# STN

# Pomocné výrobky na zachovanie integrity tkanív imobilných pacientov Časť 1: Všeobecné požiadavky (ISO 20342-1: 2022)

STN EN ISO 20342-1

84 7078

Assistive products for tissue integrity when lying down - Part 1: General requirements (ISO 20342-1:2022)

Táto norma obsahuje anglickú verziu európskej normy. This standard includes the English version of the European Standard.

Táto norma bola oznámená vo Vestníku ÚNMS SR č. 11/22

Obsahuje: EN ISO 20342-1:2022, ISO 20342-1:2022

Oznámením tejto normy sa ruší STN EN ISO 20342-1 (84 7078) z decembra 2019

# EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM

EN ISO 20342-1

August 2022

ICS 11.180.01

Supersedes EN ISO 20342-1:2019

#### **English Version**

# Assistive products for tissue integrity when lying down - Part 1: General requirements (ISO 20342-1:2022)

Produits d'assistance pour l'intégrité des tissus en position allongée - Partie 1: Exigences générales (ISO 20342-1:2022)

Hilfsmittel für die Gewebeintegrität im Liegen - Teil 1: Allgemeine Anforderungen (ISO 20342-1:2022)

This European Standard was approved by CEN on 26 June 2022.

CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without any alteration. Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN-CENELEC Management Centre or to any CEN member.

This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

CEN members are the national standards bodies of Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Türkiye and United Kingdom.



EUROPEAN COMMITTEE FOR STANDARDIZATION COMITÉ EUROPÉEN DE NORMALISATION EUROPÄISCHES KOMITEE FÜR NORMUNG

CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels

Contents	Page
European favorrand	2
European	3

# **European foreword**

This document (EN ISO 20342-1:2022) has been prepared by Technical Committee ISO/TC 173 "Assistive products" in collaboration with Technical Committee CEN/TC 293 "Assistive products and accessibility" the secretariat of which is held by SIS.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by February 2023, and conflicting national standards shall be withdrawn at the latest by February 2023.

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

This document supersedes EN ISO 20342-1:2019.

Any feedback and questions on this document should be directed to the users' national standards body/national committee. A complete listing of these bodies can be found on the CEN website.

According to the CEN-CENELEC Internal Regulations, the national standards organizations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Türkiye and the United Kingdom.

## **Endorsement notice**

The text of ISO 20342-1:2022 has been approved by CEN as EN ISO 20342-1:2022 without any modification.

# INTERNATIONAL STANDARD

ISO 20342-1

Second edition 2022-07

# Assistive products for tissue integrity when lying down —

Part 1: **General requirements** 

Produits d'assistance pour l'intégrité des tissus en position allongée —

Partie 1: Exigences générales





# **COPYRIGHT PROTECTED DOCUMENT**

© ISO 2022

All rights reserved. Unless otherwise specified, or required in the context of its implementation, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office CP 401 • Ch. de Blandonnet 8 CH-1214 Vernier, Geneva Phone: +41 22 749 01 11 Email: copyright@iso.org Website: www.iso.org

Published in Switzerland

Contents			Page
Fore	eword		<b>v</b>
Intr	oductio	on	<b>v</b> i
1	Scon	e	1
2	-	native references	
3		ns and definitions	
4		eral requirements and safety	7
	4.1 4.2	General requirementsIntended use	
	4.2	4.2.1 General requirements	
		4.2.2 Consideration regarding intended use	
		4.2.3 Intended use statement	
	4.3	APTI risk management	
	4.4	APTI usability	
		4.4.1 General	
	4.5	Design controls	
	4.6	Clinical evaluation	
	4.7	Foreseeable misuse	
	4.8	Test conditions	
	4.9	Lifting and carrying means	10
5	Safet	ty requirements	10
	5.1	Requirements for information supplied by the manufacturer	10
		5.1.1 General	
		5.1.2 APTI traceability	
		5.1.3 Education and training 5.1.4 Pre-sale information	
		5.1.5 User information	
		5.1.6 Service information and inspection	
		5.1.7 Labelling	
		5.1.8 Marking of user weight and maximum load	
	<b>F</b> 0	5.1.9 Packaging	
	5.2	APTI that can be dismantled	
		5.2.1 General requirements 5.2.2 Small parts	
		5.2.3 Fasteners and connections	
	5.3	Resistance to corrosion	
	5.4	Noise and vibration	
	5.5	Sound audible acoustic energy	
	5.6 5.7	Default indicatorsFeedback	
6		ımability	
	6.1 6.2	General Flammability	
	6.3	Moulded parts used as enclosures for electrical equipment	
7		• •	
7	Mect 7.1	nanical safetyPrevention of traps for the human body	17 17
	7.1 7.2	Safety of moving and folding parts	
	7.2	V-shaped openings	
	7.4	Surfaces, corners, edges and protruding parts	
	7.5	Folding and adjusting mechanisms	19
	7.6	Instability hazard	
	7.7	Temperature of parts that come into contact with human skin	20

