

Evaluation and routine testing in medical imaging departments - Part 2-4: Constancy tests - Hard copy cameras

EESTI STANDARDI EESSÕNA

NATIONAL FOREWORD

See Eesti standard EVS-EN 61223-2-4:2016 sisaldab Euroopa standardi EN 61223-2-4:1994 ingliskeelset teksti.	This Estonian standard EVS-EN 61223-2-4:2016 consists of the English text of the European standard EN 61223-2-4:1994.
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.
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constancy test, evaluation testing, routine testing

ENGLISH VERSION

Evaluation and routine testing in medical imaging
departments

Part 2-4: Constancy tests - Hard copy cameras
(IEC 1223-2-4:1994)

Essais d'évaluation et de
routine dans les services
d'imagerie médicale
Partie 2-4: Essais de constance
Reprographes

(CEI 1223-2-4:1994)

Bewertung und
routinemäßige
Prüfung in Abteilungen für
medizinische Bildgebung
Teil 2-4: Konstanzprüfungen
Bildokumentationssysteme
(IEC 1223-2-4:1994)

This European Standard was approved by CENELEC on 1994-03-08.

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Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the Central Secretariat or to any CENELEC member.

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CENELEC

European Committee for Electrotechnical Standardization
Comité Européen de Normalisation Electrotechnique
Europäisches Komitee für Elektrotechnische Normung

Central Secretariat: rue de Stassart 35, B-1050 Brussels

FOREWORD

The text of document 62B(CO)105, as prepared by Sub-Committee 62B: Diagnostic imaging equipment, of IEC Technical Committee 62: Electrical equipment in medical practice, was submitted to the IEC-CENELEC parallel vote in May 1993.

The reference document was approved by CENELEC as EN 61223-2-4 on 8 March 1994.

The following dates were fixed:

- latest date of publication of
an identical national standard (dop) 1995-03-01
- latest date of withdrawal of
conflicting national standards (dow) 2000-03-01

Annexes designated "normative" are part of the body of the standard. Annexes designated "informative" are given only for information. In this standard, annexes A and ZA are normative and annexes B, C, D and E are informative.

ENDORSEMENT NOTICE

The text of the International Standard IEC 61223-2-4:1994 was approved by CENELEC as a European Standard without any modification.

ANNEX ZA (normative)

OTHER INTERNATIONAL PUBLICATIONS QUOTED IN THIS STANDARD WITH THE REFERENCES OF THE RELEVANT EUROPEAN PUBLICATIONS

This European Standard incorporates by dated or undated reference, provisions from other publications. These normative references are cited at the appropriate places in the text and the publications are listed hereafter. For dated references, subsequent amendments to or revisions of any of these publications apply to this European Standard only when incorporated in it by amendment or revision. For undated references the latest edition of the publication referred to applies.

NOTE : When the international publication has been modified by CENELEC common modifications, indicated by (mod), the relevant EN/HD applies.

IEC Publication	Date	Title	EN/HD	Date
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788	1984	Medical radiology - Terminology	HD 501 S1	1988
1223-1	1993	Evaluation and routine testing in medical imaging departments Part 1: General aspects	-	-
1223-2-1	1993	Part 2-1: Constancy tests - Film processors	-	-
1223-2-2	1993	Part 2-2: Contancy tests - Radiographic cassettes and film changers - Film-screen contact and relative sensitivity of the screen-cassette assembly	-	-
1223-2-3	1993	Part 2-3: Constancy tests - Darkroom safelight conditions	-	-
1223-2-5	1994	Part 2-5: Constancy tests - Image dipslay devices	EN 61223-2-5	1994
1223-2-12	-	Part 2-12: Constancy tests - Film illuminators (under consideration)		

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INTRODUCTION

Some provisions or statements in the body of this part of IEC 1223 require additional information. Such information is presented in annex D, Rationale. An asterisk in the left margin of a clause or subclause indicates the presence of such additional information.

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EVALUATION AND ROUTINE TESTING IN MEDICAL IMAGING DEPARTMENTS –

Part 2-4: Constancy tests – Hard copy cameras

1 Scope and object

1.1 Scope

This part of IEC 1223 applies to HARD COPY CAMERAS producing images on monochrome continuous tone material (such as photographic films and materials sensitive to infrared radiation), and comprising types of cameras using a cathode ray tube, laser beam, or a thermoprinting system, as used in diagnostic imaging systems such as:

- digital radiography;
- digital subtraction angiography;
- imaging in COMPUTED TOMOGRAPHY;
- magnetic resonance imaging;
- ultrasound imaging;
- imaging in NUCLEAR MEDICINE.

* The test methods are based on the use of test patterns.

This standard does not apply to x-y (analogue) recording systems used in NUCLEAR MEDICINE.

This standard is a part of a series of Particular Publications (standards and technical reports) which give methods of tests for the constancy of properties of diagnostic imaging systems as described in IEC 1223-1 (see clause 2).

*1.2 Object

This part of IEC 1223 describes a method to check, in terms of functional parameters, the constancy of the quality of images produced by HARD COPY CAMERAS in order to ensure that the required conditions for producing consistent hard copies are maintained after the calibration and adjustment have been carried out.

The aims of the method are:

- to establish a reference level of performance for the HARD COPY CAMERA when such equipment has been accepted;
- to detect and verify any significant variation in functional parameters which may then require corrective actions.

With regard to the measurements, reference is made to methods described in related publications, which for practical reasons should be carried out prior to the application of the method described in this standard (see clause 2).