

INTERNATIONAL STANDARD

IEC 60601-2-37

Edition 1.1

2004-10

Edition 1:2001 consolidated with amendment 1:2004

Medical electrical equipment –

Part 2-37: Particular requirements for the safety of ultrasonic medical diagnostic and monitoring equipment

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CONTENTS

FOREWORD.....	4
INTRODUCTION.....	6

SECTION ONE: GENERAL

1 Scope and object.....	7
2 Terminology and definitions.....	8
3 General requirements.....	18
6 Identification, marking and documents.....	18

SECTION TWO: ENVIRONMENTAL CONDITIONS

SECTION THREE: PROTECTION AGAINST ELECTRIC SHOCK HAZARDS

19 Continuous LEAKAGE CURRENTS and PATIENT AUXILIARY CURRENTS.....	21
20 Dielectric strength.....	22

SECTION FOUR: PROTECTION AGAINST MECHANICAL HAZARDS

SECTION FIVE: PROTECTION AGAINST HAZARDS FROM UNWANTED OR EXCESSIVE RADIATION

*35 Acoustical energy (including ultrasonic).....	22
*36 Electromagnetic compatibility.....	22

SECTION SIX: PROTECTION AGAINST HAZARDS OF IGNITION OF FLAMMABLE ANESTHETIC MIXTURES

SECTION SEVEN: PROTECTION AGAINST EXCESSIVE TEMPERATURES AND OTHER SAFETY HAZARDS

42 Excessive temperatures.....	25
44 Overflow, spillage, leakage, humidity, ingress of liquids, cleaning, sterilization, and disinfection.....	28

SECTION EIGHT: ACCURACY OF OPERATING DATA AND PROTECTION AGAINST HAZARDOUS OUTPUT

50 Accuracy of operating data.....	28
51 Protection against hazardous output.....	29

SECTION NINE: ABNORMAL OPERATION AND FAULT CONDITIONS; ENVIRONMENTAL TESTS

SECTION TEN: CONSTRUCTIONAL REQUIREMENTS

Annex AA (normative) Terminology – Index of defined terms	31
Annex BB (informative) Guidance and rationale for particular subclauses	33
Annex CC (informative) Guidance in classification according to CISPR 11.....	37
Annex DD (normative) Test methods for determining the MECHANICAL INDEX and the THERMAL INDEX.....	38
Annex EE (informative) Relationships with other standards	43
Annex FF (informative) Guidance notes for measurement of OUTPUT POWER in SCANNING MODE	44
Annex GG (informative) Rationale and derivation of index models	48
Annex HH (informative) Guidance on the interpretation of <i>TI</i> and <i>MI</i> to be used to inform the OPERATOR.....	62
Annex II (informative) Example set-up to measure surface temperature for externally applied transducers	65
Bibliography.....	68
Figure FF.1 – Suggested 1 cm-wide aperture mask	46
Figure FF.2 – Suggested orientation of probe, mask slit, and RFB target.....	46
Figure FF.3 – Suggested orientation of probe, mask slit, and 1 cm RFB target	46
Figure GG.2a – Focused transducer with a large aperture	60
Figure GG.2b – Focused transducer with smaller aperture ($\geq 1 \text{ cm}^2$)	60
Figure GG.2c – Focused transducer with a weak focus ($A_{\text{eq}} > 1 \text{ cm}^2$)	61
Figure GG.2d – Weakly focused transducer.....	61
Figure II.1 – Set-up of an example test object to measure the surface temperature of externally applied transducers	67
Table 101 – Acoustic output reporting table	21
Table 102 – Overview of the tests noted under 42.3.....	27
Table DD.1 – Summary of combination formulae for each of the THERMAL INDEX categories	42
Table DD.2 – Summary of the acoustic quantities required for the determination of the indices	42
Table GG.1 – THERMAL INDEX categories and models	50
Table GG.2 – THERMAL INDEX formulae.....	54
Table HH.1 – Relative importance of maintaining low exposure indices in various scanning situations	63
Table II.1 – Acoustic and thermal properties of tissues and materials	65
Table II.2 – Weight % pure components.....	66

INTERNATIONAL ELECTROTECHNICAL COMMISSION

MEDICAL ELECTRICAL EQUIPMENT –

Part 2-37: Particular requirements for the safety of ultrasonic medical diagnostic and monitoring equipment

FOREWORD

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International Standard IEC 60601-2-37 has been prepared by subcommittee 62B: Diagnostic imaging equipment, of IEC technical committee 62: Electrical equipment in medical practice.

