABORATORY USE –**

### **Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment**

#### **1 Scope and object**

This clause of Part 1 is applicable except as follows:

##### **1.1.1 Equipment included in scope**

*Replacement:*

*Replace the text, except the first paragraph, with the following new text:*

This part of IEC 61010 applies to equipment intended for in vitro diagnostic (IVD) medical purposes, including self-test IVD medical purposes.

IVD medical equipment, whether used alone or in combination, is intended by the manufacturer to be used in vitro for the examination of specimens, including blood and tissue samples, derived from the human body, solely or principally for the purpose of providing information concerning one or more of the following:

- a physiological or pathological state; or
- a congenital abnormality;
- the determination of safety and compatibility with potential recipients;
- the monitoring of therapeutic measures.

Self-test IVD medical equipment is intended by the manufacturer for use by lay persons in a home environment.

NOTE If all or part of the equipment falls within the scope of one or more other Part 2 standards of the IEC 61010 series as well as within the scope of this document, consideration is given to those other Part 2 standards.

##### **1.1.2 Equipment excluded from scope**

*Addition:*

*Add the following new item:*

- aa) equipment within the scope of IEC 61010-2-081 unless it is specifically intended by the manufacturer to be used for in vitro diagnostic examination.

#### **1.2 Object**

##### **1.2.1 Aspects included in scope**

*Addition:*

*Add the following two new items:*

- aa) biohazards;
- bb) hazardous chemical substances.

### 1.2.2 Aspects excluded from scope

*Addition:*

*Add the following new item and note:*

- aa) the handling or manipulation outside the equipment of material under analysis.

NOTE Requirements covering these subjects are the responsibility of committees preparing the relevant standards.

## 2 Normative references

This clause of Part 1 is applicable except as follows:

*Addition:*

*Add the following new references to the list:*

ISO 14971, *Medical devices – Application of risk management to medical devices*

ISO 18113-5, *In vitro diagnostic medical devices – Information supplied by the manufacturer (labelling) – Part 5: In vitro diagnostic instruments for self-testing*

## 3 Terms and definitions

This clause of Part 1 is applicable except as follows:

### 3.1 Equipment and states of equipment

*Addition:*

*Add the following new terms:*

#### 3.1.101

##### **SAMPLE ZONE**

area where OPERATOR access is typically unintended

Note 1 to entry: The inside of this zone presents mechanical HAZARDS and a more likely probability of biohazardous human skin puncture.

#### 3.1.102

##### **LOADING ZONE**

area of automated equipment where an OPERATOR handles sample or reagent material

#### 3.5.12 RESPONSIBLE BODY

*Addition:*

*Add the following new note:*

Note 1 to entry: This is not the European Union's responsible authority.

## 4 Tests

This clause of Part 1 is applicable.