7.8 Ergonomic principles	21
7.9 Additional consideration	21
Safety of electrical equipment	21
8.1 General electrical requirements	
8.2 Electromagnetic compatibility	22
8.2.1 General	22
8.2.2 Emissions	22
8.2.3 Immunity	
8.3 Liquid ingress	22
8.5 Hold to run activation	23
8.6 Emergency stop functions	23
Biocompatibility	24
9.2 Animal tissue	
Contamination	24
10.3 Cross infection and microbial contamination	
Annex A (informative) General information	
B (informative) Environmental and consumer related guidance	30
C (informative) Periodic inspection	34
	Safety of electrical equipment 8.1 General electrical requirements 8.2 Electromagnetic compatibility 8.2.1 General 8.2.2 Emissions 8.2.3 Immunity 8.2.4 Power frequency magnetic field immunity 8.3 Liquid ingress 8.4 Interruption of power supply/supply mains to an APTI 8.5 Hold to run activation 8.6 Emergency stop functions  Biocompatibility 9.1 Biocompatibility and toxicity 9.2 Animal tissue  Contamination 10.1 Liquid ingress 10.2 Cleaning and disinfection 10.3 Cross infection and microbial contamination  A (informative) General information

#### **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 <a href="www.iso.org/directives">www.iso.org/directives</a>).

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 <a href="https://www.iso.org/patents">www.iso.org/patents</a>).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT), see <a href="https://www.iso.org/iso/foreword.html">www.iso.org/iso/foreword.html</a>.

This document was prepared by Technical Committee ISO/TC 173, *Assistive products*, in collaboration with the European Committee for Standardization (CEN) Technical Committee CEN/TC 293, *Assistive products and accessibility*, in accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).

This second edition cancels and replaces the first edition (ISO 20342-1:2019), which has been technically revised.

The main changes are as follows:

- the Scope was clarified;
- Clause 2 was updated;
- <u>Clause 3</u> was updated;
- <u>subclause 7.3</u> about V-shaped openings was amended;
- <u>subclause 7.7</u> and Table 4 were amended (regarding surface temperature);
- the bibliography was updated.

A list of all parts in the ISO 20342 series can be found on the ISO website.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at <a href="https://www.iso.org/members.html">www.iso.org/members.html</a>.

## Introduction

This document addresses Assistive Products for Tissue Integrity (APTI). As some devices can be used/reused in more than one application environment, different requirements and test methods can apply to the same APTI, depending on the application environment.

APTI play a very important role in the prevention and treatment of pressure injuries. Another important role in the prevention and treatment of pressure injury is the clinical practice and the clinical evaluation. Guidance can be found in the NPUAP/EPUAP/PPPIA Guidelines<sup>[24]</sup>.

Surfaces applied on operating theatre tables can also impact in the process of patient management and might need to be taken into consideration. It should be recognized however, patient stability and specialist equipment used during an operation often create conflicting priorities to those of an APTI.

Using this document, clinicians and manufacturers should consider the impact of other items (including additional APTI) used in conjunction with an APTI on tissue integrity and safety.

This document only covers general requirements to ensure safety of users.

# Assistive products for tissue integrity when lying down —

# Part 1:

# **General requirements**

## 1 Scope

This document specifies general requirements and related test methods that are relevant to assistive products for tissue integrity (APTI) in the lying position in different application environments such as hospitals, home care and institutions. This document applies to the safety of APTI that are intended to remain in situ during periods of lying, and to prevent and/or treat pressure injuries.

This document covers a range of different lying support surfaces intended to be used in combination with the appropriate support platform (adjustable included) or as a whole integrated system.

This document does not apply to medical beds.

This document also covers assistive products primarily intended for tissue integrity for changing a lying position and assistive products for maintaining a lying position.

This document does not apply to lying support surfaces used in combination with incubators or operating/surgical tables.

It also covers safety and performance test methods to ensure protection against injuries to the user.

This document addresses the combination of a full body support surface and an adjustable mattress support platform. It also covers safety and performance test methods to ensure protection against injuries to the user.