This consolidated version of IEC 60601-2-37 consists of the first edition (2001) [documents 62B/428/FDIS and 62B/440/RVD] and its amendment 1 (2004) [documents 62B/524/FDIS and 62B/542/RVD].

The technical content is therefore identical to the base edition and its amendment and has been prepared for user convenience.

It bears the edition number 1.1.

A vertical line in the margin shows where the base publication has been modified by amendment 1.

Annexes AA and DD form an integral part of this Particular Standard.

Annexes BB, CC, EE, FF, GG and HH are for information only.

In this Particular Standard, the following print types are used:

- requirements, compliance with which can be tested, and definitions: in roman type
- notes, explanations, advice, introductions, general statements, exceptions, and references: in smaller type
- *test specifications: in italic type*
- TERMS USED THROUGHOUT THIS PARTICULAR STANDARD WHICH HAVE BEEN DEFINED IN CLAUSE 2 AND IEC 60601-1: IN SMALL CAPITALS.

The committee has decided that the contents of the base publication and its amendments will remain unchanged until the maintenance result date indicated on the IEC web site under "<http://webstore.iec.ch>" in the data related to the specific publication. At this date, the publication will be

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

A bilingual version of this standard may be issued at a later date.

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INTRODUCTION

In this Particular Standard, safety requirements additional to those in the General Standard are specified for ULTRASONIC DIAGNOSTIC EQUIPMENT.

Guidance and a rationale for the requirements of this Particular Standard are given below.

Knowledge of the reasons for these requirements will not only facilitate the proper application of this Particular Standard but will, in due course, expedite any revision necessitated by changes in clinical practice or as a result of developments in technology.

General guidance and rationale

The approach and philosophy used in drafting this particular standard for safety of ULTRASONIC DIAGNOSTIC EQUIPMENT are consistent with those in current standards in the IEC 60601-2 series that apply to other diagnostic modalities, such as X-ray equipment and magnetic resonance systems.

In each case, the safety standard is intended to require increasing sophistication of output display indicators and/or controls with increasing energy levels in the interrogating field of diagnosis. Thus, for all such diagnostic modalities, it is the responsibility of the OPERATOR to understand the risk of the output of the equipment, and to act appropriately in order to obtain the needed diagnostic information with the minimum risk to the PATIENT.

It should be noted that although UD-3 Rev.1, 1998¹ was developed as a national standard, it has since been referenced by numerous countries worldwide and by all internationally operating manufacturers and test houses; regulatory authorities also follow the standard, as it has become a *de facto* international standard. The material taken from UD-3 Rev.1, 1998 forms only a part of this Particular Standard.

This standard contains normative measurement methodologies. These clauses may be replaced in a future revision by reference to an appropriate future measurement standard.

This standard does not cover ultrasonic therapeutic equipment. Equipment used for the imaging and diagnosis of body structures by ultrasound in conjunction with other medical procedure is covered.

¹ See reference [19] in the Bibliography.

MEDICAL ELECTRICAL EQUIPMENT –

Part 2-37: Particular requirements for the safety of ultrasonic medical diagnostic and monitoring equipment

SECTION ONE: GENERAL

The clauses and subclauses of this section of the General Standard apply except as follows:

1 Scope and object

This clause of the General Standard applies except as follows:

*1.1 Scope

Addition:

This Particular Standard specifies particular safety requirements for ULTRASONIC DIAGNOSTIC EQUIPMENT as defined in 2.1.145.

This standard does not cover ultrasonic therapeutic equipment; however, equipment used for the imaging of body structures by ultrasound in conjunction with therapeutic modalities is covered.

1.2 Object

Replacement:

The object of this Particular Standard is to establish particular requirements for the safety of ULTRASONIC DIAGNOSTIC EQUIPMENT and those aspects thereof which are directly related to safety.

1.3 Particular Standards

Addition:

This Particular Standard amends and supplements a set of IEC publications, hereinafter referred to as the “General Standard”, consisting of

IEC 60601-1:1988, *Medical electrical equipment – Part 1: General requirements for safety* and its Amendments 1 (1991) and 2 (1995)

IEC 60601-1-2:2001, *Medical electrical equipment – Part 1-2: General requirements for safety – Collateral standard: Electromagnetic compatibility – Requirements and tests*

IEC 60601-1-4:1996, *Medical electrical equipment – Part 1-4: General requirements for safety – 4. Collateral Standard: Programmable electrical medical systems* and its Amendment 1 (1999)

The numbering of sections, clauses and subclauses of this Particular Standard corresponds to that of the General Standard. The changes to the text of the General Standard are specified by the use of the following words:

“Replacement” means that the clause or subclause of the General Standard is replaced completely by the text of this Particular Standard.

“Addition” means that the text of this Particular Standard is additional to the requirements of the General Standard.

“Amendment” means that the clause or subclause of the General Standard is amended as indicated by the text of this Particular Standard.

Subclauses or figures which are additional to those of the General Standard are numbered starting from 101, additional annexes are lettered AA, BB, etc., and additional items *aa*, *bb*, etc.

Clauses and subclauses to which there is a rationale are marked with an asterisk (*). These rationales can be found in an informative annex BB. Annex BB should be used in determining the relevance of the requirements addressed, but should never be used to establish additional test requirements.

Where there is no corresponding section, clause or subclause in this Particular Standard, the section, clause or subclause of the General Standard applies without modification.

Where it is intended that any part of the General Standard, although possibly relevant, is not to be applied, a statement to that effect is given in this Particular Standard.

A requirement of this Particular Standard replacing or modifying requirements of the General Standard or of a Collateral Standard takes precedence over the corresponding general requirement(s).

1.3.101 Related international standards

IEC 60529:1989, *Degrees of protection provided by enclosures (IP Code)*

IEC 60788:1984, *Medical radiology – Terminology*

IEC 61102:1991, *Measurement and characterisation of ultrasonic fields using hydrophones in the frequency range 0,5 MHz to 15 MHz*

IEC 61157:1992, *Requirements for the declaration of the acoustic output of medical diagnostic ultrasonic equipment*

IEC 61161:1992, *Ultrasonic power measurement in liquids in the frequency range 0,5 MHz to 25 MHz*

Amendment 1 (1998)

2 Terminology and definitions

This clause of the General Standard applies except as follows:

Additional definitions:

2.1.101

ACOUSTIC ATTENUATION COEFFICIENT

coefficient intended to account for ultrasonic attenuation of tissue between the source and a specified point

Symbol: α

Unit: decibels per centimetre per megahertz, dB cm⁻¹ MHz⁻¹

2.1.102**ACOUSTIC WORKING FREQUENCY**

arithmetic mean of the most widely separated frequencies f_1 and f_2 at which the amplitude of the pressure spectrum of the acoustic signal is 3 dB lower than the peak amplitude [3.4.2 of IEC 61102, modified]

Symbol: f_{awf}

Unit: megahertz, MHz

2.1.103**ATTENUATED OUTPUT POWER**

value of the acoustic OUTPUT POWER after attenuation and at a specified distance from the transducer, and given by

$$P_{\alpha} = P 10^{(-\alpha z f_{awf}/10)}$$

where

α is the ACOUSTIC ATTENUATION COEFFICIENT in decibels per centimetre per megahertz;

z is the distance from the source to the point of interest in centimetres;

f_{awf} is the ACOUSTIC WORKING FREQUENCY in megahertz;

P_{α} is the ATTENUATED OUTPUT POWER in milliwatts;

P is the OUTPUT POWER in milliwatts measured in water.