This document specifies requirements and test methods for APTI within the following classifications of ISO 9999:2022:

04 33 06 Assistive products for tissue integrity when lying down such as but not limited to

- mattresses and mattress overlays for pressure injury prevention, and
- mattress coverings for pressure injury prevention mattresses.

12 31 03 Assistive products for sliding and turning such as but not limited to the following:

Devices for changing position or direction of a person using sliding or turning techniques. The only products included are those intended to be used in a lying position and remain in situ as part of the lying support surface. They are the following:

- sliding products that glide one way and lock the other way;
- sheets and underlays in flexible materials with low friction;
- fabric sold by the metre, cut as required for repositioning use;
- powered turning product;

This excludes sliding boards unless the product is intended to be left in situ.

09 07 06 Positioning pillows, positioning cushions and positioning systems such as but not limited to

leg positioners,

- arm positioners, and
- multipurpose body positioners.

18 12 15 Bedding such as but not limited to

draw sheets.

#### 2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements 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 554, Standard atmospheres for conditioning and/or testing — Specifications

ISO 3746, Acoustics — Determination of sound power levels and sound energy levels of noise sources using sound pressure — Survey method using an enveloping measurement surface over a reflecting plane

ISO 9614-1, Acoustics — Determination of sound power levels of noise sources using sound intensity — Part 1: Measurement at discrete points

ISO 10993-1, Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process

ISO 13732-1, Ergonomics of the thermal environment — Methods for the assessment of human responses to contact with surfaces — Part 1: Hot surfaces

ISO 14155, Clinical investigation of medical devices for human subjects — Good clinical practice

ISO 14971, Medical devices — Application of risk management to medical devices

ISO 15223-1, Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements

ISO 22442-1, Medical devices utilizing animal tissues and their derivatives — Part 1: Application of risk management

IEC 60601-1:2005+AMD1:2012+AMD2:2020, Medical electrical equipment — Part 1: General requirements for basic safety and essential performance

IEC 60601-1-2, Medical electrical equipment — Part 1-2: General requirements for basic safety and essential performance — Collateral standard: Electromagnetic compatibility — Requirements and tests

IEC 60601-1-6, Medical electrical equipment — Part 1-6: General requirements for basic safety and essential performance — Collateral standard: Usability

IEC 60601-1-8, Medical electrical equipment — Part 1-8: General requirements for basic safety and essential performance — Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems

IEC 60601-1-11, Medical electrical equipment — Part 1-11: General requirements for basic safety and essential performance — Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment

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

IEC 60695-11-10, Fire hazard testing — Part 11-10: Test flames — 50 W horizontal and vertical flame test methods

IEC 61000-3-2, Electromagnetic compatibility (EMC) — Part 3-2: Limits — Limits for harmonic current emissions (equipment input current  $\leq$ 16 A per phase)

IEC 61000-3-3, Electromagnetic compatibility (EMC) — Part 3-3: Limits — Limitation of voltage changes, voltage fluctuations and flicker in public low-voltage supply systems, for equipment with rated current  $\leq$ 16 A per phase and not subject to conditional connection

IEC 61000-4-3, Electromagnetic compatibility (EMC) — Part 4-3: Testing and measurement techniques — Radiated, radio-frequency, electromagnetic field immunity test

IEC 61000-4-8, Electromagnetic compatibility (EMC) — Part 4-8: Testing and measurement techniques — Power frequency magnetic field immunity test

IEC 61672-1, Electroacoustics — Sound level meters — Part 1: Specifications

IEC 61672-2, Electroacoustics — Sound level meters — Part 2: Pattern evaluation tests

IEC 62366-1, Medical devices — Part 1: Application of usability engineering to medical devices

IEC 60601-2-35, Medical electrical equipment — Part 2-35: Particular requirements for the basic safety and essential performance of heating devices using blankets, pads or mattresses and intended for heating in medical use

EN 716-2:2017, Furniture — Children's cots and folding cots for domestic use — Part 2: Test methods

EN 1041, Information supplied by the manufacturer of medical devices

CISPR 11, Industrial, scientific and medical (ISM) radio-frequency equipment — Electromagnetic disturbance characteristics — Limits and methods of measurement

European Commission, MEDDEV 2.7/1 CLINICAL EVALUATION: A GUIDE FOR MANUFACTURERS AND NOTIFIED BODIES UNDER DIRECTIVES 93/42/EEC and 90/385/EEC

koniec náhľadu – text ďalej pokračuje v platenej verzii STN