Symbol: P_{α}

Unit: milliwatts, mW

2.1.104**ATTENUATED PEAK-RAREFACTIONAL ACOUSTIC PRESSURE**

value of the PEAK-RAREFACTIONAL ACOUSTIC PRESSURE after attenuation and at a specified point, and given by

$$p_{ra}(z) = p_r(z) 10^{(-\alpha z f_{awf}/20)}$$

where

α is the ACOUSTIC ATTENUATION COEFFICIENT in decibels per centimetre per megahertz;

z is the distance from the source to the point of interest, in centimetres;

f_{awf} is the ACOUSTIC WORKING FREQUENCY in megahertz;

$p_r(z)$ is the PEAK-RAREFACTIONAL ACOUSTIC PRESSURE measured in water.

Symbol: p_{ra}

Unit: megapascals, MPa

2.1.105**ATTENUATED PULSE-AVERAGE INTENSITY**

value of the ACOUSTIC PULSE-AVERAGE INTENSITY after attenuation and at a specified point, and given by

$$I_{pa,\alpha} = I_{pa}(z) 10^{(-\alpha z f_{awf}/10)}$$

where

α is the ACOUSTIC ATTENUATION COEFFICIENT in decibels per centimetre per megahertz;

z is the distance from the source to the point of interest in centimetres;

f_{awf} is the ACOUSTIC WORKING FREQUENCY, at distance z in megahertz;

$I_{pa}(z)$ is the PULSE-AVERAGE INTENSITY measured in water, in milliwatts per centimetre squared.

Symbol: $I_{pa,\alpha}$

Unit: watts per centimetre squared, W cm⁻²

2.1.106**ATTENUATED PULSE-INTENSITY INTEGRAL**

value of the PULSE-INTENSITY INTEGRAL after attenuation and at a specified point, and given by

$$I_{pi,\alpha} = I_{pi} 10^{(-\alpha z f_{awf}/10)}$$

where

α is the ACOUSTIC ATTENUATION COEFFICIENT in decibels per centimetre per megahertz;

z is the distance from the source to the point of interest in centimetres;

f_{awf} is the ACOUSTIC WORKING FREQUENCY in megahertz;

$I_{pi,\alpha}$ is the ATTENUATED PULSE-INTENSITY INTEGRAL in millijoules per centimetre squared;

I_{pi} is the PULSE-INTENSITY INTEGRAL measured in water in millijoules per centimetre squared.

Symbol: $I_{pi,\alpha}$

Unit: millijoules per centimetre squared, mJ cm⁻²

2.1.107**ATTENUATED SPATIAL-PEAK TEMPORAL-AVERAGE INTENSITY**

value of the SPATIAL-PEAK TEMPORAL AVERAGE INTENSITY after attenuation and at a specified distance z , and given by

$$I_{zpta,\alpha}(z) = I_{zpta}(z) 10^{(-\alpha z f_{awf}/10)}$$

where

α is the ACOUSTIC ATTENUATION COEFFICIENT in decibels per centimetre per megahertz;

z is the distance from the source to the point of interest in centimetres;

f_{awf} is the ACOUSTIC WORKING FREQUENCY in megahertz;

$I_{zpta}(z)$ is the SPATIAL-PEAK TEMPORAL-AVERAGE INTENSITY, at a specified distance z in milliwatts per centimetre squared measured in water.

Symbol: $I_{zpta,\alpha}(z)$

Unit: milliwatts per centimetre squared, mW cm⁻²

2.1.108**ATTENUATED TEMPORAL-AVERAGE INTENSITY**

value of the TEMPORAL-AVERAGE INTENSITY after attenuation and at a specified point, and given by

$$I_{ta,\alpha}(z) = I_{ta}(z)10^{(-\alpha z f_{awf}/10)}$$

where

α is the ACOUSTIC ATTENUATION COEFFICIENT in decibels per centimetre per megahertz;

z is the distance from the source to the point of interest in centimetres;

f_{awf} is the ACOUSTIC WORKING FREQUENCY in megahertz;

$I_{ta,\alpha}(z)$ is the ATTENUATED TEMPORAL-AVERAGE INTENSITY in milliwatts per centimetre squared;

$I_{ta}(z)$ is the TEMPORAL-AVERAGE INTENSITY measured in water in milliwatts per centimetre squared.

Symbol: $I_{ta,\alpha}(z)$

Unit: milliwatts per centimetre squared, mW cm⁻²

2.1.109**BEAM AREA**

area in a specified plane perpendicular to the BEAM-ALIGNMENT AXIS consisting of all points at which the PULSE-INTENSITY INTEGRAL is greater than a specified fraction of the maximum PULSE-INTENSITY INTEGRAL in that plane [3.6 of IEC 61102, modified]

NOTE For measurement purposes the PULSE INTENSITY INTEGRAL can be taken as being proportional to the PULSE PRESSURE-SQUARED INTEGRAL

2.1.110**BEAM ALIGNMENT AXIS**

straight line joining the points of maximum PULSE INTENSITY INTEGRAL measured at several different distances in the far field. For the purposes of alignment, this line may be projected to the face of the ULTRASONIC TRANSDUCER [3.5 of IEC 61102, modified]

2.1.111**BONE THERMAL INDEX**

THERMAL INDEX for applications, such as foetal (second and third trimester) or neonatal cephalic (through the fontanelle), in which the ultrasound beam passes through soft tissue and a focal region is in the immediate vicinity of bone

Symbol: *TIB*

Unit: None

NOTE See annex DD.4.2 and DD.5.2 for methods of determining the BONE THERMAL INDEX.

2.1.112**BOUNDED OUTPUT POWER**

OUTPUT POWER emitted in SCANNING MODE from a region of the active area of the transducer whose width in the scan plane is limited to 1 cm

Symbol: P_1

Unit: milliwatts, mW

2.1.113**BREAK-POINT DEPTH**

value equal to 1,5 times the EQUIVALENT APERTURE DIAMETER, and given by

$$z_{bp} = 1,5 D_{eq}$$

where

D_{eq} is the EQUIVALENT APERTURE DIAMETER.

Symbol: z_{bp}

Unit: centimetres, cm

2.1.114**COMBINED-OPERATING MODE**

mode of operation of an EQUIPMENT which combines more than one DISCRETE-OPERATING MODE [3.6 of IEC 61157]

2.1.115**CRANIAL-BONE THERMAL INDEX**

THERMAL INDEX for applications, such as paediatric and adult cranial applications, in which the ultrasound beam passes through bone near the beam entrance into the body

Symbol: *TIC*

Unit: None

NOTE See annex DD.4.3 for methods of determining the CRANIAL BONE THERMAL INDEX.

2.1.116**DEFAULT SETTING**

specific state of control, the ULTRASONIC DIAGNOSTIC EQUIPMENT will enter upon power-up, new PATIENT select or change from non-foetal to foetal applications

2.1.117**DEPTH FOR BONE THERMAL INDEX**

distance from the plane where the -12 dB OUTPUT BEAM DIMENSIONS are determined along the BEAM ALIGNMENT AXIS to the plane where the product of ATTENUATED OUTPUT POWER and ATTENUATED PULSE-INTENSITY INTEGRAL is maximum

Symbol: z_b

Unit: centimetres, cm

2.1.118**DEPTH FOR SOFT-TISSUE THERMAL INDEX**

distance from the plane where the -12 dB OUTPUT BEAM DIMENSIONS are determined along the BEAM ALIGNMENT AXIS to the plane at which the lower value of the ATTENUATED OUTPUT POWER and the product of the ATTENUATED SPATIAL-PEAK TEMPORAL-AVERAGE INTENSITY and 1 cm^2 is maximized over the distance range equal to, or more than, 1,5 times the EQUIVALENT APERTURE DIAMETER

Symbol: z_s

Unit: centimetres, cm

NOTE In this Particular Standard, the restricted definition of SPATIAL-PEAK TEMPORAL-AVERAGE INTENSITY from 3.49 of IEC 61102 relating to a specified plane is used where SPATIAL-PEAK TEMPORAL-AVERAGE INTENSITY is replaced by ATTENUATED SPATIAL-PEAK TEMPORAL-AVERAGE INTENSITY.

2.1.119**DISCRETE-OPERATING MODE**

mode of operation of ULTRASONIC DIAGNOSTIC EQUIPMENT in which the purpose of the excitation of the ULTRASONIC TRANSDUCER or ULTRASONIC TRANSDUCER element group is to utilize only one diagnostic methodology [3.7 of IEC 61157]